Abnormal glycogen chain length pattern, not hyperphosphorylation, is critical in Lafora disease

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Abstract

Lafora disease (LD) is a fatal progressive epilepsy essentially caused by loss-of-function mutations in the glycogen phosphatase laforin or the ubiquitin E3 ligase malin. Glycogen in LD is hyperphosphorylated and poorly hydrosoluble. It precipitates and accumulates into neurotoxic Lafora bodies (LBs). The leading LD hypothesis that hyperphosphorylation causes the insolubility was recently challenged by the observation that phosphatase-inactive laforin rescues the laforin-deficient LD mouse model, apparently through correction of a general autophagy impairment. We were for the first time able to quantify brain glycogen phosphate. We also measured glycogen content and chain lengths, LBs, and autophagy markers in several laforin- or malin-deficient mouse lines expressing phosphatase-inactive laforin. We find that: (i) in laforin-deficient mice, phosphatase-inactive laforin corrects glycogen chain lengths, and not hyperphosphorylation, which leads to correction of glycogen amounts and prevention of LBs; (ii) in malin-deficient mice, phosphatase-inactive laforin confers no correction; (iii) general impairment of autophagy is not necessary in LD. We conclude that laforin’s principle function is to control glycogen chain lengths, in a malin-dependent fashion, and that loss of this control underlies LD.

Keywords glycogen chain length; glycogen phosphorylation; Lafora disease; laforin; malin
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Introduction

Lafora disease (LD) is a recessively inherited teenage-onset severe and fatal progressive myoclonus epilepsy. Its etiopathological hallmark is the Lafora body (LB), present throughout the brain in profuse ever-increasing numbers and sizes (Lafora & Glueck, 1911; Minassian, 2001). LBs are overwhelmingly composed of an abnormally formed glycogen (Sakai et al., 1970) and likely result from gradual precipitation, aggregation and accumulation of abnormal glycogen.

Normally glycogen consists of highly and regularly branched roughly sphere-shaped molecules, each composed of up to 55,000 glucose units. It is mainly synthesized by coordinated actions of glycogen synthase (GS), which elongates glucan chains forming α1-4 interglucosidic linkages, and glycogen branching enzyme, which detaches a part of each newly formed chain and reattaches it to another glucan chain through an α1-6 linkage. Glycogen is essentially degraded by glycogen phosphorylase and glycogen debranching enzyme. Glucan chains of glycogen have a wide chain length distribution with an average of ~13 units (Roach et al., 2012; Nitschke et al., 2013). High frequency of branching points and relatively short chains are critical for glycogen solubility. They prevent neighbouring chains from forming double helices, which otherwise exclude water, reduce the water solubility of the interacting chains and finally lead the entire molecule to precipitate. Incidentally, the main component of plant starch amylpectin is insoluble precisely because it is composed of longer chains that through clustered arrangement of branch points form crystalline areas of double helices (Roach et al., 2012; Cenci et al., 2014). Glycogen and amylpectin possess phosphate esters bound to a very small number of glucose units, in glycogen’s case at carbon positions C2, C3 and C6 (Tagliabracci et al., 2011; Nitschke et al., 2013). The kinase(s) responsible for glycogen phosphorylation are not known, except that GS itself has been reported to be involved in C3 phosphorylation through what so far has been considered an undesirable side reaction of the enzyme (Tagliabracci et al., 2011; Contreras et al., 2016).
The abnormal glycogen constituting LBs has three characteristics: (i) its chain length distribution contains longer chains than normal glycogen, (ii) its phosphate levels are elevated, and (iii) its quantity is high, the latter reflecting the profuse deposition as LBs (Tagliabracci et al., 2008; Nitschke et al., 2013). LD is mainly caused by loss-of-function mutations in either the EPM2A (laforn) or EPM2B (malin) gene, and knockout of either in mouse recapitulates the disease (Minassian et al., 1998; Ganesh et al., 2002; Chan et al., 2003; Turnbull et al., 2010). Laforn is a glycogen phosphatase, its absence resulting in progressive glycogen hyperphosphorylation (Worby et al., 2006; Tagliabracci et al., 2008). Malin is a ubiquitin E3 ligase (Gentry et al., 2005), which in cell culture experiments, in a laforin-dependent fashion, diminishes both GS and a protein involved in GS activation, R5/PTG, through proteasomal degradation (Vilchez et al., 2007; Worby et al., 2008). Malin, again in a laforin-dependent manner, also diminishes a second protein that participates in GS activation, R6, in this case through autophagic degradation (Rubio-Villena et al., 2013). Laforn and malin form a functional complex, the former (but not the latter) possessing a carbohydrate binding domain through which the complex is targeted to glycogen (Gentry et al., 2005; Lohi et al., 2005).

From the above results, two main hypotheses were advanced to explain the abnormal glycogen formation in LD: (i) absence of laforin or malin leads to increased GS activity, which outpacing branching leads to the formation of glycogen with long chains promoting water insolubility (Vilchez et al., 2007); and (ii) laforin deficiency results in non-removal of erratically introduced phosphate and thus hyperphosphorylation of glycogen, which via an unknown mechanism alters its structure promoting glycogen precipitation (Tagliabracci et al., 2008; Roach, 2015). Both hypotheses, however, were challenged by more recent findings. (i) Tissues from laforin and malin knockout mice were shown to have unaltered GS activity (Tagliabracci et al., 2008; DePaoli-Roach et al., 2010). (ii) When phosphatase-inactive laforin was overexpressed in laforin knockout mice, LBs no longer formed and the mice were rescued from LD (Gayarre et al., 2014). The latter result was particularly provocative. It suggested that laforin’s chief and only established physiological function (glycogen dephosphorylation) and the major and distinguishing pathological feature of LD glycogen (progressive hyperphosphorylation) were irrelevant to LB formation and LD.

Gayarre et al., however, left a number of critical questions unaddressed before the field could make a decided paradigm shift away from the centrality of glycogen phosphate in LB formation and LD towards exploring alternative disease mechanisms. They did not measure glycogen phosphate. It can, therefore, not be excluded that the overexpressed in vitro-inactive laforin had retained some phosphatase activity in vivo or that there was a small amount of wild-type (WT) laforin somehow still expressed in the transgenic mice. Laforn’s phosphatase activity is catalysed by a cysteine residue (C265 in mouse, C266 in human), which Gayarre and colleagues had mutated to serine to inactivate the enzyme. Recently, crystal structures of laforin were determined (Raththagala et al., 2015; Sankhala et al., 2015) suggesting the presence of a putative second phosphatase site in laforin (C168 in mouse, C169 in human). The C168 site could be active in vivo, especially when overexpressed, and explain the rescue by laforin mutated only at C265.

The other two features specific to LD glycogen, chain length distribution and quantity, were also outside the scope of the above study and had not been analysed. As such, conclusions about the mechanism conferring rescue from LBs remained tentative. It was possible, for example, that the rescued mice did accumulate abnormal glycogen, which, coated with a large amount of hydrophilic protein (overexpressed laforin), remained soluble and did not coalesce into pathologically observable LBs.

In this work, we address all these points by characterizing glycogen in both laforin- and malin-deficient mice as affected by overexpressed WT and phosphatase-inactive laforin. Our results combine with previous studies to allow the proposal of a unifying hypothesis of LB pathogenesis in both laforin- and malin-deficient LD.

Results

Amounts and molecular parameters of muscle glycogen in Epm2a<sup>−/−</sup> mice overexpressing phosphatase-inactive laforin

We quantified total glycogen from skeletal muscle from the actual mice used in the Gayarre et al study (Gayarre et al., 2014). In Epm2a<sup>−/−</sup> mice, muscle glycogen content was fivefold increased, as in previous studies (Tagliabracci et al., 2008). In Epm2a<sup>−/−</sup> mice overexpressing WT laforin (Epm2a<sup>−/−</sup>.Laf), muscle glycogen content was normal. In Epm2a<sup>−/−</sup> mice overexpressing phosphatase-inactive laforin (Epm2a<sup>−/−</sup>.C265SLaf), muscle glycogen content was also normal (Fig 1A), indicating that the prevention of LBs (which are aggregates of insoluble glycogen) by phosphatase-inactive laforin (Gayarre et al., 2014) is indeed due to the prevention of generating insoluble glycogen, as opposed to merely maintaining the excess abnormal glycogen in a soluble state.

We measured total glycogen phosphate in the above tissues, which in Epm2a<sup>−/−</sup> mice was 14-fold increased, as in previous studies (Tagliabracci et al., 2008). Levels were normal in Epm2a<sup>−/−</sup>.Laf mice, but in Epm2a<sup>−/−</sup>.C265SLaf mice glycogen phosphate was just as high as in Epm2a<sup>−/−</sup> mice (Fig 1B). This indicates that phosphatase-inactive laforin indeed has no phosphatase activity in vivo and does not normalize the hyperphosphorylation characteristic of LD glycogen. This in turn indicates that glycogen hyperphosphorylation does not cause the generation of insoluble glycogen.

Measurement of total glycogen phosphate requires relatively large amounts of purified glycogen (see Materials and Methods). Of the three glucosyl carbons phosphorylated in glycogen, C6 phosphorylation is the only one for which a sensitive method has been developed that allows site-specific quantification. Moreover, C6 phosphate can be measured in very small amounts of glycogen, and as all three phosphorylation sites are proportionally high in LD (DePaoli-Roach et al., 2015; Roach, 2015), it can be used as a measure of total glycogen phosphorylation. While sufficient amounts of glycogen can be purified from skeletal muscle for total glycogen phosphate measurement, this is currently unfeasible for brain glycogen, where C6 phosphorylation only would be quantified as an indicator of the total. Leading up to the brain studies (next sections), we measured C6 glycogen phosphate in skeletal muscle and found that it is, as expected, like total glycogen phosphate,
increased in both the Epm2a\(^{-/-}\) and Epm2a\(^{-/-}\).C265SLaf mice (Fig 1C).

Besides glycogen amount and phosphate content, the third feature that distinguishes LD from WT glycogen is chain length distribution. We measured skeletal muscle glycogen chain lengths in the above genotypes and found that the chain length distribution in Epm2a\(^{-/-}\) mice is abnormal with a significantly increased proportion of longer chains, while in Epm2a\(^{-/-}\).C265SLaf mice (and of course Epm2a\(^{-/-}\).Laf mice) the chain length distribution is normal, indicating that phosphatase-inactive laforin rescues the glycogen chain length abnormality characteristic of LD glycogen (Fig 1D).

To summarize, phosphatase-inactive laforin does not correct the hyperphosphorylation of LD muscle glycogen, but does normalize the glycogen’s chain length distribution preventing abnormal glycogen accumulation and LB formation.

**Brain glycogen in Epm2a\(^{-/-}\) and Epm2b\(^{-/-}\) mouse lines overexpressing phosphatase-inactive laforin**

The implications of the Gayarre et al (2014) study and our above results are major: Laforin’s phosphatase function is dispensable and glycogen hyperphosphorylation is not pathogenic. These results run counter to a leading LD hypothesis, in which glycogen phosphorylation is considered a damaging side product of GS activity that needs to be cleared by laforin (Tagliabracci et al, 2011; Raththagala et al, 2015; Roach, 2015; Contreras et al, 2016; Turnbull et al, 2016). To rule out the possibility of an artefact in the Gayarre et al mice, we generated a whole new set of Epm2a\(^{-/-}\), and this time also Epm2b\(^{-/-}\) mice overexpressing phosphatase-inactive laforin. Phosphatase-inactive human laforin (C266SLaf) was used, in contrast to the phosphatase-inactive murine laforin (C265SLaf) used by Gayarre.
et al (2014). We did so by crossing WT mice overexpressing C266SLaf available in our laboratory (Chan et al, 2004) with Epm2a/−/− (Ganesh et al, 2002) and Epm2b/−/− (Turnbull et al, 2010) mice. The resultant transgenic animals are termed Epm2a/−/−.C266SLaf and Epm2b/−/−.C266SLaf, respectively.

We simultaneously analysed glycogen and LBs from this new line and from the Gayarre et al Epm2a/−/−.C265SLaf mice, focusing on brain, the disease-relevant organ. Consistency of findings in Gayarre et al and our mice would confirm phosphatase-inactive laforin’s effect on glycogen and obviate results arising from any particularities of each mouse line studied. The differences between the mouse lines are transgene species (as mentioned above), presence or absence of a tag on the transgene protein, expression level, and mouse background (Chan et al, 2004; Gayarre et al, 2014). We had previously generated the C266SLaf mice (Chan et al, 2004) in an attempt to create an LD mouse model through a dominant-negative approach. The phosphatase-inactive laforin (C266SLaf) was supposed to outcompete WT laforin’s phosphatase activity and lead to an effective loss of the endogenous laforin function. In fact, the resultant mice did have a few LBs and we reported this mouse as a mild, only-pathological, model of LD, with no neurological phenotype (Chan et al, 2004). Compared to the laforin (and malin) knock-out mice (Epm2a/−/− and Epm2b/−/−, respectively) (Ganesh et al, 2002; Turnbull et al, 2010), the number of LBs occurring in this model is orders of magnitude lower (Fig 2), and negligible for purposes of the present experiments.

**Glycogen and LB analyses**

Brain glycogen content was threefold to fourfold increased in both Epm2a/−/− mouse lines and in Epm2b/−/− mice, as previously published (Tagliabracci et al, 2008; DePaoli-Roach et al, 2010; Turnbull et al, 2010). Brain glycogen content was normal in Epm2a/−/−.C265SLaf and Epm2a/−/−.C266SLaf mice, but remained elevated in Epm2b/−/−.C266SLaf mice (Fig 3A and B). Thus, overexpressed phosphatase-inactive laforin corrects the abnormal glycogen accumulation in laforin-deficient LD but not in malin-deficient LD. The latter indicates that the rescue in the former is not through some mechanism related to overexpressed protein (somehow physically solubilizing insoluble glycogen) as otherwise the same correction would be expected in the malin-deficient background.

![Figure 2. Phosphatase-inactive laforin prevents Epm2a/−/− but not Epm2b/−/− from LB formation.](source)

Representative PASD-stained sections of hippocampi from WT, Epm2a/−/− and Epm2b/−/− mice in the absence or presence of the human C266S mutated laforin transgene (C266SLaf). Scale bar equals 100 μm.

Source data are available online for this figure.
Consistent with the rescue of the abnormal brain glycogen accumulation in the Epm2a<sup>-/-</sup>.C265SLaf and Epm2a<sup>-/-</sup>.C266SLaf mice, there were no LBs in the brains of these mice. LBs were, however, still present in the Epm2b<sup>-/-</sup>.C266SLaf mice, and to the same extent as in their Epm2b<sup>-/-</sup> controls (Fig 2), again indicating that phosphatase-inactive laforin prevents LB formation in laforin-deficient but not malin-deficient LD.

We next measured brain glycogen phosphate (via C6 phosphate quantification). Due to particularities of brain tissue, this (and the below glycogen chain length analyses) required extensive new method development, detailed in the Materials and Methods section, and the results represent the first time these analyses have been possible in brain. Brain glycogen phosphate was threefold and twofold elevated in Epm2a<sup>-/-</sup> and Epm2b<sup>-/-</sup> mice, respectively, similar to muscle glycogen phosphate (Tagliabracci et al., 2008; DePaoli-Roach et al., 2010; Turnbull et al., 2010). This increase was not corrected in any of the three phosphatase-inactive laforin expressing mouse genotypes (Epm2a<sup>-/-</sup>.C265SLaf, Epm2a<sup>-/-</sup>.C266SLaf and Epm2b<sup>-/-</sup>.C266SLaf) (Fig 4). Taken together, the results in this section clearly indicate that hyperphosphorylation of glycogen is not pathogenic nor is it causative of abnormal glycogen accumulation and LB formation.

Similar to previous findings in muscle glycogen (Fig 1D; see also Nitschke et al., 2013), but even more pronounced, brain glycogen chain length distributions were abnormal, significantly shifted towards longer chains, in Epm2a<sup>-/-</sup> and Epm2b<sup>-/-</sup> mice. They were normalized in Epm2a<sup>-/-</sup>.C265SLaf and Epm2a<sup>-/-</sup>.C266SLaf mice (indistinguishable from WT) and remained abnormal in Epm2b<sup>-/-</sup>.C266SLaf (Fig 5). This demonstrates that phosphatase-inactive laforin can correct the brain glycogen chain length abnormality in laforin-deficient LD, but not in malin-deficient LD. These results tease apart, for the first time, the three characteristics of

![Figure 3. Phosphatase-inactive laforin rescues Epm2a<sup>-/-</sup>, but not Epm2b<sup>-/-</sup> abnormal glycogen accumulation in the brain.](image)

A Brain glycogen levels in the presence or absence of WT (Laf) or C265 mutated (C265Laf) murine laforin transgene (tissues from the Gayarre et al. 2014 study).

B Brain glycogen levels of WT, Epm2a<sup>-/-</sup> and Epm2b<sup>-/-</sup> mice in the presence or absence of human C266 mutated laforin (C266Laf) (tissues from the new mice generated in the present study).

Data information: Data are presented as means of at least five biological replicates ± SEM. Asterisks, statistical significance by ANOVA and post hoc analyses (*P < 0.05,**P < 0.01,***P < 0.001, Appendix Table S2).

![Figure 4. Phosphatase-inactive laforin does not rescue glycogen hyperphosphorylation in Epm2a<sup>-/-</sup> or Epm2b<sup>-/-</sup> brain.](image)

A Brain glycogen carbon C6 phosphate levels in the presence or absence of WT (Laf) or C265 mutated (C265Laf) murine laforin transgene.

B Brain glycogen C6 phosphate levels of WT, Epm2a<sup>-/-</sup> and Epm2b<sup>-/-</sup> mice in the presence or absence of human C266 mutated laforin (C266Laf).

Data information: Data are presented as means of at least five biological replicates ± SEM. Asterisks, statistical significance by ANOVA and post hoc analyses (*P < 0.05,**P < 0.01,***P < 0.001, Appendix Table S3).
Abnormal glycogen chain length underlies LD

Abnormal LD glycogen (glycogen accumulation, hyperphosphorylation, abnormal chain length distribution) and indicate that the altered glycogen chain length distribution, and not hyperphosphorylation, correlates with abnormal glycogen accumulation and LB formation. Moreover, phosphatase-inactive laforin’s ability to rescue the chain length abnormality (and the abnormal glycogen accumulation/LBs) depends on the presence of malin, as this correction does not occur in the malin-deficient mice (Epm2b−/−.C266SLaf). This implies that laforin (irrespective of its phosphatase domain) and malin impact glycogen chain length distribution and prevent LB formation and LD conversely. It directly links both proteins to glycogen structure in vivo and corroborates the close laforin–malin partnership previously suggested by cell culture overexpression experiments (Gentry et al., 2005; Lohi et al., 2005; Vilchez et al., 2007).

**Effect of phosphatase-inactive laforin on LD-associated general defect in autophagy**

LC3 becomes lipidated (i.e. LC3-I converts to LC3-II) as it joins proliferating autophagosomes during autophagy, and therefore decreased LC3-II indicates decreased autophagy. p62 directs ubiquitinated proteins to autophagy, and thus, accumulation of p62 indicates failing autophagy. LD mouse brains were shown to have reduced LC3-II and increased p62, suggesting that LD mice have impaired autophagy and that this defect underlies LB accumulation (Criado et al., 2012). Gayarre and colleagues reported normalization of LC3-II levels in their Epm2a−/−.C265Laf mice, suggesting that phosphatase-inactive laforin rescues laforin knockout mice by correcting the autophagic defect (Gayarre et al., 2014). We tested brain LC3-II and p62 in the new mice generated for the present study and found no differences in either protein between WT, Epm2a−/−, Epm2b−/−, Epm2a−/−.C266Laf, and Epm2b−/−.C266Laf genotypes (Fig 6). These results imply that impaired autophagy may not be a common feature to all LD models and, therefore, it is not a strict requirement for LD.

**Discussion**

The report that phosphatase-inactive laforin rescues laforin-deficient LD (Gayarre et al., 2014) runs contrary to the widely accepted view that the disease is caused by lack of laforin-mediated clearance of erratic glycogen phosphorylation (Raththagala et al., 2015; Roach,
Our results show that glycogen hyperphosphorylation does not correlate with LB formation. Hyperphosphorylation is shared by Epm2a\(^{−/−}\)Epm2b\(^{−/−}\), Epm2a\(^{−/−}\).C266Slaf (rescued), Epm2a\(^{−/−}\).C266Slaf (not rescued), and Epm2b\(^{−/−}\).C266Slaf (not rescued) genotypes and can therefore not be the cause of LB formation (Fig 7). The glycogen chain length abnormality, on the other hand, is shared by the Epm2a\(^{−/−}\), Epm2b\(^{−/−}\), and Epm2b\(^{−/−}\).C266Slaf (not rescued) mice, strictly correlating with LB formation and glycogen accumulation (Fig 7). This together with the well-established facts that longer glucan chains are less water soluble as they form stable double helices and that isolated LBs consist of poorly branched glucans (Sakai et al., 1970; Gidley & Bulpin, 1987; Hejazi et al., 2009) strongly supports the notion that chain length abnormality is the actual cause of glycogen precipitation and, over time, accumulation into LBs.

Phosphatase-inactive laforin rescues glycogen precipitation and accumulation in the Epm2a\(^{−/−}\).C266Slaf and Epm2a\(^{−/−}\).C266Slaf but not Epm2b\(^{−/−}\).C266Slaf mice. As the relevant difference between these genotypes is that the latter lacks malin and has an abnormal glycogen chain length pattern, it is safe to conclude that laforin confers its effect strictly together with malin. Previous publications afford insights into the mechanisms through which malin could prevent the generation of glycogen molecules with abnormally long chains. As mentioned, in cell culture experiments malin has consistently been shown to ubiquitinate, in a laforin-dependent fashion, GS and proteins involved in GS activation (PTG/R5 and R6), targeting them to proteasomal or autophagic degradation (Vilchez et al., 2007; Worby et al., 2008; Rubio-Villena et al., 2013). On the surface, this mechanism contrasted with the lack of overall elevated GS activity in LD mouse tissues (Tagliabracci et al., 2008; DePaoli-Roach et al., 2010). These results can, however, be reconciled if the action of the laforin–malin complex is restricted to a small subset of glycogen molecules. Glycogen is heterogeneous, in both size (Wanson & Drochmans, 1968; Marchand et al., 2002; Nitschke et al., 2013) and chain length distribution (i.e., some glycogen molecules contain longer chains) (Palmer et al., 1983; Nitschke et al., 2013). The laforin–malin system would be required only for those molecules/molecule regions in which glucan chains approach thresholds to insolubility. Consistent
with this is the observation that laforin’s carbohydrate binding domain has greater affinity to longer rather than shorter glucan chains (Chan et al., 2004; Lohi et al., 2005; Dias et al., 2016). A reduced chain elongation would prevent progression towards precipitation, and since the activity would be localized, its absence would not result in detectably increased global GS activity.

Aguado et al. (2010) reported that their murine Epm2a<sup>−/−</sup> LD model has an impairment of autophagy, and Criado et al. (2012) that this impairment presents prior to the appearance of detectable glycogen accumulations/LBs. Gayarre et al. (2014) showed that phosphatase-inactive laforin restores autophagy as it rescues LD and proposed that laforin–malin prevents LB formation via autophagy. In contrast and in line with recent publications (Irimia et al., 2015; Wang et al., 2016), we do not detect presence of an autophagic impairment in the Epm2a<sup>−/−</sup> and Epm2b<sup>−/−</sup> animals used to prepare the new transgenic mice generated here and can, therefore, not confirm autophagy correction by laforin–malin. Additionally, other recent work showed that disturbances of autophagy markers present in LD are correctable by genetic reduction of GS activity (Durán et al., 2014), suggesting that the autophagy impairment may be secondary to the glycogenosis. Notwithstanding, it is entirely possible that laforin–malin acts locally to eliminate, through autophagy, individual glycogen molecules that are “at risk” of precipitation. In fact, malin was recently shown to be able to mediate K63-linked ubiquitination, which can result in p62-mediated substrate targeting to autophagy (Sanchez-Martin et al., 2015). Since laforin preferentially binds glycogen molecules with longer chains (see above), it is conceivable that malin could ubiquitinate laforin through K63 ubiquitination and target laforin-bound “at-risk” glycogen molecules to autophagy.

As our results show that phosphorylation is not causative of glycogen precipitation and accumulation, what, if any, are the functions of glycogen phosphates? In starch metabolism, one established function of monophosphate esters of amylopectin is, through their high charges and hydrophilia, to separate amylopectin chains from each other, rendering them fully hydrated, and thus accessible to degrading enzymes (Edner et al., 2007; Hejazi et al., 2008; Köttig et al., 2009; Zeeman et al., 2010). As glycogen and amylopectin have a shared chemical basis (both consist of branched alpha-glucan chains), effects of covalent modifications, such as phosphorylation, are expected to be similar. Phosphorylation should therefore enhance glycogen solubility and may represent a third mechanism to protect against precipitation of long-chained “at-risk” molecules. It is well established that neither starch-degrading exo-amyloses nor glycogen phosphorylase can proceed beyond a phosphorylated glucosyl residue (Lomako et al., 1994; Köttig et al., 2009; Cenci et al., 2014; Emanuelle et al., 2016). Laforin’s phosphorylase role may be, similar to starch phosphatases (Gentry et al., 2007; Köttig et al., 2009), to remove the phosphates to allow reshortening of chains in “at-risk” molecules and during glycogen digestion. Consistent with this is the recent demonstration that glycogen dephosphorylation by laforin occurs principally during glycogen degradation (Irimia et al., 2015). In the absence of laforin’s phosphatase activity, cycles of partial glycogen digestion and resynthesis would lead to the time-dependent phosphate accumulation that characterizes LD glycogen.

In summary, our results combine with recent work (Gayarre et al., 2014) to suggest that laforin’s principle LD relevant function is mediated through malin and directed to preventing glycogen molecules with hyperextended chains. In the absence of either protein, some glycogen molecules at a time precipitate and gradually over time aggregate and amass into LBs, which, reaching a certain threshold profusion (at ~14 years of age in humans), initiate, and then drive the progressive myoclonus epilepsy.

Materials and Methods

Mouse lines, tissue collection, and histological stain for LBs

Tissues from Epm2a<sup>−/−</sup>·C266SLaf mice and their controls (see Results) were the same as utilized in Gayarre et al. (2014; C57BL/6J). These mice were 9–12 months old and of both sexes. Epm2a<sup>−/−</sup>·C266SLaf and Epm2b<sup>−/−</sup>·C266SLaf mice and controls (see Results) were obtained following the crossing of both Epm2a<sup>−/−</sup> (Ganesh et al., 2002; C57BL/63J) and Epm2b<sup>−/−</sup> (Turnbull et al., 2010; C57BL/6J) mice with the C266SLaf mice described previously (Chan et al., 2004; 129SvJ). Mice were housed with environmental enrichment in ventilated cages at 20–22°C, fed a high-quality, commercially available diet, water being automatically delivered. Brains of male and female mice were harvested after cervical dislocation of at least six 9- to 12-month-old mice from each genotype, split along the sagittal plane, one hemisphere immediately frozen in liquid nitrogen, the other fixed in 10% [w/v] formalin. The latter was embedded in paraffin, sectioned and stained using the periodic acid–Schiff diastase method (PASD; Chan et al., 2004). All animal procedures were approved by The Centre for Phenogenomics Animal Care Committee and are in compliance with the CCAC Guidelines and the OMAFRA Animals for Research Act.

Glycogen determination

Glycogen content was determined essentially as described previously (Suzuki et al., 2001). Briefly, frozen tissue was ground in liquid nitrogen, boiled in 30% [w/v] KOH, and precipitated thrice in 67% ethanol and 15 mM LiCl (at least 1 h at −30°C). After redissolving in water, aliquots were digested with amyloglucosidase (Megazyme) in 80 mM sodium acetate buffer pH 4.5. Liberated glucose was determined enzymatically (Lowry & Passonneau, 1972).

Glycogen purification from muscle for phosphate determination

Quantification of total glycogen-bound phosphate requires high glycogen purity, because its relative amounts are very low and non-glycogen derived phosphoesters, for example, in nucleic acids, can severely affect the determination. Extracted muscle glycogen was thus further purified as previously published (Tagliabracci et al., 2007). Subsequently, glycogen was repeatedly precipitated in 67% ethanol and 15 mM LiCl (at least 1 h at −30°C) until the absence of nucleic acids could be confirmed by UV–vis spectra of glycogen solutions at concentrations of ~100 mM glucose equivalents.

Total glycogen phosphate quantification in muscle glycogen

Total glycogen-bound phosphate was quantified as orthophosphate following enzymatic hydrolysis of all phospho-monoesters. To facilitate complete orthophosphate release, highly purified muscle
glycogen was incubated with phosphatase together with two glycogen hydrolysing enzymes. In a final volume of 65 μl, the reaction mixture contained 100 or 200 μg glycogen, 2.5 U Antarctic phosphatase (NEB), 0.1 U α-amylase (Roche), 2 U amyloglucosidase (Megazyme), 50 mM bis–Tris–propane pH 6.0, 1 mM MgCl₂, and 0.1 mM ZnCl₂. Following overnight incubation at 37°C, 40 μl of this reaction mixture, and orthophosphate standards for calibration, was used to quantify ortho-phosphatase enzymatically as described previously (Nelson & Kaufman, 1987). Following the addition of equal volumes of a buffer pH 7 (100 mM imidazole–HCl, 10 mM MgCl₂, 5 mM EDTA), in a total volume of 80 μl, orthophosphate, via G1P, is converted to G6P using 0.02 mg rabbit phosphorylase a (Sigma), 67 μg purified rabbit muscle glycogen, and 1 U phosphoglucomutase (Sigma). Fluorescence (ex340/em470) was read until stable, first, after the addition of 150 μl 20 mM Tris pH 8, 60 μM NADP, and again, after the addition of 0.4 μl G6PDH (Roche) in 5 μl 20 mM Tris pH 8. The G6PDH-dependent increase in fluorescence was used to calculate the orthophosphate amounts in samples originally containing 100 or 200 μg glycogen. Glucose-based phosphate amounts of the latter two measurements for each sample were averaged and deviated < 5% indicating that the enzymatic orthophosphate release was complete.

**Glycogen purification from brain**

Initial attempts to quantify C6 phosphate in brain glycogen were unsuccessful when following an isolation procedure similar to that used for muscle glycogen. Unexpected high values of apparent glucosyl C6 phosphate residues in hydrolysates of brain glycogen indicated the presence of a contaminant that greatly affected the measurements. The ratio of contaminant to glycogen was much higher in samples with low glycogen content, such as WT, suggesting that the contaminant is unrelated to glycogen but derives from other constituents of brain (whose glycogen content appears to be low as compared to other organs). Further experiments confirmed that the contaminant is very likely glucose 6-phosphate released during acid hydrolysis from a glycan-containing compound other than glycogen. This contamination is relevant only for tissues with low glycogen content, such as brain. Systematic experiments were conducted to remove the contaminant while retaining the little glycogen using spare WT mouse brains. After the extraction of glycogen from 100 to 400 mg ground frozen brain tissue in 30% KOH and three consecutive precipitations in 67% ethanol and KOH and three consecutive precipitations in 67% ethanol, 15 mM LiCl (see extraction of muscle glycogen for glycogen determination), the following procedure was established for all brain glycogen samples analysed in this study. The resulting pellet, containing glycogen, most likely proteins, and the contaminant, was resuspended in 100 μl water and subsequently extracted with 900 μl 2:1 methanol:chloroform [v/v] by vigorous mixing, incubation for 25 min at room temperature (RT) on a rotator, and 10 min at 85°C. After cooling to RT, the suspension was centrifuged for 20 min at 20°C and 16,000 × g, the supernatant discarded and the pellet dried briefly at 85°C. The extraction procedure was repeated twice, followed by resuspending the dry pellet in 300 μl water. After 20-min incubation at 95°C with intermittent agitation, the solubilized glycogen was recovered in the supernatant (S1) obtained by centrifugation at RT (16,000 × g for 20 min). The pellet was extracted with 100 μl 0.5 M KOH at 95°C for 45 min with intermittent agitation. Subsequently, the extract, cooled to RT, was centrifuged (20 min, RT, 16,000 × g), and the resultant supernatant (S2) was combined with the solubilized glycogen (S1). The combined supernatants (S1 + S2) were subjected to several rounds of repeated ethanol precipitation (1,400 μl 67% ethanol, 15 mM LiCl, 70 min at ~30°C), 20 min centrifugation at 4°C and 16,000 × g, carefully discarding the supernatant, and resolubilization of the pellet in water followed by 10-min heat treatment (95°C, intermittent agitation). The ethanolic supernatant harbours the contaminant, the amount of which tends to be smaller with each round of precipitation. Glycogen pellets were considered to be contaminant free when the apparent G6P amount in the last ethanolic supernatant was below the detection limit. Using the enzymatic cycling method, the contaminant (apparent G6P) was quantified after drying the ethanolic supernatants (SpeedVac), dissolving the dry contaminant pellets in water, and subsequent acid hydrolysis (see below). Glycogen concentration was determined after dissolving the purified glycogen pellet as described above. As the number of sequential precipitations was adjusted to the persistence of the contaminant in the ethanolic supernatant, glycogen recovery varied but in most cases was between 50 and 80%.

**Glycogen carbon C6 phosphate determination**

Aliquots of purified muscle or brain glycogen preparations (see above) were subjected to acid hydrolysis (3 h, 95°C, 0.7 M HCl) and subsequently neutralized using 5 M KOH. G6P was determined alongside G6P standards using the enzymatic cycling assay previously described (Nitschke et al., 2013). G6P amounts were based on glucose amounts that were liberated during acid hydrolysis and determined enzymatically (Lowry & Passonneau, 1972).

**Determination of glycogen chain length distributions**

Purified muscle or brain glycogen (10–20 μg dry weight) was completely debranched by overnight incubation with 200 U isoamylase (Sigma) in 110 μl 10 mM sodium acetate at pH 5 and 37°C. Following heat inactivation (10 min at 95°C) and centrifugation (10 min at 20,000 × g), 90 μl of the supernatant was applied to high-performance anion exchange chromatography with pulsed amperometric detection (HPAEC-PAD; ThermoFisher, ICSS5000). Separation of the oligoglucan chains was achieved on a CarboPac PA100 column and guard combination (ThermoFisher) at a constant flow rate of 1 ml/min with an elution profile of the two eluents A (150 mM NaOH) and B (150 mM NaOH, 500 mM sodium acetate) as follows: injection after 15 min 95% [v/v] A, 5% [v/v] B equilibration; 5 min 95% A, 5% B; 9 min linear gradient to 70% A, 30% B; 9 min linear gradient to 55% A, 45% B; 37 min linear gradient to 33% A, 67% B; 10 min 100% B. Chromatograms were analysed with Dionex Chromelene software (version 7.2). For each chain length (degree of polymerization), relative peak areas were determined, which were then averaged among at least five biological replicates allowing the calculation of standard deviations (SD). Note that purified glycogen from all available brain samples of the genotype *Epm2a<sup>-/-</sup>-Laf* was pooled to obtain sufficiently large amounts of glycogen for the chain length distribution analysis. SD at each chain length as a measure of biological variance for this genotype was estimated by multiplying the average SD of the other genotypes
(WT, Epm2a<sup>−/−</sup>, Epm2a<sup>−/−</sup>, C265SLaf) by two. As variances in relative peak areas between those genotypes are mostly equal, the estimated biological variance should include 95% of samples of the Epm2a<sup>−/−</sup>-Laf genotype.

Analysis of autophagy markers

For the detection of LC3 in brain tissue, a membrane fraction enriched in autophagosomes was prepared as described previously (Chu et al, 2009), and for p62 after extracting soluble protein as in Wang et al (2007); 15 µg total protein was separated on 14% SDS-PAGE using a Laemmli buffer system. Proteins of interest were detected after transfer to PVDF membranes using primary antibodies NB100-2220 (LC3, Novus Biologicals), P0067 (p62, Sigma), sc-25778, and sc-11397 (GAPDH and calnexin, respectively, Santa Cruz), all at a 1:1,000 dilution.

Statistical analyses

Except where stated, all biochemical analyses were conducted in at least five biological replicates. Standard deviations (SD) or standard error of mean (SEM) was calculated as indicated. Averages of multiple groups of biological replicates were compared using one-way analysis of variance (ANOVA) followed by post hoc analysis using homoscedastic Student’s t- or heteroscedastic Welch’s tests, both unpaired, two-tailed, and with Holm–Bonferroni correction. The adequate post hoc test was selected based on equal or unequal variances of groups tested with F-tests. Significance levels are indicated by asterisks (*<i>P</i> < 0.05, **<i>P</i> < 0.01, ***<i>P</i> < 0.001). For glycogen chain length distribution analyses, significance between genotypes was tested for each individual degree of polymerization.

Expanded View for this article is available online.

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Author contributions

BAM, FN, and MS conceived the study in collaboration with SRC. Experiments were conducted by FN, PW, and XZ with help from AMP, LI, LJL, and PB. Data were analysed by FN, EEC, and PW. The manuscript was written by FN, BAM, and MS with critical review from MAS, EEC, and SRC.

Conflict of interest

The authors declare that they have no conflict of interest.

References

Abnormal glycogen chain length underlies LD neuronal disease, ataxia, myoclonus epilepsy and impaired behavioral disruption of the Epm1 laforin. Proc Natl Acad Sci USA 102: 2765 – 2770


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