O-GlcNAc transferase missense mutations linked to X-linked intellectual disability deregulate genes involved in cell fate determination and signaling

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Running title: Characterization of XLID associated mutations in OGT-TPRs

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Keywords: post-translational modification (PTM), glycosyltransferase, O-GlcNAc transferase (OGT), O-GlcNAc, genetic disease, X-linked intellectual disability (XLID), enzyme kinetics, thermodynamic stability, transcriptomics, RNA-seq

Abstract

It is estimated that ~1% of the world’s population has intellectual disability, with males affected more often than females. OGT is an X-linked gene encoding for the enzyme O-GlcNAc transferase (OGT), which carries out the reversible addition of N-Acetylglucosamine (GlcNAc) to Ser/Thr residues of its intracellular substrates. Three missense mutations in the tetratricopeptide (TPR) repeats of OGT have recently been reported to cause X-linked Intellectual Disability (XLID). Here we report the discovery of two additional novel missense mutations (c. 775 G>A, p. A259T and c. 1016 A>G, p. E339G) in the TPR domain of OGT that segregate with XLID in affected families. Characterization of all five of these XLID missense variants of OGT demonstrates modest declines in thermodynamic stability and/or activities of the variants. We engineered each of the mutations into a male human embryonic stem cell line using CRISPR/Cas9. Investigation of the global O-GlcNAc profile as well as OGT and OGA levels by Western blotting showed no gross changes in steady-state levels in the engineered lines. However, analyses of the differential transcriptomes of the OGT variant expressing stem cells revealed shared deregulation of genes involved in cell fate determination and LXR/RXR signaling, which has been implicated in neuronal development. Thus, here we reveal 2 additional mutations encoding residues in the TPR regions of OGT that appear causal for XLID and provide evidence that the relatively stable and active TPR variants may share a common, unelucidated, mechanism of altering gene expression profiles in human embryonic stem cells.

Introduction

Intellectual disability (ID) is defined in the Diagnostic and Statistical Manual – 5 (DSM-5) as a childhood-onset neurodevelopmental disorder characterized by limitations in intellectual
functioning and adaptive behavior. The estimated world-wide prevalence of ID is ~ 1% (1). While Down Syndrome and Fragile X Syndrome are the two most common causes of ID, it is also present as a feature in a number of rare conditions (2). Only a decade ago, ID in up to 80% of affected individuals in cohorts analyzed was of unknown etiology, including non-genetic factors (2). The genetic cause of ID is now established in slightly over 70% of affected individuals, an advance made possible by the use of massive parallel sequencing approaches (2) Monogenic defects account for 40% of the identified causes as opposed to those caused by copy-number variants (deletions of duplications) or aneuploidies. 5–10% of all monogenic defects are X-linked disorders (2).

OGT is a gene on the X chromosome encoding the enzyme O-GlcNAc transferase (OGT) (3), which is responsible for the co- and post-translational addition of β-D-N-acetylglucosamine (GlcNAc) to specific serine and threonine residues on a number of nucleocytoplasmic and mitochondrial proteins (4,5). O-GlcNAc addition is reversed by the action of the enzyme O-GlcNAc hydrolase (OGA) (4). The cycling of O-GlcNAc by these two enzymes on hundreds of proteins in the cell regulates basic cellular processes such as transcription, translation, and proteostasis, and influences cell survival, division, signaling, metabolism, and autophagy (4,6–9). Both OGT and OGA are essential genes in mice (3,10,11). OGT is a multi-domain protein with the catalytic center located within its C-terminus (12). At its N-terminus are multiple tetratricopeptide repeats (TPRs) that mediate protein-protein interactions (7,13–16). OGT appears to be important for the proper functioning of the brain; it is abundantly expressed in the brain (17) and O-GlcNAc is enriched in both the brain and in synapses (18,19). OGT is also a regulator of embryonic development and patterning (20–22), a discovery first made using the model organism Drosophila melanogaster (23), in which loss of OGT, also known as super-sex combs (sxc), results in abnormal morphology and death at the pharate adult stage (24,25).

While deregulation of O-GlcNAc associated with diseases such as cancer, Alzheimer’s disease, diabetes mellitus and cardiovascular disease has been well established (4), it is only recently, as the result of an X-exome parallel sequencing effort, that a missense mutation in OGT was first directly attributed to causing intellectual disability, accompanied by dysmorphic features and congenital abnormalities (26,27). Reports have since emerged of at least three other mutations in OGT that segregate with disease in other families with intellectual disability (28,29). In this study, we report the clinical features/phenotypes of three separate patients, two of who are brothers, with what appears to be syndromic intellectual disability. Using exome sequencing to determine the etiology of the disease, we discover two separate novel missense mutations in OGT that segregate with X-linked ID (XLID) in these patients. We undertake the in vitro biochemical characterization of these and other currently known XLID missense variants of OGT using recombinant proteins, revealing that the stability and activity of these OGT variants is only modestly reduced. To use as an ex vivo human isogenic model to study the effects of the XLID mutations on early development and disease pathogenesis, we engineered a human male embryonic stem cell line using CRISPR/Cas9 to express these OGT variants. We investigate, and find no significant changes in the steady-state global O-GlcNAc profile as well as OGT and OGA levels in these cells by Western blotting. Analysis of the transcriptome of these cells, however, reveals shared dysregulation of genes and pathways involved in early development, providing a starting point for dissecting the impact of OGT variants on common phenotypes in the XLID affected individuals.

Results

Case reports of three patients with intellectual disability

At the time of diagnosis, Patient 1 was a 7-year-old boy with moderate ID, microcephaly, hypothyroidism, abnormal sleep pattern, nystagmus and epilepsy. No other family member was similarly affected (Fig. 1a). During pregnancy, there was concern for intrauterine growth retardation. Patient 1 was born at 39 weeks’ gestation with APGAR scores of 8 at 1 minute and 9 at 5 minutes, weighing 6 lbs. 8 oz. (>10th centile). Postnatal course was significant...
Characterization of XLID associated mutations in OGT-TPRs for failure to thrive, with short stature and delayed milestones. The height and weight since early infancy were always below 3rd centile. Patient 1 had microcephaly since birth with head size consistently below 2nd centile. He walked at 3 years of age and was nonverbal until 5 years of age. At 7 years of age, it was estimated that the patient functioned at the level of an 18 to 24 month old. The patient was diagnosed with hypothyroidism at infancy and continued to receive thyroid hormones at diagnosis. He had mild conductive hearing loss despite use of myringotomy tubes. At the age of 4 years, the patient began to experience episodes of full body stiffening, cyanosis and unresponsiveness. His seizures included atonic seizures, tonic seizures, generalized tonic clonic seizures and staring spells. His seizures were treated with Oxcarbazepine, lamotrigine and clonazepam and were well-controlled at the time of diagnosis. His EEG at onset of seizures showed multifocal spikes. The patient also had insomnia and his EEG revealed disorganized background during sleep with no sleep spindles and vertex waves during sleep. A brain MRI at 8 months of age demonstrated thin corpus callosum. At age 7 years, his brain MRI revealed extremely thin corpus callosum particularly the posterior body and splenium, mild ventriculomegaly and abnormal periventricular T2 and FLAIR signal (Fig. 1b). In addition, the clivus appeared short and the posterior arch of first cervical vertebra was relatively hypoplastic (Fig. 1c). MRI of spine at 7 years of age revealed diminished vertebral body heights at multiple levels, especially the thoracic and lumbar levels. Mild disc bulge was also seen at multiple levels in the lumbar region. Patient 1’s karyotype, comparative genomic hybridization and DNA test for Fragile X did not reveal any abnormalities. Several random glucose levels were mildly elevated. Routine metabolic testing including serum lactate and ammonia, serum amino acids, urine organic acids, serum acylcarnitine profile, plasma very long chain fatty acids, lysosomal enzyme assays and transferrin testing for congenital disorders of glycosylation were normal. Patient 2 and Patient 3 are siblings and unrelated to Patient 1 (Fig. 1d). There was a maternal family history of intellectual impairment with the mother’s two male siblings and a maternal great uncle being affected (Fig. 1d). The parents’ first child is healthy with no history of developmental or learning difficulties (Fig. 1d). Patient 2 was born at 39 weeks after an uncomplicated pregnancy. His birth weight was 2820 g (10th centile). In early childhood his motor development was within normal limits but he was noted to have language delay. He was later diagnosed with mild to moderate ID. He also had behavioral difficulties. He experienced recurrent middle ear infections and required multiple tympanostomy tube insertion. Examination findings in Patient 2 at age 8 years 4 months were weight 22.4 kg (10-25th centile), height 127.4 cm (50th centile and head circumference 49.5 cm (2nd centile). He had a dolicocephalic face shape with a high arched palate and malalignment of the teeth (with absence of two upper lateral incisors. His ears were long (length > 97th centile). On neurological examination he had generalised mild hypotonia and subtle asymmetry of deep tendon reflexes. Patient 3 (Fig. 1d) was born after an uncomplicated pregnancy. He had a dolicocephalic face shape with broad nasal root and downslanted palpebral fissures. His philtrum was long and smooth and upper lip was thin. His palate was high arched. He had bilateral fifth finger clinodactyly. G-banded and molecular karyotypes and Fragile X testing were normal in both Patient 2 and Patient 3.

Identification of two distinct missense mutations in the TPR region of OGT causal for XLID To investigate the etiology of ID in Patient 1, full exome sequencing was performed using genomic DNA isolated from the patient, his parents and a healthy brother. Initially, 20 unique alterations in 20 genes were identified. Manual review to rule out sequencing artifacts and polymorphisms as
well as genes lacking clinical overlap with patient’s clinical course resulted in a short list of 4 genes with 4 unique alterations. Further analysis considering likely autosomal and X-linked inheritance models and familial segregation revealed a hemizygous threonine to alanine (p.A259T) variant in exon 7 of the OGT gene as the causal mutation (Fig 2a-c). Based on data from the Exome Aggregation Consortium (ExAC) (30), the OGT c.775G>A alteration was not observed among 60,706 individuals tested. Allele frequency data for this nucleotide position are not currently available from the 1000 Genomes Project (31) and the alteration is not currently listed in the Database of Single Nucleotide Polymorphisms (dbSNP) (32). The p. A259 amino acid is invariable across available vertebrate species. The p.A259T alteration is predicted to be probably damaging by Polyphen (33) (score 1.00) and deleterious by SIFT (34) (score 0.01) in silico analyses. Co-segregation analysis of the c.775G>A (p.A259T) alteration revealed that it is present in the patient’s mother and absent in his father and brother (Fig. 1a).

To find the cause of ID in Patients 2 and 3, the genomic DNA of Patient 2 was subjected to X-chromosome exome sequencing, following which confirmation of variants and segregation analysis in further family members, including Patient 3, was performed by Sanger sequencing (see Methods). Resultant output variants (8 unique alterations in 7 genes were found, with one of these genes harboring a single silent mutation in addition to a missense mutation) were then assessed based on presence in known SNP databases including the Genome Aggregation Database (gnomAD) (30), predicted functional impact, nucleotide and amino acid conservation, and biochemical nature of amino acid substitutions. Only one variant remained after the filtering process; a hemizygous mutation c.1016A>G (p.E339G) located in exon 8 of the OGT gene (Fig 2a-c). This variant is not present in ExAC or gnomAD, nor listed in dbSNP (32). The amino acid p.E339 is invariable across available vertebrate species, and is predicted to be probably damaging by Polyphen (33) (score 1.00) and deleterious by SIFT (34) (score 0). Confirmation of OGT c.1016A>G and segregation analysis in further family members by sequencing confirmed that this variant was present in the proband (Patient 2), as well as the patient’s mother and affected brother (Patient 3) (Fig. 1d). It was absent in his father and unaffected brother (Fig. 1d).

The identification of these two novel mutations in OGT now result in a total of five known missense mutations in the TPR domain of OGT causal for XLID (Fig. 2a-b) (26-29). Patient phenotypes resulting from each of these mutations are summarized in Table 1. Modeling of the XLID-TPR mutations within the structure of OGT (Fig. 2c) shows that substitutions at these residues would likely result in steric clashes with side chains of neighboring residues (as in the case of L254F, A259T, R284P, A319T) and the loss of secondary structure elements (in the case of R284P) or of stabilizing interactions (in the case of E339G), which could lead to changes in the structure and thermodynamic stability of OGT.

OGT XLID-TPR mutants form functional dimers but demonstrate decreased thermal stability

Having identified the mutations causal for ID in our patients, we wanted to biochemically characterize these and other known XLID OGT variants. It is known that OGT forms a functional dimer. Hydrophobic contact between the TPR residues Trp208 in one monomer and Ile211 in the second monomer causes OGT to dimerize (14). Given that the XLID mutations lie in the TPR domain of OGT, we wanted to investigate the impact of these amino acid substitutions on OGT dimerization. To do this, we used purified recombinant variant TPR domains alone. Samples of the proteins were run on a Superdex 200 gel-filtration column and all of the mutants eluted at the same elution volume as wild-type (WT) TPR domain, indicating that the mutations do not affect the dimerization of OGT (Fig. 3a). Given that the modeling of the mutations into the structure of OGT predicted changes in the structure/stability of the protein, we proceeded to analyze the effect of the mutations on the thermodynamic stability of the TPR domain using Differential Scanning Fluorimetry (Thermofluor assay). Samples of the proteins combined with the Sypro Orange dye were subjected to increasing temperature as a function of time in a real-time PCR machine, which measures the increase in fluorescence of the dye as it binds to hydrophobic regions of the proteins.
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that become exposed as they denature at higher temperatures. This assay revealed that all of the XLID OGT mutant TPR domains are less stable than OGT-WT, with OGT-R284P and OGT-L254F being most impacted, i.e. least stable. This is in agreement with previously published data for OGT-R284P (29) (Fig. 3b). The high degree of purity of the protein preparations used for this assay are demonstrated by Comassie G-250-staining following SDS-PAGE separation (Supplementary Fig. 1a).

**XLID-TPR variants of OGT are active enzymes but exhibit impaired glycosyltransferase kinetics**

Having established that the XLID OGT-TPR mutants form functional dimers like OGT but that their thermal stability is modestly reduced, we went on to assess the function of all of the enzyme variants, except for L254F, which we previously characterized (27), using recombinant full length OGT variants. We performed an end-point assay wherein we incubated wild-type or mutant OGTs with the well-studied acceptor protein substrate CK2α (12) in the presence or absence of the donor substrate UDP-GlcNAc and then probed the samples for O-GlcNAc by Western blotting using the anti-O-GlcNAc antibody RL2 (Fig. 4a). We found that all of the XLID OGT-TPR variants glycosylated CK2α to an extent comparable to OGT-WT (Fig. 4a).

OGT is not only a glycosyltransferase, but also carries out the proteolytic processing of the transcriptional cofactor HCF1 (35). Interestingly, mutations in HCFC1 have been implicated in causing XLID (36-38). We therefore evaluated the protease activity of OGT using the short GST-tagged substrate HCF1-rep1 (35,39,40) in an endpoint assay similar to the one for CK2α O-GlcNAcylation (Fig. 4b). This assay revealed that all of the XLID OGT-TPR mutants are active HCF-1 proteases, with activity comparable to OGT-WT, with the exception of OGT-R284P, which showed visibly reduced cleavage of the substrate compared to OGT-WT, although it still glycosylated the cleavage product efficiently (Fig. 4b). This is consistent with previously published data for OGT-R284P (29).

Having observed that the XLID OGT-TPR variants are competent enzymes, at least with respect to the conditions used in the end-point assays, we asked if the enzyme kinetics of OGT would be affected by these mutations to explain the disease association. This time we titrated in increasing concentrations of the protein substrate CK2α into reaction mixtures containing a fixed concentration of UDP-GlcNAc in excess of the $K_m$, and OGT-WT or one of the XLID variants and measured the Michaelis-Menten kinetics using a luminescence-based assay for the detection of UDP released (Fig. 4c). All of the XLID OGT-TPR variants showed a modest increase in $K_m$ accompanied by a reduction in turnover, resulting in a ~ 2-fold decrease in catalytic efficiency (Fig. 4c, Table 2). The purity of the protein preparations used for these assays is demonstrated by the Comassie G-250-stained SDS-PAGE gel (Supplementary Fig. 1b).

**Human embryonic stem cells expressing the XLID OGT-TPR variants show no gross changes in basal O-GlcNAc, OGT and OGA levels**

After characterizing the properties of the XLID OGT-TPR variants using recombinantly isolated proteins in *in vitro* assays, we wanted to dissect the effects of the mutations in the context of a human isogenic *ex vivo* model in an attempt to identify and understand the molecular basis of the disease. To this end, we engineered the mutations into the RUES-1 male human embryonic stem cell line using CRISPR/Cas9 (Supplementary Fig. 1). We chose stem cells so that we could investigate both differentiation pathways as well as multiple cell types. We were able to generate cell lines containing all of the mutations except for OGT-A319T, which presented as a technical challenge, likely because the gRNA design with the highest predicted efficiency did not straddle the site of the intended mutation and no alternative gRNAs (with or without high off-target efficiency) were designable. All of the cell lines obtained were confirmed to contain the intended mutation by Sanger sequencing (Supplementary Fig. 2) and no noticeable changes were observed in cell growth or maintenance of pluripotency (data not shown).

We previously showed that OGT steady-state levels in XLID-derived lymphoblasts containing the OGT-L254F mutation were lower than in unaffected controls from relatives, however, global O-GlcNAc levels assessed by Western blotting, remained unchanged (27). This was demonstrated to be the consequence of a
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compensation mechanism whereby OGT, in complex with the mSin3A-HDAC1 complex, transcriptionally downregulated OGA to maintain O-GlcNAc homeostasis (27). A similar mechanism was proposed to exist in patient-derived fibroblasts expressing OGT-R284P, however, as the authors noted, significant differences in the levels of OGT and OGA in the two separate control fibroblast lines expressing OGT-WT used in that study confounded the findings (29). In light of this, we set out to analyze and compare the levels of O-GlcNAc, OGT and OGA in our RUES-1 cells expressing the different OGT variants by Western blotting using two antibodies against each of the targets (Fig. 5a-d). It was unsurprising that we found no significant changes (CTD 110.6: p > 0.4, RL2: p > 0.7; Mann-Whitney U test, data not shown) in the levels of O-GlcNAc in the XLID OGT-TPR mutant-expressing lines compared to wild-type (Fig. 5a-b). We also, however, saw no significant changes in the levels of OGT (DM17: p > 0.2, F-12: p > 0.1, Mann-Whitney U test, data not shown) or OGA (G-12: p > 0.7, 14711-1-AP: p > 0.4, Mann-Whitney U test, data not shown) (Fig. 5c-d), an observation suggesting that there may be cell-type specific differences in the regulation, and indeed function, of these enzymes.

Differential transcriptomes of the hES cells expressing XLID OGT-TPR variants reveal changes in genes associated with neural development

Given the developmental phenotypes of the patients and the fact that OGT, belonging to the polycomb group of developmental genes (24), regulates the transcription of a number of these genes (22), we went on to investigate changes in gene expression between OGT-WT and XLID OGT-TPR mutant-expressing stem cells by RNA-seq. A total of 20930 genes were identified across all five cell lines combined (Supplementary Dataset 1). We found that a larger number of genes were 2-fold downregulated than 2-fold upregulated in all of the mutant lines compared to WT (Fig. 6a-b and Supplementary Datasets 2-5). Interestingly, Gene Ontology (GO) analysis (PANTHER Classification System) (41,42) (Fig. 6c, Supplementary Datasets 6-9) also showed that genes involved in DNA-dependent transcription and the regulation of transcription by RNA pol II were significantly overrepresented (p < 0.005, Bonferroni correction for multiple testing applied, Supplementary Dataset 9) within the subset of downregulated genes in the mutant cell lines, perhaps explaining why more genes are downregulated than upregulated in the mutants. It is to be noted, however, that in general, over- or underrepresentation of downregulated genes within a Biological Process-GO term does not directly imply that the process itself is downregulated. Significantly, genes involved in mesoderm and ectoderm development were also differentially expressed (downregulated) in all of the mutants (Fig. 6c), indicating that cellular differentiation of the mutant cell lines along these lineages may proceed at a rate that is altered (faster or slower than wild-type cells), which could then result in defects that cause intellectual disability (as a consequence of effects on differentiation of neural stem cells derived from ectoderm) and the dysmorphic facial features and clinodactyly seen in many of the patients (as a consequence of effects on differentiation of neural crest cells and mesenchymal stem cells derived from the ectoderm and mesoderm respectively). Other significantly represented (p < 1 x 10^-7, Bonferroni correction for multiple testing applied) biological processes include development and pattern/segment specification, as would be expected of a developmental disease such as XLID and RNA and nucleobase metabolic processes, which would influence cellular transcription (Fig. 6c). Ingenuity Pathway Analysis (Fig. 6d and Supplementary Datasets 10-13) revealed components of LXR/RXR activation pathway to be the most significantly represented in all of the mutants compared to wild-type (p < 0.001) and predicted the pathway itself to be downregulated (z-score, L254F: -2.71; A259T: -0.80; R284P: -
1.7; E339G: -3.89). This is interesting given that LXR has been shown to have a role in the generation of midbrain dopaminergic neurons, which have roles in memory and learning, from human embryonic stem cells (43).

Other significantly represented (p < 0.001) pathways predicted to be downregulated (z-score ≤ -2) in at least two of the four mutants were the Acute Phase Response pathway and Osteoarthritis pathway (Fig. 6d and Supplementary Datasets 6-9). RT-qPCR analysis was performed on GPR18, which is only slightly upregulated (3 to 7-fold by RNA-Seq), from biological triplicates of the wild-type and mutant cell lines, to orthogonally validate the directionality and fold change of the expression data (Supplementary Fig. 4).

Discussion

Advances in genome sequencing and the reduction in associated costs have made it possible in the past decade or so to investigate the etiology of rare (genetic) disorders (2). Etiological studies on cohorts of patients with ID, however, show that 29% of these individuals still have an unknown (genetic or otherwise) cause of ID, suggesting that there remains scope for the use of massive parallel sequencing approaches in the clinic for the investigation of disease causality (2). This facilitates not only genetic counseling for families but may, in some cases, inform strategies for therapeutic intervention for patients. From a biochemical standpoint, large-scale sequencing studies reveal the involvement of novel genes/proteins in disease, as in the case of OGT and XLID, allowing for a detailed examination and thereby understanding of the roles of such genes/proteins in specific cellular processes/pathways in both disease and normal physiology.

Three missense mutations were previously shown to segregate in patients with XLID (26-29) and in this study we describe two more novel mutations (Figs. 1-2). OGT-dependent XLID appears to be syndromic, however, not all phenotypes found in patients with one mutation are present in patients with another mutation (Table 1). This is conceivably due to the fact that OGT has a plethora of substrates/interactors and that while all of the mutations may affect a common pool of these substrates, they may also each affect a unique subset, as all are present in slightly different regions of the protein-protein interaction domain, TPRs, of OGT. Our previous study on the family with the OGT-L254F mutation (27) and the family with the OGT-E339G mutation reported here show that related patients with the same mutations also exhibit a range of severity of phenotypes as well as unique phenotypes, indicating there is variable expressivity and incomplete penetrance. Nonetheless, a phenotype does emerge across all patients consisting of intellectual disability, developmental delay, dysmorphic features and eye abnormalities. Many of the clinical findings are not uncommon in and of themselves in the XLID population, but a male presenting with this constellation of clinical findings should perhaps be considered for OGT testing.

In an attempt to study the direct effects of the XLID mutations on OGT stability and function, we purified recombinant OGT variants and subjected these to size-exclusion chromatography, the themofluor assay and both end-point and kinetic activity assays. We observed no adverse effects on the ability of the OGT variants to dimerize (Fig. 3a) and found modest decreases in their thermodynamic stability compared to OGT-WT (Fig. 3b). The end-point activity assays showed that all of the OGT variants investigated in this study were able to glycosylate the substrate CK2α. They were also able to glycosylate and proteolytically cleave HCF1 (Fig. 4a-b), although this assay may not detect subtle changes in the reaction rates, which could be enough to cause some of the patient phenotypes observed, given mutations in the gene encoding HCF1 have previously been implicated in XLID pathogenesis (36-38). Analysis of the kinetics of CK2α glycosylation revealed that while catalytically competent, the OGT variants demonstrated reduced turnover (Fig. 4c, Table 2). Overall, these findings, while important to establish, are perhaps not surprising given that the mutations all cluster to the TPR domains and not the catalytic domains of the enzyme.

To delineate the molecular mechanisms underpinning this developmental disease, we introduced all of the patient mutations into a male human embryonic stem cell line. We chose this over deriving iPSCs from patient-fibroblasts in order to eliminate effects arising from the different genetic backgrounds, which could
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confound comparative analyses. Examination of the steady-state global O-GlcNAc, OGT and OGA levels revealed no significant changes in these isogenic cells expressing the OGT variants compared to OGT-WT (Fig. 5a-d). This, combined with the contrasting observation that there are changes in the rate of catalysis by the OGT variants (Fig. 4c) suggests the need to investigate temporal dynamics of an in-depth O-GlcNAcome beyond what can be observed by a steady-state 1-D Western blot. We are currently in the process of developing such an approach. One hypothesis to be tested is that alterations in the kinetics of O-GlcNAc cycling on certain substrates, rather than the complete loss of O-GlcNAc on these substrates, is disability-causing.

OGT interacts with a number of proteins in the cell through its N-terminal TPRs (44). While the TPRs are also implicated in the recognition of certain substrates for O-GlcNAcylation (15), it has been shown using hypomorphic OGT mutants of the fly that post-pupal development proceeds in the near-absence of the catalytic activity of OGT (45), suggesting that the scaffolding function of OGT in forming functional protein complexes is also critical. Given that the XLID mutations lie in the TPR domain of OGT, we also hypothesize that an altered OGT-interactome in these patients may be responsible for disease, and are in the process of defining the differential OGT-interactome using multiple parallel approaches including looking at cell type specific interactomes derived from the human embryonic stem cells harboring the disorder-causing mutations.

Because OGT and protein O-GlcNAcylation have roles in transcription (4,7,16), we subjected mRNA samples from the OGT variant and OGT-WT expressing stem cells to a differential transcriptomics analysis using RNA-seq (Fig. 6a-d, Supplementary Datasets 1-11). Our data show, as predicted from the differences in patient phenotypes, that a common pool of genes is regulated by all of the mutants but that each mutant also regulates the expression of a unique subset of genes (Fig. 6a-b). While all of the cell lines expressed comparable levels of the pluripotency markers POU5F1 (Oct4), SOX2 and NANOG (Supplementary Dataset 1), GO analysis on genes that were found to be differentially regulated in all of the mutants revealed them to be involved in ectoderm and mesoderm development (Fig. 6c). Pathway analysis of all of the differentially expressed genes in each of the mutants led to the identification of common pathways affected in OGT-dependent XLID although the members of the pathway affected or the extent to which members found in all mutants were affected was different between each of the mutants (Fig. 6d). Modulation of the LXR/RXR transcriptional network is of particular interest given the established role of this pathway in cell fate specification and neurogenesis (46,47).

In conclusion, we have identified a list of genes that are differentially expressed in the XLID OGT-TPR variant expressing cells that clearly correlate with patient phenotypes. We are now taking both biased and unbiased approaches using our isogenic stem cells and derived neuronal lineages to reveal the impact of these variants on the cell type-specific dynamic O-GlcNAcome and OGT interactome to provide mechanistic insight into these XLIDs.

**Experimental Procedures**

**Study samples**

Individuals with XLID and control relatives with normal cognitive function were recruited from the Children’s Hospital of Michigan (Detroit, Michigan) and the Genetic Health Services, Northern Hub, New Zealand (Auckland, New Zealand). Human subject research protocols for these studies were approved by the respective institutional review boards and abide by the Declaration of Helsinki principles. Informed consent was obtained from each study member and/or their parents or legal guardians. All individuals were found to have a normal karyotype and a negative molecular test for Fragile X syndrome.

**Full exome sequencing and validation**

Whole exome sequencing of genomic DNA from Patient 1, his parents and an unaffected brother was performed through Ambry’s (Ambyr Genetics) clinical diagnostic exome assay. DNA was isolated from whole blood of the subjects and short tandem repeat markers were used to confirm the relationship of proband and parents. Samples were prepared using the SeqCap EZ VCRome 2.0 (Roche NimbleGen). Each DNA sample was sheared, adaptor ligated, PCR-
amplified and incubated with the exome baits. Captured DNA was eluted and PCR amplified. Final quantified libraries were seeded onto an Illumina flow cell and sequenced using paired-end, 100 cycle chemistry on the Illumina HiSeq 2000 or HiSeq 2500.

Initial data processing, base calling, alignments and variant calls were generated using various bioinformatics tools. Approximately 90% of the bases had quality score of Q20 or higher, which translated to an expected base-calling rate of 99% and error rate of 1:100. Exons and at least 2 bases into the 5' and 3' ends of all introns were analyzed. 90% of exome was covered at 10 times or higher depth, sufficient for high quality heterozygous or homozygous variant calling for germline variants. Exons plus at least 2 bases into the 5' and 3' ends of all the introns were analyzed and reported. Sequencing of the mitochondrial genome and screening for characterized mutations were performed. The mean depth of coverage for targeted mitochondrial bases was greater than 1,000 x. Variants were analyzed for population frequency, predicted functional impact, nucleotide and amino acid conservation, and biochemical nature of amino acid substitution. Human Gene Mutation Database (HGMD), the Single Nucleotide Polymorphism database (dbSNP), 1000 genomes, HapMap data and online search engines (e.g. Pubmed) sites were used to search for previously described gene mutations and polymorphisms. Variants were then filtered based on possible inheritance models and family cosegregation.

Alteration(s) detected via exome sequencing with Q-score and read depth above established confidence thresholds did not systematically undergo confirmation by Sanger sequencing. But Sanger sequencing was used for validation of candidate variant and family cosegregation studies. Candidate mutations were evaluated from among the genes in Ambry’s (Ambry Genetics) internal, dynamic gene database, which classifies genes as characterized or novel. Characterized genes include those currently understood to be associated with a clinical phenotype based on data from the medical literature, human gene mutation databases, and novel gene findings identified through clinical exome. Novel genes are those not currently known to underlie a genetic condition. Characterized genes were analyzed first. As no positive findings were identified in characterized genes, reflex analysis of novel genes occurred. Each candidate mutation was analyzed by a board certified molecular geneticist/laboratory director to identify the most likely causative mutation(s). The evidence to support the likelihood that a novel gene alteration was involved in the patient’s phenotype included knowledge of the gene function based on in vitro or in vivo functional studies, knowledge of the gene family or gene pathway, and familial cosegregation analysis.

_X-chromosome exome sequencing and validation_

DNA from Patient 2 was extracted from blood using the QIAamp DNA blood maxi kit (Qiagen). From this, X-exome sequencing, followed by data analysis, was performed as described before (48). Confirmation of variants and segregation analysis in further family members, including Patient 3, was performed by Sanger sequencing using the following oligos: g1F: AGC ATT ACC AGC CAT TAG GC; g1R: TCT CTT GAA CTG TGG GTT AAT GT.

_Plasmids and site-directed mutagenesis_

Constructs used in this study have been described before (27). The plasmid encoding the TPR domain of OGT was obtained from Elena Conti (Max Planck Institute for Biochemistry). Site-directed mutagenesis was performed as described before (49).

_Protein expression and purification_

All proteins were expressed in _E. coli_ and purified as described before (14,27). Briefly, all proteins except for ncOGT were purified to homogeneity using affinity chromatography followed by size-exclusion chromatography. ncOGT was purified just using affinity chromatography. Protein preparations were quantified using a NanoDrop™ Spectrophotometer (Thermo). For ncOGT, relative quantification using BSA standards was performed, whereby Coomassie G-250-stained SDS-PAGE gels were scanned on an Odyssey ® CLx Imaging System (LI-COR Biosciences) and bands of interest quantified using the Image Studio™ software (LI-COR Biosciences).
Thermal stability assay

The thermal stability of the proteins was measured using Differential Scanning Fluorimetry (Thermofluor assay). Briefly, 45 µl of 1 mg/ml solutions of protein in a buffer comprised of 25 mM Tris-HCl pH 7.5 and 150 mM NaCl were combined with 5 µl of 200 x SYPRO Orange diluted from a 5000 x concentrate (Thermo) in the aforementioned buffer. The mixture was allowed to incubate at room temperature for 15 min. The sealed PCR plate containing the mixtures was exposed to a temperature gradient program (25-95 °C, 0.5 °C intervals) with fluorescence intensity measured after a 30 s dwell time at each particular temperature in a MyIQ™ Single Color Real-Time PCR Detection System (485/20 nm for excitation and 530/30 nm for emission) equipped with an iCycler (Biorad). The fluorescence intensities plotted against their corresponding temperatures were exported to Microsoft Excel and then to Prism (GraphPad). Wells containing no protein, but only buffer and SYPRO Orange were used for background subtraction. Fluorescence values obtained between 40-65 °C were fitted to the Boltzmann Sigmoidal equation to obtain T_m values. All experiments were performed in technical triplicate.

CK2α glycosylation and HCF1 proteolysis

The Western blotting assays for CK2α glycosylation and HCF1 proteolysis were carried out using full length OGT as described before (27). Reactions were run either on a 10% or 4-15% gradient Mini-PROTEAN® TGX Stain-Free™ Precast Gel (Biorad) before being blotted onto nitrocellulose membranes, explaining the differences in the appearance of specific bands across independent experiments. Blots were probed with the antibodies described before (27). All experiments were performed in biological triplicate (with three separate preparations of nOGT).

Enzyme kinetics

The UDP-Glo™ Glycosyltransferase Assay kit (Promega) was used to measure the steady state kinetics. Recombinantly purified full length OGT was used in these experiments. Reaction mixtures contained 50 nM of the OGT variants (20 nM of OGT-WT), 500 µM of ultrapure donor substrate UDP-GlcNAc (Promega) and varying concentrations (0.234 µM to 60 µM of the acceptor substrate, human recombinant CK2α (1-365), in a final volume of 18 µl. The reaction buffer was made up of 20 mM Tris-HCl pH 8.0, 20 mM MgCl₂, 150 mM NaCl and 1 mM DTT. Reactions were incubated for 90 min at room temperature and stopped by the addition of 18 µl of the UDP detection reagent prepared as recommended by the manufacturer. 10 µl of the samples were transferred to a white 384-well plate and the luminescence measured using the GloMax® Multi Detection System (Promega). Relative luminescence units were converted to amount of UDP released using a UDP standard curve. All assays were performed in technical triplicate and measurements were corrected for background emission from reactions containing no acceptor substrate. For all assays performed, UDP-GlcNAc turnover was under 10% and reactions were linear past the 90 min incubation time used. Non-linear regression curves were fitted with Prism (GraphPad).

Cell lines and culture maintenance

RUES-1 male human ES cells were obtained from the Laboratory of Molecular Embryology at the Rockefeller University. XLID OGT-TPR point mutations in these cells were introduced using CRISPR/Cas9 as a pay-for-service by Applied StemCell. Cells were maintained in TeSR™ E8™ media (STEMCELL Technologies) in 5% CO₂. Media was changed every day and at appropriate confluency, cells were split 1:3 or 1:5 using an enzyme-free dissociation buffer (1X DPBS – Ca²⁺ and Mg²⁺ free (Thermo) supplemented with sterile-filtered 0.5 mM EDTA and 150 mM NaCl).

Cell lysis and Western blotting

Harvested stem cells were lysed using 50 mM Tris-HCl pH 7.5, 150 mM NaCl, 1% IGEPAL® CA-630 and 1 mM DTT supplemented with 10 µM PUGNAc (hexosaminidase inhibitor), 1 x EDTA-free Protease Inhibitor Cocktail Set V (Millipore) and 1 x Phosphatase inhibitor Cocktail Set II (Millipore). Lysates were resolved on Bolt™ 4-12% Bis-Tris Plus Gels (Thermo) and blotted onto nitrocellulose membranes.
Membranes were probed with the following antibodies: O-GlcNAc CTD 110.6 (1:500, ascites generated in the lab of Gerald Hart at Johns Hopkins University), O-GlcNAc RL2 (1:1000, Thermo, MA1-072), OGT DM17 (1:1000, Sigma, O6264), OGT F-12 (1:1000, Santa Cruz, sc-74546), OGA/MGEA5 (1:1000, Proteintech, 14711-1-AP), OGA/NCOAT G-12 (1:1000, Santa Cruz, sc-376429) and β-Actin 13E5 (1:1000, Cell Signaling, 4970). Secondary antibodies were purchased from LI-COR and used at a dilution of 1:10000. Experiments were conducted in biological triplicate.

**RNA-seq and bioinformatics**

Total RNA was isolated from cells harvested from confluent 60 mm dishes using the RNeasy Plus Mini Kit (Qiagen). The concentration and integrity of the extracted total RNA were estimated using the Qubit® 2.0 Fluorometer (Invitrogen) and Agilent 2100 Bioanalyzer (Applied Biosystems), respectively. Five hundred nanograms of total RNA was required for downstream RNA-seq applications. Polyadenylated RNAs were isolated using NEBNext Magnetic Oligo d(T)25 Beads. Next, first strand synthesis was performed using NEBNext RNA first strand synthesis module (New England BioLabs). Immediately, directional second strand synthesis was performed using NEBNext Ultra Directional second strand synthesis kit. The NEBNext® DNA Library Prep Master Mix Set for Illumina® was then used to prepare individually bar-coded next-generation sequencing expression libraries as per the manufacturer's recommended protocol. Library quality was assessed using the Qubit 2.0 Fluorometer, and the library concentration was estimated by utilizing a NanoDrop™ spectrophotometer (Thermo). iScript cDNA synthesis kit (Biorad) was used for cDNA synthesis and iTaq™ Universal SYBR®Green Supermix (Biorad) or iQ™ SYBR® Green Supermix (Biorad) was used for qPCR, performed as described before (50,51). Briefly, the ΔΔCt method was used to quantify differences in GPR18 gene expression and ACTB (β-Actin) was used as the housekeeping gene. Experiments were performed in biological triplicate and data analysis was performed as described before (50,51) Primer sequences are provided in the Supporting Information (Supplementary Table 1).
We thank Dr. Gerald Hart (Johns Hopkins School of Medicine), Dr. Suzanne Walker (Harvard University), Dr. Alison Nairn (University of Georgia), and Dr. Winship Herr (University of Lausanne) for providing plasmids, and/or technical advice. This work was funded in part by a grant from the W.M. Keck Foundation (L. W., co-PI); a NIGMS, National Institutes of Health (NIH) Grant P41GM103490 (L. W., co-PI); a grant from the South Carolina Department of Disabilities and Special Needs (SCDDSN) (C. E. S.); a NINDS, National Institute of Health (NIH) grant RO1N5073854 (C. E. S.); and grants from the National Health and Medical Research Council of Australia APP1091593 and APP1041920 (J. G.). The data reported in this paper have been deposited in the Gene Expression Omnibus (GEO) database, (accession no. GSE110616).

Conflict of Interests
The authors declare that there are no conflicts of interests. The content of this article is solely the responsibility of the authors and does not necessarily represent the official views of the National Institutes of Health.

Author contributions
N. S. and L. W. conceived and coordinated the study and wrote the paper. All authors designed, performed, and/or analyzed the experiments presented, and reviewed and approved the final submitted version of the manuscript.
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Table 1: Patient phenotypes caused by the XLID-associated variants of OGT

| Genotype | Vaidyanathan et al. (27) | This study | Willems et al. (29) | Bouazzi et al. (28) | This study | Total |
|----------|--------------------------|------------|---------------------|---------------------|------------|-------|
| cDNA (NM_181672.2) | 762 G>T | 775 G>A | 851 G>C | 955 G>A, cosegregates with MED12 c. 5922 G>T | 1016 A>G |
| Skewing in X-chromosome inactivation in carrier | + | ND | NR | - | ND |
| Protein | L254F | A259T | R284P | A319T, cosegregates with MED12 p. Q1974H | E339G |
| Phenotype | | | | | |
| Affected males | 3 | 1 | 1 | 3 | 2 | 10 |
| Intellectual disability | 3/3 | 1 | 1 | 3/3 | 2/2 | 10/10 |
| Developmental delay | 3/3 | 1 | 1 | 3/3 | 2/2 | 10/10 |
| Behavioral problems | 0/3 | 1 | 1 | 2/3 | 1/2 | 5/10 |
| Brain abnormalities | NA | 1 | 1 | 0/1 | 0/2 | 2/5 |
| Cardiac symptoms | 1/3 | 0 | 1 | NA | 0/2 | 2/7 |
| Dysmorphic features | 3/3 | 1 | 1 | 2/3 | 2/2 | 9/10 |
| Clinodactyly | 3/3 | 0 | 1 | 1/3 | 1/2 | 6/10 |
| Eye abnormalities | 3/3 | 1 | 1 | 2/2 | 0/2 | 7/9 |
| Genital/reproductive abnormalities | 2/3 | 0 | 1 | NA | 0/2 | 3/7 |
| Ear abnormalities/hearing impairment | 0/2 | 1 | 0 | 1/3 | 2/2 | 4/9 |
| Microcephaly | 1/3 | 1 | 1 | 0/3 | 0/2 | 3/10 |
| Hypothyroidism | NA | 1 | NA | NA | NA | 1/1 |
| Epilepsy/seizures | 0/3 | 1 | NA | 0/1 | 0/2 | 1/7 |

NA: Not available. Numbers indicate number of affected patients over number of patients examined for the phenotype.
Table 2: Michaelis-Menten kinetics of the XLID-associated TPR variants of OGT

| OGT variant | $K_m$ (µM)   | $k_{cat}$ (s$^{-1}$)    | $k_{cat} / K_m$ (s$^{-1}$M$^{-1}$) |
|-------------|------------|------------------------|----------------------------------|
| WT          | 7.4 ± 0.6  | 4.5 x 10$^{-3}$ ± 0.1 x 10$^{-3}$ | 640 ± 80                        |
| L254F       | 13 ± 3     | 3.5 x 10$^{-3}$ ± 0.3 x 10$^{-3}$ | 270 ± 200                       |
| A259T       | 9.2 ± 0.6  | 3.5 x 10$^{-3}$ ± 0.1 x 10$^{-3}$ | 390 ± 30                        |
| R284P       | 8.9 ± 1.1  | 3.5 x 10$^{-3}$ ± 0.1 x 10$^{-3}$ | 390 ± 100                       |
| A319T       | 15 ± 2     | 3.3 x 10$^{-3}$ ± 0.1 x 10$^{-3}$ | 230 ± 50                        |
| E339G       | 12 ± 2     | 4.0 x 10$^{-3}$ ± 0.2 x 10$^{-3}$ | 330 ± 100                       |
Figure legends

Figure 1: Pedigrees and phenotypes of the patients.
(a) Three-generation pedigree of Patient 1 (with OGT c. 775 G>A, p. A259T). Proband is identified by the arrow. Circles denote females; circles with dots, carrier females confirmed using exome sequencing or Sanger sequencing; open squares, unaffected males and closed squares, males with XLID.
(b) MRI of brain of Patient 1 at 7 years of age. Sagittal T1 image reveals extremely thin corpus callosum particularly the posterior body and splenium (short thin arrow), short appearing clivus (thick arrow), and relatively hypoplastic posterior arch of first cervical vertebra (long thin arrow).
(c) MRI of brain of patient 1 at 7 years of age. Axial T2 FLAIR image reveals mild ventriculomegaly (arrow).
(d) Four-generation pedigree of patients 2 and 3 (with OGT c. 1016 A>G, p. E339G). Probands are identified by arrows and diagram has same coding as in (a).

Figure 2: All known missense mutations in the TPR domain that are causal for XLID.
(a) Schematic of the domain organization of OGT. The N-terminal TPR domain is depicted in green with XLID variants noted, the intervening domain in pink and the lobes of the catalytic domain in purple.
(b) Model of full length OGT based on the structure of the first 11.5 TPRs (PDB ID: 1W3B (14)) and the structure of the catalytic domain with the last 2.5 TPRs (PDB ID: 4AY5 (52)). The domains in the model are colored as in the schematic and the side chains of the residues in the TPR that are mutated in XLID are depicted as spheres with black carbons.
(c) Close-up of residues that are mutated in XLID. Panels on the left show the wild-type residue in the amino acid position indicated, while panels on the right show the replacing residue modeled into the structure (side-chains shown as sticks with black carbons). Surface representation shown in both panels is from OGT-WT. Side-chains of neighboring residues that could clash are shown as sticks with yellow carbons.

Figure 3: OGT XLID-TPR mutants form functional dimers but demonstrate decreased thermal stability
(a) The TPR domain is a constitutive homodimer in solution. Samples of wild-type and mutant TPR domains were analyzed on a Superdex 200 size-exclusion column. The elution profiles are plotted as A280 against elution volume. Elution time of standards of known molecular weight are marked above; 1: IgG – 158 kD, 2: human albumin – 66 kD and 3: ovalbumin – 44 kD. The 6 x His-tagged TPR dimer has an expected molecular weight of ~90 kD and an apparent molecular weight of ~120 kD (14) due to its elongated, non-globular structure.
(b) Thermal denaturing curves of wild-type or mutant OGT TPR domains. The melting temperatures (Tm, the temperature at which both the folded and unfolded states of a protein are equally populated at equilibrium) of the proteins are indicated. Fluorescence of Sypro orange is plotted against temperature. The data were fitted to Boltzmann sigmoidal curve equation using Prism (GraphPad). Experiments were performed in triplicate and error bars represent mean ± S.E.

Figure 4: XLID-TPR variants of OGT are active enzymes but exhibit impaired glycosyltransferase kinetics
(a) O-GlcNAcylation of recombinant CK2α (1-365) protein substrate by wild-type or mutant OGT detected using the anti-O-GlcNAc antibody RL-2. Reactions were carried out at 37°C for 1.5 h. Reactions containing no UDP-GlcNAc were included as a negative control. Samples were resolved in a 10% (A259T, R284P and E339G) or a 4-15% gradient (A319T) SDS-PAGE gel, explaining the differences in the appearance of specific bands across independent experiments. Blots shown are representative of three independent replicates.
Proteolytic cleavage of a recombinant HCF-1 derived construct (HCF1-rep1) by wild-type or mutant OGT. Uncleaved and cleaved HCF1-rep1 were detected using an anti-GST tag antibody while the O-GlcNAcylation of the substrate/products were detected using the anti-O-GlcNAc antibody RL-2. Reactions were carried out at 37°C for 6 h. Reactions containing no UDP-GlcNAc were included as a negative control. Samples were resolved in a 10% (A259T and R284P) or a 4-15% gradient (A319T and E339G) SDS-PAGE gel, explaining the differences in the appearance of specific bands across independent experiments. Reactions containing OGT-R284P were run on the opposite end of the same gel as those containing OGT-WT. OGT-A319T and OGT-E339G were tested in parallel and were therefore run alongside a single set of the OGT-WT containing control reactions. Blots shown are representative of three independent replicates.

(c) Michaelis-Menten kinetics of wild-type or mutant OGT measured using varying amounts of CK2α (1-365) protein substrate and a fixed concentration of UDP-GlcNAc in excess of the \( K_m \). Reactions were carried out for 90 min at room temperature and read using the UDP-Glo assay system (Promega). Data points were fitted to the Michaelis-Menten equation using Prism (GraphPad). Experiments were performed in triplicate and error bars represent mean ± S.E.

**Figure 5:** Human embryonic stem cells expressing the XLID OGT-TPR variants show no gross changes in steady-state O-GlcNAc, OGT and OGA levels

(a) Endogenous wild-type or variant OGT expressing RUES-1 hES cell lysates were probed for O-GlcNAc using the anti-O-GlcNAc antibody, CTD 110.6 or

(b) a second anti-O-GlcNAc antibody, RL2.

(c) Samples were also probed for OGT and OGA using the antibodies DM-17 and G-12 respectively or

(d) the antibodies F-12 and 14711-1-AP respectively.

All membranes were probed using an anti-β-actin antibody as a loading control. Blots shown are representative of three biological replicates.

**Figure 6:** Differential transcriptomes of the hES cells expressing XLID OGT-TPR variants reveal changes in genes associated with neural development

(a) Number of genes that are >2-fold upregulated in each of the TPR mutants analyzed compared to wild-type.

(b) Number of genes that are >2-fold downregulated in each of the TPR mutants analyzed compared to wild-type.

(c) Gene Ontology (GO) analysis (PANTHER Classification System) showing significantly represented (over- or underrepresented - \( p < 1 \times 10^{-7} \), Bonferroni correction for multiple testing applied) biological processes in genes that are differentially expressed (up- or down-regulated) in at least three mutants compared to wild-type.

(d) Pathway analysis (Ingenuity Pathway Analysis, Qiagen) showing significantly represented (over- or underrepresented - \( p < 0.001 \)) pathways in genes found in each of the mutants compared to wild-type. Z-scores for the pathways in each of the mutants is shown to the right of the bars. A z-score of \(< -2\) denotes the pathway as a whole is downregulated while a score of \( > +2\) denotes it is upregulated. Only those significantly represented pathways with a z-score of \( \pm 2\) in at least two mutants are shown.
Figure 1

a. OGT c. 775 A>G, p. A259T

b. [Brain MRI images]

c. [Brain MRI images]

d. OGT c. 1016 A>G, p. E339G

[Family trees showing genetic inheritance patterns and medical conditions such as depression, breast cancer, ovarian tumor, and miscarriage.]
Figure 2

(a) Schematic representation of the TPR and GT41 domains of the protein, with labeled amino acid variants.

(b) Detailed view of the molecular structure, highlighting mutations at specific positions.

(c) Enlarged views of the amino acid residues A259, T259, R284, L254, and E339, showing the effects of variant mutations L254F, A259T, R284P, A319T, and E339G.
Figure 4

a

| UDP-GlcNAc | - | + | - | + | - | + | - | + | - | + | - | + |
|------------|---|---|---|---|---|---|---|---|---|---|---|---|
| CK2α       | + | + | + | + | + | + | + | + | + | + | + | + |
| OGT        | WT | A259T | WT | R284P | WT | A319T | WT | E339G |
| kD         | 37 | 37 | 37 | 37 | 37 | 37 | 37 | 37 |
| IB: O-GlcNAc |
| IB: CK2α   |
| IB: OGT    |

b

| UDP-GlcNAc | - | + | - | + | - | + | - | + | - | + | - | + |
|------------|---|---|---|---|---|---|---|---|---|---|---|---|
| HCF1-rep1  | + | + | + | + | + | + | + | + | + | + | + | + |
| OGT        | WT | A259T | WT | R284P | E339G | WT | A319T |
| kD         | 50 | 50 | 50 | 50 | 50 | 50 | 50 |
| IB: GST (HCF1-rep1) |
| IB: O-GlcNAc |
| IB: OGT    |

c

$k_{cat}$ (s$^{-1}$) vs. CK2α 1-365 [µM]

- WT
- L254F
- A259T
- R284P
- A319T
- E339G
Figure 5

(a) Western blot analysis showing the expression levels of O-GlcNAc (CTD 110.6) and β-Actin proteins in WT, L254F, A259T, R284P, and E339G samples.

(b) Western blot analysis showing the expression levels of O-GlcNAc (RL2) and β-Actin proteins in WT, L254F, A259T, R284P, and E339G samples.

(c) Western blot analysis showing the expression levels of OGT (DM17) and OGA (G-12) proteins in WT, L254F, A259T, R284P, and E339G samples.

(d) Western blot analysis showing the expression levels of OGT (F-12) and OGA (14711-1-AP) proteins in WT, L254F, A259T, R284P, and E339G samples.
Figure 6

(a) Upregulated

(b) Downregulated

(c) GO - Biological Processes

(d) Ingenuity Pathway Analysis
O-GlcNAc transferase missense mutations linked to X-linked intellectual disability
deregulate genes involved in cell fate determination and signaling
Nithya Selvan, Stephan George, Fatema J Serajee, Marie Shaw, Lynne Hobson, Vera M
Kalscheuer, Nripesh Prasad, Shawn E Levy, Juliet Taylor, Salim Afitmos, Charles E.
Schwartz, Ahm M Huq, Jozef Gecz and Lance Wells

J. Biol. Chem. published online May 16, 2018

Access the most updated version of this article at doi: 10.1074/jbc.RA118.002583

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