A Drosophila screen identifies NKCC1 as a modifier of NGLY1 deficiency

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Abstract N-Glycanase 1 (NGLY1) is a cytoplasmic deglycosylating enzyme. Loss-of-function mutations in the NGLY1 gene cause NGLY1 deficiency, which is characterized by developmental delay, seizures, and a lack of sweat and tears. To model the phenotypic variability observed among patients, we crossed a Drosophila model of NGLY1 deficiency onto a panel of genetically diverse strains. The resulting progeny showed a phenotypic spectrum from 0 to 100% lethality. Association analysis on the lethality phenotype, as well as an evolutionary rate covariation analysis, generated lists of modifying genes, providing insight into NGLY1 function and disease. The top association hit was Ncc69 (human NKCC1/2), a conserved ion transporter. Analyses in NGLY1-/- mouse cells demonstrated that NKCC1 has an altered average molecular weight and reduced function. The misregulation of this ion transporter may explain the observed defects in secretory epithelium function in NGLY1 deficiency patients.

Introduction

NGLY1 deficiency (OMIM 615273) is a rare, autosomal recessive disorder caused by loss-of-function mutations in the NGLY1 gene. Patients with NGLY1 deficiency have a variety of symptoms, including developmental delay, seizures, liver dysfunction, central and peripheral nervous system abnormalities, sweat gland abnormalities, and a lack of tears (alacrima) (Enns et al., 2014; Lam et al., 2017). While the first NGLY1 deficiency patient was only recently identified (Need et al., 2012), there have been rapid research advances thanks to the support of two patient organizations (NGLY1.org and Grace Science Foundation). Even though a great deal has been learned about the genetic disorder in a short amount of time, there are currently no cures or approved treatments for NGLY1 deficiency.

The NGLY1 gene encodes the N-Glycanase protein (NGLY1). NGLY1 functions as part of the Endoplasmic Reticulum (ER) Associated Degradation (ERAD) pathway as evidenced by its association with other ERAD components (Katiyar et al., 2005; McNeill et al., 2004; Park et al., 2001). The ERAD pathway retrotranslocates misfolded proteins from the ER lumen to the cytoplasm where they are degraded by the proteasome (reviewed in Qi et al., 2017). NGLY1 is localized to the cytoplasm where it is thought to remove N-linked glycans from misfolded proteins prior to their degradation (Hirsch et al., 2003). Recent evidence suggests that this deglycosylation is required for retrotranslocation for at least some protein substrates (Galeone et al., 2020). Nevertheless, it remains unclear whether NGLY1 is required to deglycosylate all misfolded proteins, or just a subset, or if it is...
necessary for protein degradation at all. It has been shown that model substrates can be degraded regardless of glycosylation state (Hirsch et al., 2003; Kario et al., 2008). While a recent report showed that ER stress markers were increased in NGLY1 -/- MEFs (Galeone et al., 2020), other experiments such as RNAi knockdown (KD) of NGLY1 in Drosophila (Owings et al., 2018) and loss of NGLY1 function in mouse, rat, and human cells (Asahina et al., 2020; Mueller et al., 2020; Tambe et al., 2019) have shown no evidence of ER stress. ER stress is often observed when there are mutations in proteins that are necessary for ERAD due to the accumulation of misfolded proteins in the ER. It may be that NGLY1 is not necessary for ERAD, or it is involved in a non-canonical ERAD function, or it may be deglycosylating cytoplasmic proteins for an entirely different purpose. These hypothesized functions are not mutually exclusive.

NGLY1 has been shown to deglycosylate various exogenous model substrates such as TCR-\(\alpha\) (Hirsch et al., 2003) and RNaseB (Kario et al., 2008). To identify endogenous substrates several mass spectrometry experiments have been performed (Fujihira et al., 2017; Hosomi et al., 2016; Maynard et al., 2020; Zolekar et al., 2018). Yet, the first high-confidence substrate of NGLY1 deglycosylation, NRF1 (gene: NFE2L1), was discovered in a Caenorhabditis elegans genetic screen (Lehrbach and Ruvkun, 2016). NRF1 mediates a proteasome 'bounce-back' response. NRF1 is constitutively degraded by the proteasome through the ERAD pathway, until the proteasome is inhibited or overwhelmed by protein load. During this proteasome stress, NRF1 accumulates and is deglycosylated by NGLY1 (Tomlin et al., 2017). Rather than targeting the protein for degradation, the deglycosylation activates NRF1 by converting asparagine to aspartic acid residues (Lehrbach et al., 2019). NRF1 can then be imported into the nucleus to act as a transcription factor for proteasome subunits. The lack of NRF1 activation in NGLY1-deficient patients likely explains some of the disorder’s symptoms such as motor dysfunction (Kobayashi et al., 2011) and cognitive deficits (Lee et al., 2011). Recently, it was found that the protein BMP4 is deglycosylated by NGLY1 when overexpressed in either Drosophila or mammalian cells (Galeone et al., 2020). BMP4 is a signaling molecule and could explain several of the developmental symptoms of NGLY1 deficiency. However, not all symptoms can be explained by these two targets and therefore there is a pressing need to identify and validate more substrates of NGLY1 deglycosylation.

In addition to discovering new NGLY1 targets, there is a need to understand how background genetic variants affect the number and severity of symptoms in patients. While the majority of patients harbor two complete loss-of-function mutations in NGLY1 (He et al., 2015), there are many symptoms such as seizures and scoliosis that are only reported in a subset of the patients (Enns et al., 2014). All patients experience developmental delay, but it ranges from slightly below average IQ to completely non-verbal (Lam et al., 2017). This variability based on background genetics was also observed in the lab when an NGLY1 deficiency mouse model was crossed onto an out-bred mouse strain which partially rescued the lethality of the model (Fujihira et al., 2017). In order to identify components of the genetic background that affect the range of symptoms and severity of disease, we have utilized a collection of genetically diverse Drosophila strains known as the Drosophila Genetic Reference Panel (DGRP) (Mackay et al., 2012). By crossing a fly model of NGLY1 deficiency onto the panel, we recapitulated the variable phenotype seen in the human population. Here, we report the results of this cross and a list of candidate modifier genes derived from the genome-wide association (GWA) of the cross. To contextualize the candidate modifier list, we also performed an evolutionary rate covariation (ERC) analysis to identify genes that are co-evolving with NGLY1. Together these two genetic analyses have generated a list of genes that (1) may explain some of the variation seen between NGLY1 patients, (2) may encode proteins that physically interact with NGLY1 in ERAD or other cellular processes, and (3) may be direct deglycosylation targets of NGLY1. The top candidate modifier gene from the GWA is NKCC1, a conserved Na/K/Cl ion co-transporter. We found that NKCC1 modifies multiple phenotypes in Drosophila, and in NGLY1 -/- mammalian cells, NKCC1 displays abnormal average molecular weight and has reduced activity. The misregulation of NKCC1 likely explains several prominent secretory epithelium-related phenotypes observed in NGLY1 deficiency patients.
Results

Variation in lethality associated with NGLY1 deficiency

We crossed a fly model of NGLY1 deficiency (Pngl in flies, hereon referred to as NGLY1) onto 163 strains of the DGRP in order to assess the effect of natural variation on loss of NGLY1 function. We have previously validated this NGLY1 deficiency model where an NGLY1 RNAi reduces NGLY1 transcript by >95% when driven by the ubiquitous Tubulin-GAL4 driver transgene (Tubulin > NGLY1 RNAi) (Owings et al., 2018). In order to cross a ubiquitously expressed NGLY1 RNAi onto the DGRP strains in a single cross, we needed to overcome the lethality associated with loss of NGLY1 (Owings et al., 2018). To do this, a Tubulin-GAL80 transgene, which represses the effect of GAL4, was crossed onto the Tubulin > NGLY1 RNAi background, such that RNAi is not expressed and flies from this parent strain are healthy and viable (Figure 1A). This donor strain was crossed to each DGRP strain to generate F1 flies that have both ubiquitous KD of NGLY1 and 50% of their genome from each respective DGRP strain (Figure 1B). In this way, analyzing the F1 progeny was a direct measurement of the dominant effect of the DGRP genetic variants on the NGLY1 KD phenotype.

The phenotypic outcome used for this screen was adult survival through eclosion. We simply scored all adults emerging from each cross in the four balancer categories: CyO, Sb, double balanced, or no balancers, with the no balancer flies being the NGLY1 KD. If no lethality is present, Mendelian segregation should produce the expected 1:1:1:1 ratio of the genotypes. Given that there is a very low level of lethality associated with each balancer, the largest balancer class is the closest to the expected, and was used to calculate the ratio of lethality for NGLY1 KD. Results of the screen reveal that survival to adulthood was strongly influenced by DGRP genetic background (Figure 1B; Supplementary file 1), with proportion of surviving flies ranging from 0.0 to 0.967. Survival to adulthood was not correlated with efficiency of RNAi, as there was no difference in KD efficiency in flies from either end of the phenotypic distribution (low surviving: 92.0% ± 3.7; high surviving: 94.8% ± 3.7; p=0.4). There was no correlation between proportion of surviving flies and the absolute number of flies in the balancer class (R² = 0.02; p=0.14), indicating that the ratio is not driven by the number of the balancer control flies.

Genome-wide association

We hypothesized that the observed variable survival to adulthood in NGLY1 KD flies was due to the underlying genetic variation in the DGRP. Therefore, genome-wide association (GWA) analysis of the fully sequenced DGRP was used to identify variants that associated with NGLY1 KD survival. We used a linear mixed model to test 2,007,145 single-nucleotide polymorphisms (SNPs; Supplementary file 2). We recognize that our study suffers from a multiple

![Figure 1](https://doi.org/10.7554/eLife.57831)

**Figure 1.** Lethality phenotype of NGLY1 knockdown is highly modifiable by strain background. (A) Drosophila cross for NGLY1 knockdown in each Drosophila genetic reference panel (DGRP) strain. (B) Proportion of NGLY1 knockdown flies surviving for each cross was calculated based on the number eclosing compared to the expected number. Expected number was based on the largest control balancer class for each cross. The online version of this article includes the following figure supplement(s) for figure 1:

**Figure supplement 1.** Ncc69 expression is not correlated with survival.
testing problem, making it difficult to interpret the role of any single SNP identified. Instead, the location of SNPs was used to identify candidate modifier genes. This type of approach has worked well in the past for other disease models (Ahlers et al., 2019; Chow et al., 2013a; Chow et al., 2013b; Lavoy et al., 2018; Palu et al., 2019) and provides an unbiased list of candidate genes that can be functionally tested for interactions with NGLY1.

At a nominal p-value of p<10^{-5}, 125 variants are associated with survival to adulthood. Of these 125 variants, 21 fall outside of a gene region (+/- 1 kb from the 5’ or 3’ UTRs) (Supplementary file 3). The remaining 104 variants map to 61 protein coding candidate genes (Table 1). Eighty-five of these 104 variants are in noncoding regions (UTRs, introns, or upstream or downstream) and 19 are in coding regions. Of these 19, 12 are synonymous changes and 7 are nonsynonymous (exp, hiw, CG30048, SP2353, CG31690, Hrd3, and blue). When multiple testing correction is applied to all the variants, the top 12 remain significant. Nine of these SNPs reside in an intron of the Ncc69 gene. All nine SNPs are in strong linkage disequilibrium with each other, which is quite unusual for the DGRP. When we analyzed Ncc69 expression levels using previously published RNAseq data from the DGRP (Everett et al., 2020), we found there was no correlation with survival (Figure 1—figure supplement 1).

Gene ontology (GO) enrichment analysis of the 61 candidate genes did not identify enrichment in any biological process or molecular function. However, GO enrichment was identified for the cellular component categories ‘cell periphery’ (GO:0071944; 19/61; q < 0.0016) and ‘plasma membrane’ (GO:0005886; 17/61; q < 0.004). At least 12/61 candidate genes are involved in protein homeostasis: three are involved in ERAD (CG8405, CG42383, and Hrd3), six are ER resident or membrane proteins (CG33012, CG30043, CG31690, CG4341, Hrd3, and CG8405), four are involved in ubiquitination or the proteasome (hiw, blue, CG42383, and Hrd3), one regulates heatshock responses (Hsromega), and one regulates N-linked glycosylation (sff).

Three of the identified ERAD genes already have known interactions with NGLY1. CG8405 is the Drosophila ortholog of human TMEM259, which physically interacts with NGLY1 in co-immunoprecipitation experiments (Zhu et al., 2017). CG42383 is the Drosophila ortholog of human NSFL1C (cofactor p47). NSFL1C and NGLY1 interact with the VCP/P97 AAA-ATPase complex involved in delivering misfolded proteins from the ERAD complex to the proteasome for degradation (Kondo et al., 1997; McNeill et al., 2004). Hrd3 is the Drosophila ortholog of SEL1L. SEL1L is a component of the ERAD complex required for retrotranslocation of misfolded proteins from the ER to the cytoplasm for degradation. Recently, the C. elegans orthologs of NGLY1 and SEL1L were both identified as modifiers of NRF1 function (Lehrbach and Ruvkun, 2016). These candidate genes are a proof-of-principle that this screen has identified functionally relevant modifiers.

The four candidate genes that encode ER resident proteins are particularly interesting. CG31690 and CG4341 are both Drosophila orthologs of human TMTC2, an ER transmembrane protein that regulates calcium homeostasis. CG33012 and CG30043 are both Drosophila orthologs of human ERMP1, an ER metalloprotease. It is striking that in both cases, both Drosophila orthologs of a single human gene were identified as candidate modifiers, suggesting that the function of TMTC2 and ERMP1 might be particularly important for NGLY1 lethality. It is not clear how these genes might modify NGLY1 lethality, but their physical localization to the ER makes sense and suggests a possible role in protein homeostasis as well.

Gene set enrichment analysis

The rank-order candidate modifiers identified in our GWA ignores the majority of the association data by only considering one variant at a time, rather than all the variants associated with a particular gene. Therefore, we performed a gene set enrichment analysis (GSEA), which assigns each variant to the closest gene and generates a per gene metric for p-value enrichment (Palu et al., 2019; Subramanian et al., 2005). Given a defined set of genes annotated with a certain GO function, GSEA determines whether the members of that set are randomly distributed throughout the ranked list or if they are found primarily at the top or bottom of that list. We identified 21 gene sets positively associated with the ranked gene list (>5 genes;>0.25 enrichment score; p<0.05) (Figure 2; Supplementary file 4). These data suggest that these GO categories are closely linked to NGLY1 activity and variation in individual genes in these categories contribute to the distribution of lethality observed in our screen.
Table 1. Candidate modifier genes identified from GWA. Rank order of candidate genes was established based on the most significant associated SNP in the respective gene.

| Rank order | Gene | FBgn     | Human ortholog | Periphery/membrane | Proteostasis |
|------------|------|----------|----------------|--------------------|--------------|
| 1          | exp  | FBgn0033668 | —              | no                 | no           |
| 2          | Ncc69| FBgn0036279 | NKCC1/2        | yes                | no           |
| 3          | CG5888| FBgn0028523 | —              | yes                | no           |
| 4          | CG16898| FBgn0034480 | —              | no                 | no           |
| 5          | bru3 | FBgn0264001 | CELF2/3/4/5/6  | no                 | no           |
| 6          | CG31690| FBgn0051690 | TMTC2          | yes                | no           |
| 7          | CG7227| FBgn0031970 | SCARB1         | no                 | no           |
| 8          | CR44997| FBgn0266348 | —              | no                 | no           |
| 9          | rgn  | FBgn0261258 | Many           | no                 | no           |
| 10         | M6   | FBgn0037092 | GPM6A          | yes                | no           |
| 11         | Rab26| FBgn0086913 | RAB26          | yes                | yes          |
| 12         | ObpS6i| FBgn0043532 | —              | no                 | no           |
| 13         | S-HT1A | FBgn0004168 | HTR1A          | yes                | no           |
| 14         | CG33012| FBgn0053012 | ERMP1          | no                 | yes          |
| 15         | rst  | FBgn0003285 | —              | yes                | no           |
| 16         | CR43926| FBgn0264547 | —              | no                 | no           |
| 17         | CG7337| FBgn0031374 | WDR62          | no                 | no           |
| 18         | hiw  | FBgn0030600 | MYCBP2         | yes                | yes          |
| 19         | fid  | FBgn0259146 | TRMT9B         | no                 | no           |
| 20         | nmo  | FBgn0011817 | NLK            | no                 | no           |
| 21         | Sirup| FBgn0031971 | SDHAF4         | no                 | no           |
| 22         | tst  | FBgn0039117 | SKIV2L         | no                 | no           |
| 23         | Mdr50 | FBgn0010241 | many           | yes                | no           |
| 24         | Cpr49Aa| FBgn0050045 | —              | no                 | no           |
| 25         | COX7C| FBgn0040773 | COX7C          | no                 | no           |
| 26         | Eip63E| FBgn0005640 | CDK14/15       | yes                | no           |
| 27         | CG30048| FBgn0050048 | PKD1           | no                 | no           |
| 28         | CG15040| FBgn0030940 | —              | no                 | no           |
| 29         | SP2353| FBgn0034070 | EGFLAM         | no                 | no           |
| 30         | Mf   | FBgn0038294 | —              | no                 | no           |
| 31         | ome  | FBgn0259175 | many           | no                 | no           |
| 32         | esn  | FBgn0263934 | PRICKLE1-3     | no                 | no           |
| 33         | haf  | FBgn0261509 | many           | no                 | no           |
| 34         | dally| FBgn0263930 | GPC3/5         | yes                | no           |
| 35         | robo2| FBgn0002543 | ROBO1/2/3/4   | no                 | no           |
| 36         | Gyc32E| FBgn0010197 | NPR1/2         | yes                | no           |
| 37         | CG8170| FBgn0033365 | many           | no                 | no           |
| 38         | CG8405| FBgn0034071 | TMEM259        | no                 | yes          |
| 39         | scaf | FBgn0033033 | —              | yes                | no           |
| 40         | borr | FBgn0032105 | CDCA8          | yes                | no           |
| 41         | Syx7 | FBgn0267849 | STX7/12        | yes                | no           |
| 42         | DIP-delta| FBgn0085420 | many           | yes                | no           |
| 43         | cv-c | FBgn0285955 | DLC1           | yes                | no           |

Table 1 continued on next page
Some of the most significantly enriched categories such as nuclear transport, rRNA processing and signal transduction are broad categories that could have wide reaching implications for NGLY1 function. These processes, however, are difficult to test and require long-term investigation, beyond the scope of this study. Circadian rhythm, on the other hand, is a specific and testable category. The enriched category for circadian rhythm function contains a number of genes that directly modulate circadian rhythm, including, \textit{clock}, \textit{period}, \textit{timeless}, and \textit{cycle}. We hypothesized that if variation in circadian rhythm function modifies lethality associated with loss of NGLY1 function then NGLY1 must affect the circadian rhythm. To test this, we knocked down \textit{NGLY1} in the LNv pacemaker neurons in the central nervous system using the \textit{Pdf-GAL4} driver (Renn et al., 1999) and assayed rhythmicity of locomotor activity in constant darkness over 8 days in \textit{Drosophila} Activity Monitors (DAM). Compared to \textit{Pdf-GAL4}/+ and UAS-NGLY1\textsubscript{RNAi} controls, flies with \textit{NGLY1} KD exhibited a significantly longer period length (Figure 2—figure supplement 1), supporting the idea that NGLY1 function can affect sleep. Indeed it has been reported that patients with NGLY1 deficiency experience disturbed sleep patterns (Enns et al., 2014; Lam et al., 2017).

**Evolutionary rate covariation**

Many of the GWA and GSEA results are intriguing, but appear far removed from the currently known functions of NGLY1. We hypothesized that we could contextualize some of the gene and network results by discovering which of them might be co-evolving with NGLY1. Therefore, we employed evolutionary rate covariation (ERC) analysis (Wolfe and Clark, 2015). Gene pairs with high ERC values have correlated rates of substitution and are thought to function together in protein complexes or related pathways. ERC analysis identified hundreds of protein-coding genes with integrated ERC scores exceeding two with NGLY1 (column ‘sumnlogpvbest’ in Supplementary file 5). Of the 38 GWA candidates that have human orthologs, two were found in this group with elevated NGLY1 ERC values, CG4374 (\textit{many}) and \textit{esn} (\textit{PRICKLE1}). While this overlap is not enriched above background, co-evolution suggests that these two genes might have a particularly important interaction with NGLY1.

GO analysis was used to determine if there was enrichment in any biological pathways among NGLY1 co-evolving genes. Among the top enriched pathways were ‘rRNA/ncRNA/ribosome
biogenesis/metabolism-related functions’ and ‘functions related to nuclear pore complex’. This is particularly exciting as both processes overlap with the top GO enrichment categories observed in the GSEA analysis, suggesting that the same functional categories that contribute to variation in NGLY1-related lethality also appear to contain genes that co-evolve with NGLY1. The rRNA processing category (GO:0006364) contained six genes overlapping between the two analyses. This overlap is higher than expected, given two equally sized random groups of genes (GSEA: 23 genes; ERC: 37 genes; p<2.6 × 10⁻¹²). Among other ncRNA-related enriched GO categories from the ERC analysis are ncRNA metabolic process (GO:0034660), ncRNA processing (GO:0034470), tRNA metabolic process (GO:0006399), ribosome biogenesis (GO:0042254), and tRNA modification (GO:0006400). The functions related to the nuclear pore included nuclear export (GO:0051168), nuclear pore organization (GO:0006999), nuclear transport (GO:0051169), and nuclear pore complex assembly (GO:0051292). While there was no overlap between ERC and GSEA for exact nuclear pore function categories, GSEA results were enriched for functions related to nuclear import (GO:0042306 and GO:0006606). Together, these observations suggest previously unknown roles for NGLY1 in ncRNA and nuclear pore functions.

NGLY1 deficiency is part of a larger category of disorders known as Congenital Disorders of Glycosylation (CDG), with NGLY1 being the only protein that actually deglycosylates substrates. There are 151 known CDG genes. GO analysis of the ERC results identified enrichment of the GPI anchor biosynthetic process, which contains several of these CDG genes, leading us to believe that other CDG genes may have been ERC hits. However, the 151 CDG genes do not fall into one functional

Figure 2. Gene set enrichment analysis. Top significant ontological categories identified by GSEA. p-values are indicated by red-to-blue gradient, with red the lowest p-values and blue the highest p-values. Gene number identified in each category is indicated by the size of the circle.

The online version of this article includes the following figure supplement(s) for figure 2:

Figure supplement 1. NGLY1 knockdown causes circadian rhythm defect.
GO category, therefore, we manually curated the ERC list and identified 26 CDG genes that co-evolve with NGLY1 (see color coding in Supplementary file 5). This represents a significant overlap above what is expected by chance ($p<7.6 \times 10^{-10}$). In particular, 5 of the 21 genes involved in N-linked glycosylation and 9 of the 29 genes involved in GPI-anchor biogenesis are co-evolving with NGLY1. The remaining 10 genes are spread across the CDG functional spectrum. The identification of a number of CDG genes that co-evolve with NGLY1, suggests that NGLY1 function might be important to the broader glycosylation pathways.

Genetic interaction between NGLY1 and Ncc69 in Drosophila

While these genetic analyses revealed many promising modifying and co-evolving genes which should be investigated, we began by investigating Ncc69 because it was the top hit with a human ortholog in our GWA analysis. Further, Ncc69 is a glycoprotein, making it a potential target of NGLY1 deglycosylation. Ncc69 has two mammalian orthologs, NKCC1 and NKCC2. While Ncc69 is ubiquitously expressed in Drosophila, NKCC1 (gene: SLC12A2) is highly expressed in secretory epithelia and NKCC2 (gene: SLC12A1) is primarily expressed in the kidney (Delpire and Gagnon, 2018). In all cases, the protein is a 12-pass transmembrane cation-chloride co-transporter that brings Na$^+$, K$^+$, and Cl$^-$ into the cell (Delpire and Gagnon, 2018). Mutations in NKCC2 are known to cause type I Bartter syndrome (Simon and Lifton, 1996) and a recent clinical report shows homozygous loss-of-function mutations in NKCC1 cause the novel disease Kilquist syndrome (Macnamara et al., 2019).

To validate the genetic interaction observed between NGLY1 and Ncc69 in the GWA, we generated ubiquitous double knockdown (DKD) Drosophila and scored offspring that survived to eclosion. NGLY1 KD caused a decrease in survival to ~25%, in accordance with our previous report of this RNAi line (Owings et al., 2018). Ncc69 KD did not cause any significant decrease in survival ($\chi^2=1.002$, $p=0.3168$), as previously reported (Leiserson et al., 2011). The DKD, however, caused complete lethality. This synthetic lethality confirms Ncc69 as a hit from the NGLY1 modifier screen.

KD of cation-chloride cotransporters in glia has been shown previously to cause seizures in Drosophila (Rusan et al., 2014), and we wanted to test whether this phenotype could be modified by NGLY1 KD. We performed single and double knockdowns of NGLY1 and Ncc69 in glial cells using the repo-GAL4 driver. This Ncc69 RNAi line is the same as the one described above. Drosophila

![Figure 3. NGLY1 and Ncc69 interact genetically in Drosophila. (A) Proportion of flies surviving to eclosion in ubiquitous knockdowns. NGLY1 knockdown (KD) are UAS-PnglRNAi/+; Tubulin-GAL4/+; Ncc69 KD are UAS-Ncc69RNAi/+; Tubulin-GAL4/+; NGLY1 Ncc69 double knockdown (DKD) are UAS-PnglRNAi/+; UAS-Ncc69RNAi/Tubulin-GAL4/+. Four separate matings were performed for each cross with at least 40 offspring generated for the balancer control for each. Fraction surviving is calculated compared to balancer offspring. Chi-square analysis was performed for the total number of flies compared to expected Mendelian numbers. NGLY1 KD $\chi^2=109.7$, $p<0.0001$; Ncc69 KD $\chi^2=1.002$, $p=0.3168$, and NGLY1 Ncc69 DKD $\chi^2=186$, $p<0.0001$. (B) Bang sensitivity assay to assess seizures in glial knockdown flies. WT flies are attP2 and attP40. NGLY1 KD are UAS-PnglRNAi/+; repo-GAL4/+; Ncc69 KD are UAS-Ncc69RNAi/+; repo-GAL4/+; NGLY1 Ncc69 DKD are UAS-PnglRNAi/+; UAS-Ncc69RNAi/repo-GAL4. For each genotype, at least 45 4- to 7-day-old females were used to calculate the percent seizing at a given time after vortexing. Repeated measures ANOVA p-value=0.000176.](https://doi.org/10.7554/eLife.57831)
were assessed for seizure phenotype using the bang sensitivity assay (Figure 3B). Control, wild-type flies show immediate recovery, as expected. In NGLY1 KD flies, ~30% showed severe seizures in the form of complete immobility 5 s following vortex. However, by 10 s following vortex, NGLY1 KD flies were completely recovered. This is the first report of seizure phenotype in any NGLY1 deficiency model, mimicking what is observed in patients. Ncc69 KD flies showed severe seizures with 75% seizing at 5 s following vortex, in line with previous reports (Rusan et al., 2014). In the DKD, there was a partial rescue of the severe Ncc69 phenotype. At all time-points between 5 and 60 s, the DKD flies showed an intermediate phenotype relative to NGLY1 and Ncc69 single KDs, confirming a genetic interaction between NGLY1 and Ncc69.

Functional analysis of NKCC1 in NGLY1 null MEFs

To understand the cell biology behind the genetic interaction that was observed in Drosophila, we utilized NGLY1 knockout (-/-) mouse embryonic fibroblasts (MEFs) (jax.org/strain/027060). Fibroblasts should only express the ubiquitous ortholog, NKCC1 (Haas and Forbush, 1998). When the membrane fraction of NGLY1 -/- MEFs was analyzed by immunoblot for NKCC1 there was a noticeable shift in the average molecular weight of the band compared to wildtype, control cells (+/+)(Figure 4A). Using the molecular weight marker to calculate the size of the proteins (un-cropped blot in Figure 4—figure supplement 1), the upper limits of the bands were ~170 kDa for both +/- and -/- cells (Figure 4B). The lower limit of the bands, however, were ~140 kDa for +/- cells and ~150 kDa for the -/- cells (Figure 4C).

To determine if a glycosylation event might be responsible for this size difference, cell lysates were treated with deglycosylating enzymes (Figure 4D). PNGase F removes all N-linked glycans and this treatment caused a large decrease in the molecular weight, to ~125 kDa in NKCC1 proteins from both NGLY1 +/- and -/- cells. The expected weight of mouse NKCC1 without any post-translational modifications is 130 kDa indicating that all post-translational modifications are likely N-linked.

Figure 4. Endogenous NKCC1 is altered in NGLY1-deficient MEFs. (A) Control (+/+ ) and NGLY1 null (-/-) MEFs were grown to confluency and then lysed to isolate the membrane and cytoplasmic fractions. Three separate membrane lysates for both genotypes were analyzed by immunoblotting for NKCC1 compared to a molecular weight marker (MWM). Blot was used for molecular weight calculations of the upper-most limit (see un-cropped blot in Figure 4—figure supplement 1) (B) And the lower-most limit (C) of the protein band. Red bar represents the mean. Two-tailed t-test was used to calculate p-values. (D) Membrane lysates from MEFs were treated with N-Glycosidase F (PNGase F), O-Glycosidase (O-Gly), or Endoglycosidase H (Endo H) for 1 hr then analyzed by immunoblot. Control (C)Samples were treated in all the same conditions but without the added enzyme. (E) MEFs were treated with 500 nM bortezomib (Bz) or equal volume of vehicle control (DMSO) for 4 hr then lysed to collect membrane and cytoplasmic fractions. Lysates were analyzed by immunoblotting for NKCC1. NRF1 was analyzed as a positive control of proteasome inhibition. Dark band at about 140 kDa in both (A), (D), and (E) is believed to be non-specific.

The online version of this article includes the following figure supplement(s) for figure 4:

Figure supplement 1. Full western blot of NKCC1 in NGLY1 +/- and -/- MEFs.
glycans. This is in accordance with the prediction of two canonical N-linked glycosylation sites (Payne et al., 1995). The fact that there is no difference in molecular weight between the +/+ and -/- after treatment indicates the difference observed in the untreated state was eliminated by the PNGase enzyme. Treatment with O-Glycosidase had no effect on the molecular weight of the band in either the +/+ or -/- lysates. Although O-Glycosidase does not cleave every type of O-linked glycan, these results coupled with the PNGase results indicate there are likely no O-linked glycans on NKCC1. Finally, to determine the maturation state of the N-linked glycans Endoglycosidase H was used. No change in molecular weight was seen, indicating the N-linked glycans are no longer in the high-mannose state in both the +/+ and -/- cells.

The most well-studied substrate of NGLY1, NRF1, is degraded by the proteasome under normal conditions, and only when the proteasome is stressed or inhibited does NRF1 become active. Indeed, NGLY1 is thought to act in the ERAD pathway and therefore all its substrates may be regulated in some way by proteasomal degradation. To test if NKCC1 abundance is affected by the proteasome, MEFs were treated with the proteasome inhibitor bortezomib (Bz). As expected, there was an increase in NRF1 abundance during proteasome inhibition (Figure 4E). However, there was no noticeable increase in NKCC1 protein for either the +/+ or -/- MEFs, indicating proteasomal degradation is not a major regulator of NKCC1.

Given the altered glycosylation state of NKCC1 observed in NGLY1 -/- MEFs, we wanted to determine the functionality of NKCC1 in these cells. Previous reports have shown that inhibiting N-linked glycosylation can decrease functionality for both NKCC1 (Singh et al., 2015) and for NKCC2 (Paredes et al., 2006), and therefore, we hypothesized that the misglycosylation might also decrease function. The NKCC proteins and the Na⁺/K⁺-ATPase can both transport Rb⁺ in place of K⁺, so we incubated cells with radioactive ⁸⁶Rb and measured cellular uptake. Cells were assayed in the presence or absence of 10 μM bumetanide, an NKCC inhibitor, or 100 μM ouabain, a Na⁺/K⁺-ATPase inhibitor, as compared to vehicle control. The ouabain-sensitive and bumetanide-sensitive activities were summed, they accounted for all of the ⁸⁶Rb flux observed (Figure 5—figure supplement 1). We found that bumetanide-sensitive ⁸⁶Rb flux, but not ouabain-sensitive flux, was impaired by ~50% in the -/- MEFs, indicating a specific defect in NKCC1 activity without impairment in the Na⁺/K⁺-ATPase (Figure 5). We assayed ion transport activity under three conditions of isotonic, hypertonic, or hypotonic baths. Although both hypertonic and hypotonic low chloride baths can stimulate NKCC1 activity in other cell types (Darman and Forbush, 2002), we did not see any

Figure 5. NGLY1 -/- MEFs show decreased NKCC1-specific ion flux. (A) Bumetanide-sensitive ⁸⁶Rb flux was measured in NGLY1 +/+ and NGLY1 -/- MEFs to measure NKCC1 activity. Flux was examined in three bath conditions, isotonic (iso), hypotonic (hypo), and hypertonic (hyper). There was a significant effect of genotype (p<0.0001) in two-way ANOVA, with no significant effect of condition (p=0.5756) or interaction (p=0.8075). Adjusted p-values for Sidak’s multiple comparisons test between NGLY1 +/+ and NGLY1 -/- are shown in the figure. (B) Ouabain-sensitive ⁸⁶Rb flux was measured in NGLY1 +/+ and NGLY1 -/- MEFs to measure Na⁺/K⁺-ATPase activity in the same three conditions as in A. There were no significant effects of genotype (p=0.0516), condition (p=0.3047) or interaction (p=0.4711) by two-way ANOVA, indicating the NGLY1 knockout has a specific effect on NKCC1 activity without affecting Na⁺/K⁺-ATPase activity. The online version of this article includes the following figure supplement(s) for figure 5:

Figure supplement 1. ⁸⁶Rb uptake in MEFs occurs through bumetanide-sensitive and ouabain-sensitive pathways.
effect of bathing medium in the MEFs, consistent with a recent report of lack of hypertonic stimulation of NKCC1 in human fibroblasts (Delpire et al., 2016). Together, these data demonstrate that loss of NGLY1 results in a change in the glycosylation state of NKCC1 and a significant reduction in NKCC1 function.

Discussion

Like many rare diseases, research into the pathogenesis of NGLY1 deficiency has been narrowly focused, based on early hypotheses. This often limits how we understand the connection between a particular disease and other pathways. For NGLY1 deficiency specifically, basic research and potential therapies have focused intensely on NRF1, the first well-established substrate of NGLY1 deglycosylation. Motivated by the extensive phenotypic variation among NGLY1 deficiency patients, we took advantage of natural genetic variation in Drosophila to identify modifiers of NGLY1 deficiency. This unique screen demonstrated that (1) we can model the extensive phenotypic variation observed and (2) that genetic variation can cause this phenotypic variability. Association analysis then identified a number of exciting candidate modifier genes. Here, we have validated the novel and conserved modifier NKCC1 (Drosophila Ncc69), a new potential therapeutic target for NGLY1 deficiency.

A major advantage of screens is the identification of previously unanticipated biological connections. First, our association analysis of the lethality screen has generated a list of 61 genes that we hope the scientific and patient communities will be able to use. Second, GSEA identified several pathways, including rRNA metabolism and nuclear transport that are surprising based on known NGLY1 functions. Third, ERC analysis identified genes that are coevolving with NGLY1 across the animal kingdom, including both rRNA/mRNA pathways and nuclear transport. It appears that rRNA metabolism and nuclear transport are likely important to NGLY1 function, yet it remains unclear how NGLY1 is connected to these pathways. Components of the ribosome and the nuclear pore are often O-glycosylated. While there is no direct connection between NGLY1 and O-GlcNAcylation, we have previously demonstrated that loss of NGLY1 impacts UDP-GlcNAc levels (Owings et al., 2018). It is highly plausible that a misregulation of UDP-GlcNAc levels could affect O-GlcNAcylated proteins. More work is needed to determine exactly how NGLY1 is connected to these unexpected pathways. Finally, the ERC analysis also identified 26/151 known Congenital Disorders of Glycosylation (CDG) genes. While NGLY1 is also classified as a CDG, it is unclear why there might be co-evolution with other CDG genes. Perhaps, there is a feedback mechanism, again, related to UDP-GlcNAc biosynthesis that connects these genes. These results suggest that there is a previously unknown connection between these loosely connected CDG genes.

When analyzing the list of modifier genes, it is apparent that many of the candidates are involved in ERAD. This offers a proof-of-principle that this screen is well suited for identifying bona fide biologically relevant modifiers. Several previous studies linked NGLY1 with the ERAD process (Bebök et al., 1998; Katiyar et al., 2005; Park et al., 2001). Yet, NGLY1 does not appear to be required for proper ERAD function (Hirsch et al., 2003; Misaghi et al., 2004). While perturbations to ERAD often result in ER stress, we have previously reported that there was no functional or transcriptome evidence for ER stress in a Drosophila model of NGLY1 deficiency (Owings et al., 2018). Others have reported no ER stress in NGLY1-/- human cells, mice, and rats (Asahina et al., 2020; Mueller et al., 2020; Tambe et al., 2019). However, there is conflicting evidence for ER stress as it was recently reported that ER stress markers were upregulated in NGLY-/- MEFs (Galeone et al., 2020). Nevertheless, in our current screen, we did not identify any genes involved in canonical ER stress responses, suggesting that ER stress might not play a large role in the pathogenesis of the disease. Here, we have reported that NKCC1 is altered in NGLY1-/- cells, however we found that inhibiting the proteasome had no effect on the protein abundance. Thus, it may be that NGLY1 functions to regulate proteins in various ways that are closely related to ERAD, but that do not result in direct proteasomal degradation, and thereby, do not cause an accumulation of misfolded protein and ER stress.

As with most hypothesis-free approaches, our analyses produced many exciting new avenues for exploration. In order to keep our work relevant and translatable to the clinic, we have focused our follow up experiments on genes with human orthologs. In this report we began with our top hit, Ncc69 (human NKCC1/2), which encodes an SLC12 Na+/K+/2Cl⁻ transporter. In cells derived from an
NGLY1 -/- mouse model, we found that NKCC1 protein migrated at a higher average molecular weight relative to +/+ cells. We found this altered glycosylation was accompanied by a ~50% reduction in NKCC1 activity.

NKCC1 contains two canonical N-linked glycosylation sites (Payne et al., 1995) similar to those validated in NKCC2 (Paredes et al., 2006). As expected, both sites are in an extracellular loop, and therefore, these sites face the lumen of the ER during protein translation and maturation. Because of this arrangement, these two sites are not predicted to be accessible by NGLY1, which is localized to the cytoplasm. However, recent work indicates that NGLY1 can deglycosylate proteins prior to their complete retrotranslocation out of the ER (Galeone et al., 2020). NGLY1 is recruited to the cytosolic surface of the ER where it deglycosylates BMP4, which is in fact necessary for its retrotranslocation to the cytoplasm. Perhaps NKCC1 is being deglycosylated on one of its two canonical N-linked glycosylation sites through a similar mechanism. Alternatively, it may be that NGLY1 is acting on a non-canonical, cytoplasmic N-linked glycosylation site. Sequence analysis reveals three other asparagine residues within the necessary N-X-S/T sequence for N-linked glycosylation, however, two are predicted to be in transmembrane domains; the third (human NKCC1 residue N168) is located in the amino-terminal cytoplasmic tail of the protein. The recent Cryo-EM structure determined that the amino-terminal tail is disordered and that the carboxy-terminal tail acts as a regulatory domain (Chew et al., 2019). Although rare, there have been reports of cytosolic N-linked glycosylation, including on the dog kidney Na+, K+-ATPase pump (reviewed in Hart et al., 2017). In all these cases, and in our case here, the mechanism for cytoplasmic glycosylation remains unknown. Determining if one of the two canonical sites is altered or if a non-canonical, cytoplasmic site is altered is a top priority for future work.

While we observe a glycosylation difference on NKCC1, and NGLY1 is a deglycosylating enzyme, we cannot eliminate the possibility that this NKCC1 regulation is a secondary effect. NGLY1 may be regulating an intermediary protein that in turn, regulates NKCC1. This was recently found to be the case for aquaporins in NGLY1-deficient cells (Tambe et al., 2019). NGLY1 was found to regulate the abundance of transcription factors Atf1/Creb1 independent of its enzymatic activity. Atf1/Creb1 then, in turn, regulates the transcription of several aquaporin subunits. Given that we show the abundance of NKCC1 does not change, but rather NKCC1 has a molecular weight shift, it is likely that the altered state is due to some difference in a post-translational modification. If this effect is secondary, it may be that NGLY1 is directly affecting Golgi-localized glycosyltransferases that in turn modify the already present glycans. Or that NGLY1 is altering the function of a sialyltransferase, thereby altering sialic acid residues on NKCC1 which are known to cause significant changes in migration on SDS-PAGE. This hypothesis is supported by the fact that Endo H treatment did not affect NKCC1 from either +/+ or -/- cells, indicating that NKCC1 has been fully processed through the Golgi. Future work to test this hypothesis could entail mass spectrometry analysis to determine the specific glycan structures on NKCC1.

Identification of targets and modifier genes should provide insight into the pathogenesis of a disease and help explain some of the patient phenotypes. When NRF1 was identified as the first target of NGLY1, it provided insight into some of the molecular defects observed in NGLY1-deficient cells, including deficits in proteasomal function and expression. These cellular phenotypes, however, did not translate well into insight into the complex patient symptoms. In contrast, decreased NKCC1 activity may explain some of the prominent features of NGLY1 deficiency. NKCC1 functions in many secretory epithelia, such as the salivary, sweat, and lacrimal glands, to allow basolateral ion uptake and subsequent secretion (Delpire and Gagnon, 2018). Therefore, a decrease in NKCC1 activity could well explain the alacrima and reduced saliva and sweat production seen in NGLY1 deficiency. Strikingly, a recent clinical report describes a patient with a homozygous deletion in NKCC1 (null) who has many overlapping features with NGLY1 deficiency patients, including absence of saliva, tears, and sweat (Kilquist syndrome) (Macnamara et al., 2019). The NKCC1 null mouse also displays defects in salivation (Evans et al., 2000). Other notable, but perhaps less specific, features in the NKCC1-deficient child, including developmental delay and gastrointestinal problems, also overlap with those observed in NGLY1 deficiency. The NKCC1-deficient patient also had severe hearing loss, cochlear defects, and abnormal auditory brainstem responses (ABRs). Several patients with missense mutations in NKCC1, rather than complete loss-of-function mutations, display bilateral sensorineural hearing loss (McNeill et al., 2020). NGLY1 deficiency patients do not have severe hearing loss, but do report abnormal ABRs. This difference may be explained by a 50% reduction in NKCC1 activity,
rather than complete loss of activity. The overlap in a majority of the symptoms between this new syndrome and NGLY1 deficiency strengthens the case for both a genetic and functional connection between NGLY1 and NKCC1.

NKCC1 may be a promising target for the development of NGLY1 deficiency therapies. Given that it is a transporter and partially exposed to the extracellular space, NKCC1 could be particularly amenable to modulation by small molecules. Based on our work reported here, we predict that increasing function of NKCC1 may ameliorate some symptoms. Quercetin, a flavonoid, is a readily available molecule that has been shown to enhance NKCC1 activity (Asano et al., 2009; Nakajima et al., 2011). These studies demonstrated that quercetin significantly increased 86Rb uptake in cell culture and that this increase was bumetanide-sensitive, indicating specificity to NKCC1. This is similar to other studies showing flavonoids increasing the activity of channels, such as the flavonoid genistein increasing activity of the cystic fibrosis transmembrane conductance regulator (CFTR) (Sugawara and Nikaido, 2014), and several flavonoids targeting cardiovascular channels (Scholz et al., 2010). These quercetin studies, however, are in the context of normal functioning NKCC1 protein. In our study, NKCC1 is altered from the WT state and it is unknown whether quercetin would be able to modulate the activity of an altered NKCC1. Strikingly, however, quercetin was recently discovered in a drug screen to provide benefit to NGLY1-deficient C. elegans (Iyer et al., 2019). These results coupled with our discovery of NKCC1 as a NGLY1 substrate offer an exciting new avenue of treatment for NGLY1 deficiency patients. Targeted studies are needed to determine if quercetin or other molecules could specifically enhance NKCC1 function in the context of NGLY1 deficiency.

In this study, we took a series of unbiased approaches in Drosophila to identify modifiers of NGLY1 deficiency. This resulted in a number of new insights into the potential pathogenesis of NGLY1 that we hope others will also investigate. With rare diseases like NGLY1 deficiency, unbiased and forward genetic approaches are an efficient method for expanding possible avenues of investigatory and therapeutic development. This study also highlights the power of using model organisms like Drosophila to uncover pathways and genes that can be validated in mammalian systems and targeted for therapeutic development.

### Materials and methods

#### Key resources table

| Reagent type (species) of resource | Designation | Source or reference | Identifiers | Additional information |
|-----------------------------------|-------------|---------------------|-------------|------------------------|
| Gene (Drosophila melanogaster)    | Pngl        | GenBank             | ID:35527 Dmel_CG7865 |                        |
| Gene (Drosophila melanogaster)    | Ncc69       | GenBank             | ID:39410 Dmel_CG4357 |                        |
| Gene (Mus musculus)              | NGLY1       | GenBank             | ID: 59007    |                        |
| Gene (Mus musculus)              | NKCC1       | GenBank             | ID: 20496 Slc12a2 |                        |
| Genetic reagent (Drosophila melanogaster) | Pngl-RNAi   | Bloomington Drosophila Stock Center | RRID:BDSC_54853 | y1 v1; P(y+t7.7 v+t1.8=TRiP HMUJ21590)attP40 |
| Genetic reagent (Drosophila melanogaster) | Tubulin-GAL4 | Bloomington Drosophila Stock Center | RRID:BDSC_5138 | y1 w*; P[w+mC=tubP-GAL4]LL7/TM3, Sb1 Ser1 |
| Genetic reagent (Drosophila melanogaster) | Tubulin-GAL80 | Bloomington Drosophila Stock Center | RRID:BDSC_5190 | y1 w*; P[w+mC]=tubP-GAL80]LL9 P(w[+mW.hs]=FRT[w[hs]])2A/TM3, Sb1 |

Continued on next page
**Drosophila lines**

Flies were maintained at 25°C on a 12 hr light/dark cycle and raised on a standard diet based on the Bloomington Stock Center standard medium with malt. All flies were aged 3–5 days old for experiments. For the DGRP screen, the following *D. melanogaster* stocks were used: *Pngl*<sup>RNAi</sup> (Bloomington *Drosophila* Stock Center: 54853) and *Tubulin*-GAL4 driver (5138). The *Tubulin*-GAL80 strain was provided by Dr. Carl Thummel (University of Utah). The DGRP strains are available at the Bloomington Stock Center. To measure circadian rhythm, the following stocks were used: a *w*-Berlin control strain, a *w; Pdf*-GAL4 strain (outcrossed to *w-* Berlin), and a *yv; UAS-Pngl*-RNAi strain (Bloomington stock center #42592). For bang sensitivity assays, the following stocks were used: *UAS-Pngl*-RNAi (BL #54853), *UAS-Ncc69*-RNAi (BL #28682), and *repo-GAL4*. These stocks were obtained from the Bloomington Stock Center and Adrian Rothenfluh (University of Utah) respectively.

**DGRP screen**

Virgin females from the DGRP strains were fed yeast overnight and then crossed with males from the donor strain UAS-*Pngl*<sup>RNAi</sup>/CyO,*Tubulin*-GAL80; *Tubulin*-GAL4/TM3,Sb in two replicate bottles.
Progeny were collected and scored for the four balancer classes: CyO, Sb, double balanced, or no balancers, with the no balancer flies being the NGLY1 KD. This cross should produce the expected 1:1:1:1 ratio of the four genotypes. Given that there is always a very low level of lethality associated with each balancer, the largest balancer class was considered the closest to the expected number. We scored at least 200 flies per DGRP cross. Males and females were combined for a single count. To calculate the proportion of NGLY1 KD flies by generating a ratio of NGLY1 knockdown/largest balancer class. This metric was used for the GWA.

**Genome wide association**

GWA was performed as previously described (Chow et al., 2016). DGRP genotypes were downloaded from the website, http://dgrp.gnets.ncsu.edu/. Variants were filtered for minor allele frequency (>0.05), and non-biallelic sites were removed. A total of 2,007,145 variants were included in the analysis. The proportion of NGLY1 KD flies surviving was regressed on each SNP. To account for cryptic relatedness (He et al., 2014; Huang et al., 2014), GEMMA (v. 0.94) (Zhou and Stephens, 2012) was used to both estimate a centered genetic relatedness matrix and perform association tests using the following linear mixed model (LMM):

\[
y = \alpha + x\beta + u + \epsilon
\]

\[
u \sim MVN_n(0, \lambda I_n + \Sigma(\text{K})^{-1})
\]

\[
\epsilon \sim MVN_n(0, \Sigma(\text{I}_n))
\]

where, as described and adapted from Zhou and Stephens, 2012, y is the n-vector of proportion lethality for the n lines, \( \alpha \) is the intercept, x is the n-vector of marker genotypes, \( \beta \) is the effect size of the marker. u is a n x n matrix of random effects with a multivariate normal distribution (MVN_n) that depends on \( \lambda \), the ratio between the two variance components, \( \Sigma(-1) \), the variance of residuals, and where the covariance matrix is informed by K, the calculated n x n marker-based relatedness matrix. K accounts for all pairwise non-random sharing of genetic material among lines. \( \epsilon \), is a n-vector of residual errors, with a multivariate normal distribution that depends on \( \Sigma(-1) \) and \( I_n \), the identity matrix. Genes were identified from SNP coordinates using the BDGP R54/dm3 genome build. An SNP was assigned to a gene if it was +/- 1 kb from a gene body.

**Gene set enrichment analysis**

GSEA was run to generate a rank-list of genes based on their enrichment for significantly associated polymorphisms as previously described (Palu et al., 2019). Polymorphisms within 1 kb of more than one gene were assigned to one gene based on a priority list of exon, UTR, intron, and upstream or downstream. Genes were assigned to GO categories, and calculation of enrichment score was performed as described (Subramanian et al., 2005). Only gene sets with \( \geq 5 \) genes, \( >0.25 \) enrichment score, and a \( p<0.05 \) were considered.

**Evolutionary rate covariation**

ERC is a method to examine the similarity of evolutionary histories of pairs of genes (Clark et al., 2012). The method examines the variation over time of a gene’s rate of sequence evolution. Using estimates of evolutionary rate over the branches of a gene’s phylogenetic tree, the method measures the correlation between genes of these branch-specific rates. Genes within correlated rate variation tend to be functionally related and have been used to discover new genes within pathways and diseases (Brunette et al., 2019; Friedlkeit et al., 2015; Raza et al., 2019).

ERC values in this study were taken from a compilation of ERC correlations calculated separately for three taxonomic groups: 62 mammals, 39 non-mammalian vertebrates, and 22 Drosophila species. Mammal and non-mammalian vertebrate alignments were taken from the multiz alignment available from the UCSC Genome Browser (Haeussler et al., 2019). For each alignment, we filtered out low-quality orthologs containing fewer than 50 non-gap amino acid sites or less than 70% non-gap sites and removed alignments with fewer than 15 species. Alignments were made for the Drosophila species after downloading protein-coding genome sequences from FlyBase and NCBI. Orthologous groups were identified using Orthofinder and alignments made with PRANK (Emms and Kelly, 2015; Löytynoja and Goldman, 2008). For each amino acid alignment, we estimated branch lengths using aaml in the phylogenetic analysis using maximum likelihood (PAML).
package (Yang, 2007). ERC values (correlation coefficients) for all genes with NGly1 were calculated using the RERconverge package (Kowalczyk et al., 2019). We report the ERC results for the mamalian group as the negative log of their p-values for each gene pair (Supplementary file 5 ‘nlogpvbest1’). Each gene pair also incorporated results from the vertebrate and Drosophila datasets by summing their negative log p-values, when orthologs were present for their respective datasets (Supplementary file 5 ‘sumnlogpvbest’). The resulting taxonomically integrated results of ERC with NGly1 were sorted and used for gene set enrichment analysis (GSEA).

Drosophila circadian rhythm assay
Male flies with the following genotypes were used in circadian rhythm assays: w/Y;Pdf-GAL4/+, yv/Y; UAS-Png1RNAi/+; and yv/Y; Pdf-GAL4/UAS-Png1RNAi/+ Two- to 5-day-old flies were entrained for at least 3 days to a 12 hr light: 12 hr dark regimen (LD) within a Drosophila Activity Monitor (DAM; Tri-Kinetics, Waltham, MA) filled with standard fly food. After entrainment, flies were monitored in complete darkness (DD) for 8 days. The data was collected in 30 min bins, and analyzed for period length using ClockLab, Version 6. Graphs were generated and one-way ANOVA performed, with Tukey’s multiple comparison of all three genotypes, using GraphPad Prism, Version 8.

Drosophila seizure assay
The Bang Sensitivity Assay (BSA) was performed on the following genotypes: UAS-Png1RNAi; repo-GAL4, UAS-Ncc69RNAi; repo-GAL4, and UAS-Png1RNAi+/; UAS-Ncc69RNAi/repo-GAL4. Females 4–7 days old were assayed. Flies were not exposed to CO2 for 3 days prior to BSA testing. Flies were flipped into empty vials and allowed to rest for 2 hr. They were then vortexed on a Thermo Scientific LP Vortex Mixer for 10 s at maximum speed. The video was used to score seizures at 5, 10, 30, and 60 s.

Mammalian cell culture and proteasome inhibition
MEFs were generated by Jackson Laboratory (Bar Harbor, Maine) from NGly1 knockout mice and littermate controls (C57BL/6J-Ngly1em4Lutzy/J, #027060). MEFs were immortalized in the laboratory of Dr. Hamed Jafar-Nejad (Baylor College of Medicine) and then gifted to us. MEFs were grown in DMEM (Gibco 11965) supplemented with 10% fetal bovine serum (FBS) and penicillin/streptomycin in 5% CO2 at 37˚C. For proteasome inhibition, MEFs were incubated with 500 nM bortezomib (EMD Millipore) or an equivalent volume of DMSO as a vehicle control, for 4 hr under standard conditions.

Western blotting
MEFs were grown to 80–90% confluency then collected. Cell pellets were weighed and then resuspended in a proportional volume of phosphate buffered saline (PBS). Equivalent volumes of resuspension were always used for each lysis. Cells were lysed using a cell fractionation kit (Cell Signaling Technologies, #9038) with each buffer supplemented with 1 mM PMSF and 1x protease inhibitor cocktail (Cell Signaling Technologies, #5871).

Lysates were separated by SDS-PAGE on 3–8% Tris-acetate gels (BioRad #3450129) for 2.5 hr at 150V, then transferred to PVDF membrane by wet transfer at 50V for 1 hr. Membranes were blocked in either 5% milk or 5% BSA according to the recommendations of the primary antibody manufacturer. Primary antibodies were as follows: anti-NKCC1 (Cell Signaling Technologies #14581), anti-TCF11/NRF1 (Cell Signaling Technologies #8052). Membranes were incubated in primary antibody at 1:1000 in blocking buffer overnight. IRDye secondary antibody (Abcam #216773) was used for infrared detection at 1:10,000 dilution in blocking buffer for 1 hr. Membranes were scanned on an Odyssey CLx (Li-cor) and analyzed with the accompanying software, Image Studio.

Deglycosylation reactions
MEFs were lysed in the same manner as described for western blotting. The membrane fraction was then incubated with one of the three deglycosylation enzymes: O-Glycosidase (New England Biolabs, #P0733), PNGase F (New England Biolabs, #P0704), Endoglycosidase F (New England Biolabs, #P0702) according to the manufacturer’s directions. Reactions were incubated at 37˚C for 1 hr. Controls were treated with all the same buffers and reaction conditions but without the added enzyme.
**Rb⁺ flux assay**

20,000 cells/well of immortalized MEFs from control or Ngly1 -/- mice were seeded into a 96-well plate. The following day, media was removed and the cells were washed with 1x PBS. Of pre-incubation medium (in mM, 135 Na gluconate, 5 K gluconate, 1 Ca gluconate, 1 Mg gluconate, 15 HEPES pH 7.4, 5 glucose), 100 μl was added to each well and the cells were incubated for 30 min at 37°C. Next, 100 μl of pre-incubation medium containing either DMSO, bumetanide, or ouabain was added to each well to achieve final concentrations of 0.1% (DMSO), 10 μM (bumetanide), or 0.1 mM (ouabain) and incubated for 30 min at room temperature. Then, 150 μL of medium containing DMSO (0.1%), bumetanide (10 μM) or ouabain (0.1 mM), as well as ⁸⁶Rb (10 mCi/μl), was added to each well. Three different media were used. Isotonic media contained (in mM): 140 NaCl, 5 KCl, 2 CaCl₂, 1 MgCl₂, 5 glucose, 15 HEPES pH 7.4. Hypertonic medium was the same as isotonic medium, with the addition of 75 mM sucrose. For hypotonic medium, isotonic medium was diluted 1:2 in water. The cells were incubated for 7 min at room temperature. Medium was removed and the cells were washed three times with ice-cold 1x PBS. Cells were lysed in 100 μl 2% SDS and incubated for 15 min at room temperature. Radioactivity was measured in a liquid scintillation counter.

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Additional files

Supplementary files
• Supplementary file 1. NGLY1 DGRP cross progeny counts. The number of eclosed flies were scored for each resulting genotype. The ‘no marker’ column represents flies expressing the NGLY1 RNAi. The largest balanced genotype was used as ‘expected’ for percent survival.
• Supplementary file 2. GWA analysis for survival in NGLY1 DGRP screen. Single-nucleotide polymorphisms (SNPs) are listed by chromosome position and rs ID.
• Supplementary file 3. Top associated SNPs. The top 125 variants. SNPs are listed in rank order of significance.
• Supplementary file 4. Gene set enrichment analysis (GSEA). Gene Ontology (GO) terms are listed by rank significance. Individual genes within each category are listed with the FBgn#.
• Supplementary file 5. Evolutionary rate covariance (ERC). Co-evolving genes are listed by rank significance (sumnlogpvbest). Genes that are known to cause a Congenital Disorder of Glycosylation (CDG) are highlighted in red.
• Transparent reporting form

Data availability
All data generated by this study are included in the manuscript and supporting files.

References
Ahlers LRH, Trammell CE, Carrell GF, Mackinnon S, Torrevillas BK, Chow CY, Luckhart S, Goodman AG. 2019. Insulin potentiates JAK/STAT signaling to broadly inhibit Flavivirus replication in insect vectors. Cell Reports 29: 1946–1960. DOI: https://doi.org/10.1016/j.celrep.2019.10.029, PMID: 31722209
Asahina M, Fujinawa R, Nakamura S, Yokoyama K, Tozawa R, Suzuki T. 2020. Ngly1-/- rats develop neurodegenerative phenotypes and pathological abnormalities in their peripheral and central nervous systems. Human Molecular Genetics 29:1635–1647. DOI: https://doi.org/10.1093/hmg/ddaa059, PMID: 32259258
Asano J, Niisato N, Nakajima K, Miyazaki H, Yasuda M, Iwasaki Y, Hama T, Dejima K, Hisa Y, Marunaka Y. 2009. Quercetin stimulates na+/K+/2Cl- cotransport via PTK-dependent mechanisms in human airway epithelium. American Journal of Respiratory and Critical Care Medicine 179:688–695. DOI: https://doi.org/10.1165/rcmb.2008-0338OC, PMID: 19251944
Beböök Z, Mazzochi C, King SA, Hong JS, Sorscher EJ. 1998. The mechanism underlying cystic fibrosis transmembrane conductance regulator transport from the endoplasmic reticulum to the proteasome includes
Sec61beta and a cytosolic, deglycosylated intermediary. *Journal of Biological Chemistry* **273**:29873–29878. DOI: https://doi.org/10.1074/jbc.P203.309317

Brunette GJ, Jamaluddin MA, Baldock RA, Clark NL, Bernstein KA. 2019. Evolution-based screening enables genome-wide prioritization and discovery of DNA repair genes. *PNAS* **116**:19593–19599. DOI: https://doi.org/10.1073/pnas.1906559116, PMID: 31501324

Chew TA, Orlando BJ, Zhang J, Latorraca NR, Wang A, Hollingsworth SA, Chen DH, Dror RO, Liao M, Feng L. 2019. Structure and mechanism of the cation-chloride cotransporter NKCC1. *Nature* **572**:488–492. DOI: https://doi.org/10.1038/s41586-019-1438-2, PMID: 31367042

Chow CY, Wolfrin MF, Clark AG. 2013a. Large neurological component to genetic differences underlying biased sperm use in *Drosophila*. *Genetics* **193**:177–185. DOI: https://doi.org/10.1534/genetics.112.146357, PMID: 23105014

Chow CY, Wolfrin MF, Clark AG. 2013b. Using natural variation in *Drosophila* to discover previously unknown endoplasmic reticulum stress genes. *PNAS* **110**:9013–9018. DOI: https://doi.org/10.1073/pnas.1307125110, PMID: 23667151

Chow CY, Kelsey KJ, Wolfrin MF, Clark AG. 2016. Candidate genetic modifiers of retinitis pigmentosa identified by exploiting natural variation in *Drosophila*. *Human Molecular Genetics* **25**:651–659. DOI: https://doi.org/10.1093/hmg/ddv302, PMID: 26662796

Clark NL, Alani E, Aquadro CF. 2012. Evolutionary rate covariation reveals shared functionality and coevolution of genes. *Genome Research* **22**:714–720. DOI: https://doi.org/10.1101/gr.132647.111, PMID: 22287101

Darwin RB, Forbush B. 2002. A regulatory locus of phosphorylation in the N terminus of the Na-K-Cl cotransporter, NKCC1. *Journal of Biological Chemistry* **277**:37542–37550. DOI: https://doi.org/10.1074/jbc.M202693200, PMID: 12145304

Delpire E, Wolfe L, Flores B, Koumangoye R, Schornak CC, Omer S, Pusey B, Lau C, Markello T, Adams DR. 2016. A patient with multisystem dysfunction carries a truncation mutation in human SLC12A2, the gene encoding the Na-K-2Cl cotransporter, NKCC1. *Molecular Case Studies* **2**:a001289. DOI: https://doi.org/10.1101/mcs.a001289, PMID: 27900370

Delpire E, Gagnon KB. 2018. Na’+K’-2Cl’ Cotransporter (NKCC) Physiological Function in Nonpolarized Cells and Transporting Epithelia. *Comprehensive Physiology* **8**:871–901. DOI: https://doi.org/10.1002/cphy.c170018, PMID: 29687903

Emms DM, Kelly S. 2015. OrthoFinder: solving fundamental biases in whole genome comparisons dramatically improves orthogroup inference accuracy. *Genome Biology* **16**:157. DOI: https://doi.org/10.1186/s13059-015-0721-2, PMID: 26243257

Enns GM, Shashi V, Bainbridge M, Gambello MJ, Zahir FR, Bast T, Crimian R, Schoch K, Platt J, Cox R, Bernstein JA, Scavina M, Walter RS, Bibb A, Jones M, Hegde M, Graham BH, Need AC, Ovieco A, Schaaf CP, et al. 2014. Mutations in NGLY1 cause an inherited disorder of the endoplasmic reticulum-associated degradation pathway. *Genetics in Medicine* **16**:751–758. DOI: https://doi.org/10.1038/gim.2014.22, PMID: 24651603

Evans RL, Park K, Turner RJ, Watson GE, Nguyen HV, Dennett MR, Hand AR, Flagella M, Shull GE, Melvin JE. 2000. Severe impairment of salivation in na+/K+/2Cl- cotransporter (NKCC1)-deficient mice. *Journal of Comparative Physiology B* **275**:26720–26726. DOI: https://doi.org/10.1007/s003600050730, PMID: 10831596

Everrett LJ, Huang W, Zhou S, Carbone MA, Lyman RF, Arya GH, Geisz MS, Ma J, Morgante F, St Armstrong G, Turlapati L, Anholt RRH, Mackay TFC. 2020. Gene expression networks in the *Drosophila* genome browser database: 2019 update. *Genome Research* **30**:836–844. DOI: https://doi.org/10.1101/gr.257592.119, PMID: 32782039

Galeone A, Adams JM, Matsuda S, Presa MF, Pandey A, Han SY, Tachida Y, Hirayama H, Vaccari T, Suzuki T, Lutz CM, Affolter M, Zuberi A, Jafar-Nejad H. 2020. Regulation of BMP4/Dpp retrotranslocation and signaling by deglycosylation. *eLife* **9**:e55596. DOI: https://doi.org/10.7554/eLife.55596, PMID: 32720893

Haas M, Forbush B. 1998. The Na-K-Cl cotransporters. *Journal of Bioenergetics and Biomembranes* **30**:161–172. DOI: https://doi.org/10.1023/a:1020521308985, PMID: 9672238

Haeussler M, Zweig AS, Tyner C, Speir ML, Rosenblum KR, Raney BJ, Lee CM, Lee BT, Hinrichs AS, Gonzalez JB, Gibson D, Diekhans M, Clawson H, Casper J, Barber GP, Haussler D, Kuhn RM, Kent WJ. 2019. The UCSC genome browser database: 2019 update. *Nucleic Acids Research* **47**:D853–D858. DOI: https://doi.org/10.1093/nar/gky1095, PMID: 30407534

Hart G, West C, Varki A, Cummings R, Esko J, Stanley P, Hart G, Aebl M, Seegerer P. 2017. Essentials of Glycobiology. Third Edition. Cold Spring Harbor Laboratory Press.

He BZ, Ludwig MZ, Dickerson DA, Barse L, Arun B, Vilhjalmsson BJ, Jiang P, Park SY, Tamirina NA, Selleck SB, Wittkopp PJ, Bell GI, Kreitman M. 2014. Effect of genetic variation in a *Drosophila* model of diabetes-associated misfolded human proinsulin. *Genetics* **196**:557–567. DOI: https://doi.org/10.1534/genetics.113.157800, PMID: 24281155

He P, Grotzke JE, Ng BG, Gunel M, Jafar-Nejad H, Cresswell P, Enns GM, Freeze HH. 2015. A congenital disorder of deglycosylation: biochemical characterization of N-glycanase 1 deficiency in patient fibroblasts. *Glycobiology* **25**:836–844. DOI: https://doi.org/10.1093/glycob/cwv042, PMID: 25909030

Hirsch C, Blom D, Ploegh HL. 2003. A role for N-glycanase in the cytosolic turnover of glycoproteins. *The EMBO Journal* **22**:1036–1046. DOI: https://doi.org/10.1093/emboj/cdg107, PMID: 12606569
Hosomi A, Fujita M, Tomioka A, Kaji H, Suzuki T. 2016. Identification of PNGase-dependent ERAD substrates in Saccharomyces cerevisiae. Biochemical Journal 473:3001–3012. DOI: https://doi.org/10.1042/BCJ20160453

Huang W, Massouras A, Inoue Y, Peiffer J, Rámírez M, Tarone AM, Turlapati L, Zichner T, Zhu D, Lyman RF, Magwire MM, Blankenburg K, Carbore MA, Chang K, Ellis LL, Fernandez S, Han Y, Highnam G, Hjelmén CE, Jack JR, et al. 2014. Natural variation in genome architecture among 205 Drosophila melanogaster genetic reference panel genomes. Genome Research 24:1193–1208. DOI: https://doi.org/10.1101/gr.171546.113, PMID: 24714809

Iyer S, Mast JD, Tsang H, Rodriguez TP, DiPrimio N, Prangley M, Sam FS, Parton Z, Perlstein EO. 2019. Drug screens of NGLY1 deficiency in worm and fly models reveal catecholamine, NRF2 and anti-inflammatory-pathway activation as potential clinical approaches. Disease Models & Mechanisms 12:mm040576. DOI: https://doi.org/10.1242/dmm.040576, PMID: 31615832

Kario E, Tirosh B, Ploegh HL, Navon A. 2008. N-linked glycosylation does not impair proteasomal degradation but affects class I major histocompatibility complex presentation. Journal of Biological Chemistry 283:244–254. DOI: https://doi.org/10.1074/jbc.M706237200, PMID: 17951257

Katiyar S, Yoshi S, Lennarz WJ. 2005. The retrotranslocation protein Derlin-1 binds peptide-n-glycanase to the endoplasmic reticulum. Molecular Biology of the Cell 16:4584–4594. DOI: https://doi.org/10.1091/mbc.e05-04-0345, PMID: 16055502

Kobayashi A, Tsukide T, Miyasaka T, Morita T, Mizoroki T, Saito Y, Ihara Y, Takashima A, Noguchi N, Fukamizu A, Hirotsu Y, Ohtsuji M, Katsuoka F, Yamamoto M. 2011. Central nervous system-specific deletion of transcription factor Nr1f1 causes progressive motor neuronal dysfunction. Genes to Cells 16:692–703. DOI: https://doi.org/10.1111/j.1365-2443.2011.01522.x, PMID: 21554501

Kondo H, Rabouille C, Newman R, Levine TP, Pappin D, Freemont P, Warren G. 1997. p47 is a cofactor for p97-mediated membrane fusion. Nature 388:75–78. DOI: https://doi.org/10.1038/40411, PMID: 9214505

Kowalczyk A, Meyer WK, Partha R, Mao W, Clark NL, Chikima M. 2019. RERconverge: an R package for associating evolutionary rates with convergent traits. Bioinformatics 35:4815–4817. DOI: https://doi.org/10.1093/bioinformatics/btz468, PMID: 31192356

Lam C, Ferreira C, Krasnewich D, Toro C, Latham L, Zein WM, Lehky T, Brewer C, Baker EH, Thurm A, Farmer CA, Rosenzweig SD, Lyons JJ, Schreiber JM, Gropman A, Lingala S, Ghany MG, Solomon B, Macnamara E, Davids M, et al. 2017. Prospective phenotyping of NGLY1-CDGD, the first congenital disorder of deglycosylation. Genetics in Medicine 19:160–168. DOI: https://doi.org/10.1038/gim.2016.75, PMID: 27388694

Lavey S, Chittoor-Vinod VG, Chow CY, Martin I. 2018. Genetic modifiers of neurodegeneration in a Drosophila Model of Parkinson’s Disease. Genetics 209:1345–1356. DOI: https://doi.org/10.1534/genetics.118.301119, PMID: 29907646

Lee CS, Lee C, Hu T, Nguyen JM, Zhang J, Martin MV, Vawter MP, Huang EJ, Chan JY. 2011. Loss of nuclear factor E2-related factor 1 in the brain leads to dysregulation of proteasome gene expression and neurodegeneration. PNAS 108:8408–8413. DOI: https://doi.org/10.1073/pnas.1019201108, PMID: 21536885

Lahrbach NJ, Breen PC, Ruvkun G. 2019. Protein sequence editing of SKN-1A/Nrf1 by peptide-n-glycanase controls proteasome gene expression. Cell 177:737–750. DOI: https://doi.org/10.1016/j.cell.2019.03.035, PMID: 31002798

Lahrbach NJ, Ruvkun G. 2016. Proteasome dysfunction triggers activation of SKN-1A/Nrf1 by the aspartic protease DDI-1. eLife 5:e17721. DOI: https://doi.org/10.7554/eLife.17721, PMID: 27528192

Leiserson WM, Forbush B, Keshishian H. 2011. Drosophila Glia use a conserved cotransporter mechanism to regulate extracellular volume. Glia 59:320–332. DOI: https://doi.org/10.1002/glia.21103, PMID: 21125654

Löytynoja A, Goldman N. 2008. Phylogeny-aware gap placement prevents errors in sequence alignment and evolutionary analysis. Science 320:1632–1635. DOI: https://doi.org/10.1126/science.1158395, PMID: 18566285

Mackay TF, Richards S, Stone EA, Barbadilla A, Ayroles JF, Zhu D, Casillas S, Han Y, Magwire MM, Cridland JM, Richardson MF, Anholt RR, Barrón M, Bess C, Blankenburg KP, Carbore MA, Castellano D, Chaboub L, Duncan L, Harris Z, et al. 2012. The Drosophila melanogaster genetic reference panel. Nature 482:173–178. DOI: https://doi.org/10.1038/nature10811, PMID: 22318601

Macnamara EF, Koehler AE, D’Souza P, Estwick T, Lee P, Vazina G, Fauni H, Braddock SR, Torti E, Holt JM, Sharma P, Malicdan MCV, Tiff CJ, Undiagnosed Diseases Network. 2019. Kif15/ku70 deficiency causes host-mediated membrane fusion. Nature 570:3001–3012. DOI: https://doi.org/10.1038/s41586-019-1398-7, PMID: 31002798

Maynard JC, Fujihira H, Dolgonos GE, Suzuki T, Burlingame AL. 2020. Cytosolic N-GlcNAc proteins are formed by the action of endo-β-N-acetylglucosaminidase. Biochemical and Biophysical Research Communications 530:719–724. DOI: https://doi.org/10.1016/j.bbrc.2020.06.127, PMID: 32782141

McNeill H, Knebel A, Arthur JS, Cuenda A, Cohen P. 2004. A novel UBA and UBX domain protein that binds polyubiquitin and VCP and is a substrate for SAPKs. Biochemical Journal 384:391–400. DOI: https://doi.org/10.1042/BJ20041498, PMID: 15362974

McNeill A, Iovino E, Mansard L, Vache C, Baud D, Bedoukian E, Cox H, Dean J, Goudie D, Kumar A, Newbury-Ecob R, Falleni C, Renieri A, Lopergolo D, Mari F, Blanchet C, Willems M, Roux AF, Pippucci T, Delpeire E. 2020. SLC12A2 variants cause a neurodevelopmental disorder or cochleovestibular defect. Brain 143:2380–2387. DOI: https://doi.org/10.1093/brain/awaa176, PMID: 32659971

Misaghi S, Pacold ME, Bloem D, Ploegh HL, Kebrel GA. 2004. Using a small molecule inhibitor of peptide-n-glycanase to probe its role in glycoprotein turnover. Chemistry & Biology 11:1677–1687. DOI: https://doi.org/10.1016/j.chembiol.2004.11.010, PMID: 15610852
PDF neurons each cause severe abnormalities of behavioral circadian rhythms in
for gene prioritization and construction of informative gene-based networks.
Radhakrishnan SK, Bertozzi CR. 2017. Inhibition of NGLY1 inactivates the transcription factor Nrf1 and
charge in cl- transport.
renal Na+-K+-2Cl- cotransporter is reduced by mutagenesis of N-glycosylation sites: role for protein surface
transcriptional and translational regulation in K562 cell lines.
and implicate this hydrolase in the proteasome-dependent pathway for protein degradation. PNAS 98:11163–
implicate this hydrolase in the proteasome-dependent pathway for protein degradation. PNAS 98:11163–
activity of the renal Na+-K+-2Cl- cotransporter is reduced by mutagenesis of N-glycosylation sites: role for protein surface
charge in cl- transport. American Journal of Physiology. Renal Physiology 290:F1094–F1102. DOI: https://doi.
goal for gene prioritization and construction of informative gene-based networks. PLOS Genetics 11:e1004967.
DOI: https://doi.org/10.1037/aji.2017.1055–1066. DOI: https://doi.org/10.1093/hmg/ddy026, PMID: 29346549
Palu RAS, Ong E, Stevens K, Chung S, Owings KG, Goodman AG, Chow CY. 2019. Natural genetic variation
section in Drosophila Identifies Wnt Signaling, Mitochondrial Metabolism, and Redox Homeostasis Genes as
Modifiers of Apoptosis. G3: Genes, Genomes, Genetics 9:3995–4005. DOI: https://doi.org/10.1534/g3.119.
400722, PMID: 31570502
Paredes A, Plata C, Rivera M, Moreno E, Vázquez N, Muñoz-Clares R, Hebert SC, Gamba G. 2006. Activity of the renal Na+-K+-2Cl-
cotransporter is reduced by mutagenesis of N-glycosylation sites: role for protein surface
change in cl- transport. American Journal of Physiology. Renal Physiology 290:F1094–F1102. DOI: https://doi.
Identification of proteins that interact with mammalian peptide:n-glycanase and implicate this hydrolase in the proteasome-dependent pathway for protein degradation. PNAS 98:11163–
Identification of proteins that interact with mammalian peptide:n-glycanase and implicate this hydrolase in the proteasome-dependent pathway for protein degradation. PNAS 98:11163–
PF, Haase B, Benes V, Paulsen M, Sehr P, Lewis J, Drewes G, Steinmetz LM, Drewes G, Steinmetz LM. 2020. Loss of N-Glycanase 1 alters
compared with those of the AcrAB-TolC system of
enzymatic activity.
radicchio as a molecular target of flavonoids. Cardiovascular Therapeutics 28:e46–e52. DOI: https://doi.org/10.1016/j.
Cardiovascular ion channels as a molecular target of flavonoids. Cardiovascular Therapeutics 28:e46–e52. DOI: https://doi.org/10.1016/j.
58:7250–7257. DOI: https://doi.org/10.1371/journal.pone.0101117, PMID: 24971529
Scholz EP, Zitron E, Katus HA, Karle CA. 2010. Cardiovascular ion channels as a molecular target of flavonoids. Cardiovascular Therapeutics 28:e46–e52. DOI: https://doi.org/10.1016/j.
DOI: https://doi.org/10.1371/journal.pgen.1007720, PMID: 30763317
DOI: https://doi.org/10.1037/aji.2017.1055–1066. DOI: https://doi.org/10.1093/hmg/ddy026, PMID: 29346549
Paredes A, Plata C, Rivera M, Moreno E, Vázquez N, Muñoz-Clares R, Hebert SC, Gamba G. 2006. Activity of the renal Na+-K+-2Cl-
cotransporter is reduced by mutagenesis of N-glycosylation sites: role for protein surface
charge in cl- transport. American Journal of Physiology. Renal Physiology 290:F1094–F1102. DOI: https://doi.
Identification of proteins that interact with mammalian peptide:n-glycanase and implicate this hydrolase in the proteasome-dependent pathway for protein degradation. PNAS 98:11163–
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PDF neurons each cause severe abnormalities of behavioral circadian rhythms in
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activity of the renal Na+-K+-2Cl- cotransporter is reduced by mutagenesis of N-glycosylation sites: role for protein surface
charge in cl- transport. American Journal of Physiology. Renal Physiology 290:F1094–F1102. DOI: https://doi.
Identification of proteins that interact with mammalian peptide:n-glycanase and implicate this hydrolase in the proteasome-dependent pathway for protein degradation. PNAS 98:11163–
Identification of proteins that interact with mammalian peptide:n-glycanase and implicate this hydrolase in the proteasome-dependent pathway for protein degradation. PNAS 98:11163–
activity of the renal Na+-K+-2Cl- cotransporter is reduced by mutagenesis of N-glycosylation sites: role for protein surface
charge in cl- transport. American Journal of Physiology. Renal Physiology 290:F1094–F1102. DOI: https://doi.
potentiates proteasome inhibitor cytotoxicity. ACS Central Science 3:1143–1155. DOI: https://doi.org/10.1021/acscentsci.7b00224, PMID: 29202016

Wolfe NW, Clark NL. 2015. ERC analysis: web-based inference of gene function via evolutionary rate covariation. Bioinformatics 31:3835–3837. DOI: https://doi.org/10.1093/bioinformatics/btv454, PMID: 26243019

Yang Z. 2007. PAML 4: phylogenetic analysis by maximum likelihood. Molecular Biology and Evolution 24:1586–1591. DOI: https://doi.org/10.1093/molbev/msm088, PMID: 17483113

Zhou X, Stephens M. 2012. Genome-wide efficient mixed-model analysis for association studies. Nature Genetics 44:821–824. DOI: https://doi.org/10.1038/ng.2310, PMID: 22706312

Zhu B, Jiang L, Huang T, Zhao Y, Liu T, Zhong Y, Li X, Campos A, Pomeroy K, Masliah E, Zhang D, Xu H. 2017. ER-associated degradation regulates alzheimer’s amyloid pathology and memory function by modulating γ-secretase activity. Nature Communications 8:1472. DOI: https://doi.org/10.1038/s41467-017-0199-4, PMID: 29133892

Zolekar A, Lin VJT, Mishra NM, Ho YY, Hayatshahi HS, Parab A, Sampat R, Liao X, Hoffmann P, Liu J, Emmitte KA, Wang YC. 2018. Stress and interferon signalling-mediated apoptosis contributes to pleiotropic anticancer responses induced by targeting NGLY1. British Journal of Cancer 119:1538–1551. DOI: https://doi.org/10.1038/s41416-018-0265-9, PMID: 30385822