Congenital disorders of glycosylation

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Abstract: Congenital disorders of glycosylation are a genetically and clinically heterogeneous group of >130 diseases caused by defects in various steps along glycan modification pathways. The vast majority of these monogenic diseases are autosomal recessive and have multi-systemic manifestations, mainly growth failure, developmental delay, facial dysmorphisms, and variable coagulation and endocrine abnormalities. Carbohydrate deficient transferrin (CDT) and protein-linked glycan analysis with mass spectrometry can diagnose some subtypes of congenital disorders of glycosylation (CDG), while many currently rely on massively parallel genomic sequencing for diagnosis. Early detection is important, as a few of these disorders are treatable. Molecular and biochemical techniques continue to further our understanding of this rapidly expanding group of clinically and genetically diverse disorders.

Keywords: Congenital disorders of glycosylation; carbohydrate deficient transferrin; carbohydrate deficient glycoprotein syndrome; PMM2-CDG; MPI-CDG; ALG6-CDG; N-linked glycosylation; O-linked glycosylation; lipid glycosylation defects; congenital disorders of glycosylation (CDG)


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Overview

Glycosylation is the process of adding sugar residues to proteins and lipids in different cellular pathways. Congenital disorders of glycosylation (CDG) are a genetically and clinically heterogeneous group of over a hundred diseases caused by defects in various steps along glycan synthesis or modification pathways. Most of these monogenic diseases are autosomal recessive in inheritance, but autosomal dominant and X-linked forms have also been described.

CDG typically present with multi-systemic manifestations, most commonly developmental delay, failure to thrive, hypotonia, neurologic abnormalities, hepatopathy, and coagulopathy. Affected individuals may also present with eye, skin, and cardiac disease, as well as facial dysmorphisms. Although neurologic changes and cognitive delays are seen in the majority of affected individuals, there are certain cases and even types that do not have neurologic manifestations. Given the broad clinical and genetic etiology of CDG, clinical diagnosis relies on high index of suspicion in multi-systemic disease.

Serum carbohydrate deficient transferrin (CDT) analysis is the first-line screening test in patients with suspected CDG, but is limited in detection to N-glycosylation defects with sialic acid deficiencies. Next-line testing includes dolichol-linked glycan analysis and genetic testing. Early diagnosis of this group of exponentially growing diseases is important, as some CDGs are treatable. Treatment for glycosylation defects is mainly supportive, although targeted therapies are available for MPI-CDG, SLC35C1-CDG, PIGM-CDG, and PGM1-CDG. Details about these treatments are under the “Targeted Therapies and Prognosis” section below. The focus of this review will be on the most common types of CDG with recognizable
phenotypes or treatments, with the target audience being primary care providers.

**Epidemiology**

The incidence and prevalence of all types of CDG in aggregate have not been well established, although patients have been reported worldwide from almost every ethnic background and both sexes are equally affected. The estimated prevalence in European and African American populations is 1/10,000 based on carrier frequencies of known pathogenic variants in 53 genes (1-4). The prevalence of the most commonly diagnosed CDG, PMM2-CDG, ranges from 1/20,000 in Dutch populations and 1/77,000 in Estonia based on isolated reports (5,6). To date, fewer than 100 cases have been reported for most CDG types.

**Biochemical classification and nomenclature**

Broadly, CDG currently are classified into four categories—(I) N-linked glycosylation, (II) O-linked glycosylation, (III) combined N- and O-linked/multiple glycosylation, and (IV) lipid and glycosylphosphatidylinositol (GPI) anchor biosynthesis defects.

The N-linked protein glycosylation defect PMM2-CDG, previously known as CDG type Ia, was the first CDG to be reported by Jaeken in 1980 and remains by far the most common CDG to date (7). PMM2-CDG was initially named “carbohydrate-deficient glycoprotein syndrome” due to multiple serum glycoprotein abnormalities seen by isoelectric focusing of serum transferrin in affected individuals. Historically, CDG were classified by patterns of transferrin isoform analysis—type I patterns were attributed to dolichol-linked glycan assembly and transfer defects localizing to the cytoplasm or ER, and type II patterns were attributed to processing defects in the Golgi apparatus. From this branch point, CDGs were then named alphabetically in order of discovery.

With the advent of widespread molecular diagnostics, CDG nomenclature was updated in 2008 to specify molecular etiology of disease, reflecting the exponential growth of pathways and disorders that did not neatly fit in the previously established dichotomous categories. Presently, CDG nomenclature is denoted by the affected gene name (non-italicized, gene names at www.genenames.org), followed by -CDG (e.g., PMM2-CDG) (8).

**Genetics**

The vast majority of congenital disorders of glycosylation are inherited in an autosomal recessive fashion, with one mutation inherited from each asymptomatic (carrier) parent. Molecular testing, usually with next-generation sequencing methods, is necessary to establish a genetic diagnosis. Parental testing for the known variant can confirm inheritance versus *de novo* occurrence. For autosomal recessive inheritance, the recurrence risk for siblings and each pregnancy of an affected individual is 25% for being affected, 50% for being an asymptomatic carrier, and 25% for being unaffected.

A handful of CDG have autosomal dominant inheritance (N-linked: GANAB-CDG, PRKCSH-CDG; O-linked: EXT1/EXT2-CDG, POFUT1-CDG, POGLUT1-CDG). Fewer are X-linked (ALG13-CDG, SSR4-CDG, PIGA-CDG, SLC35A2-CDG, ATP6AP1-CDG). Most dominant and some X-linked forms of CDG are due to *de novo* mutations. Specific diseases and genes are described below in the “pathophysiology” section.

Mutation data for all published genes for CDG are available at the Human Gene Mutation Database (http://www.hgmd.cf.ac.uk/ac/index.php). Information about specific gene variants is available at the Leiden Open Variation Database with integrated in silico pathogenicity tools (http://www.lovd.nl/3.0/home). Clinical synopses for specific genes can be found at the Online Mendelian Inheritance in Man (http://www.omim.org/) or in a more limited scope at GeneReviews (http://www.ncbi.nlm.nih.gov/books/NBK1116/). Given the small number of affected patients for most CDG subtypes, genotype-phenotype correlation is difficult to establish.

**Pathophysiology**

Over 130 types of CDG have been reported to date (9,10). Given the ubiquitous presence of glycosylation pathways, CDG are extremely diverse in their biochemical pathogenesis. Numerous proteins and lipids (i.e., sphingolipids and glycolipids) undergo glycosylation with monosaccharides and/or oligosaccharides, collectively termed glycans, in different cellular compartments. Their subcellular locations are diverse, but most defects occur within the ER or Golgi apparatus. Clinical features and genetic etiology of more common CDG by pathway is summarized in Table S1.
Amongst proteins, glycans are described by their linkage to the polypeptide chain—N-glycans are attached to the amide group of asparagine (Asn) while O-glycans are attached to the hydroxyl group of either serine or threonine. N-glycan synthesis requires the stepwise construction of nucleotide-linked sugars in the cytosol, assembly in the endoplasmic reticulum, and processing in the Golgi apparatus. In contrast, O-glycan synthesis requires assembly but no processing, therefore O-glycosylation defects occur predominantly in the Golgi apparatus.

**N-linked protein glycosylation defects**

N-glycosylation involves the covalent attachment of carbohydrate structures to the side chain amide group of Asn residues within a consensus Asn-X-Ser/Thr acceptor site, translocation of the substrate polypeptide to the endoplasmic reticulum for remodeling, and further modification of the N-glycan chain within the Golgi (11,12). Defects anywhere along the synthesis, assembly, and processing pathway can lead to clinical disease.

PMM2-CDG is caused by pathogenic variants in the phosphomannomutase 2 (PMM2) gene, leading to deficiency of the PMM2 enzyme that catalyzes the cytosolic conversion of mannose-6-phosphate to mannose-1-phosphate in the second step of guanosine diphosphate (GDP) mannose synthesis. Most patients harbor compound heterozygous pathogenic missense mutations (www.lovd.nl/PMM2). The most common recurrent pathogenic variant p.Arg141His is found in approximately 40% of affected individuals of European ancestry, and p.Phe119Leu is also frequently found in northern Europe (1). Genotype-phenotype correlations have been reported for PMM2-CDG (3,13,14).

MPI-CDG is an autosomal recessive disorder caused by pathogenic variants in the mannose phosphate isomerase (MPI) gene leading to deficient phosphomannose isomerase (MPI). MPI normally catalyzes the first step of GDP-mannose synthesis (i.e., the conversion of fructose-6-phosphate to mannose-6-phosphate), but fructose-6-phosphate does not accumulate intracellularly since it can also be metabolized by the glycolytic pathway. Therefore, although biochemically similar to PMM2-CDG, MPI-CDG does not cause as significant neurologic and multi-systemic involvement. CDT is also the screening test of choice for MPI-CDG, which shows a type 1 pattern. The diagnosis can then be confirmed molecularly or by fibroblast/leukocyte MPI activity.

ALG6-CDG is a recessive disease caused by mutations in ALG6, leading to abnormal attachment of three glucose molecules to dolichol-linked mannose intermediates and downstream hypoglycosylation of serum glycoproteins (15).

**O-linked glycosylation defects and combined N- and O-linked glycosylation defects**

O-glycosylation comprises the stepwise addition of carbohydrate chains to serine, threonine, and hydroxylsine residues of proteins by glycosyltransferases in the Golgi apparatus (16). Several types of O-linked glycans have been associated with human disease, named by the first sugar attached to the amino acid residue [e.g., O-N-acetylglucosaminyl for GlcNAc and O-N-acetylgalactosaminyl for N-acetylgalactosamine (GalNAc)] (17).

**Lipid glycosylation and GPI anchor biosynthesis defects**

GPI anchors are glycolipids that undergo sequential assembly in the endoplasmic reticulum and modifications within the Golgi. GPI anchor biosynthesis disorders due to enzyme deficiencies are named alphabetically by order of discovery and not chronologically by step of assembly. Once synthesized, GPI anchors reside on plasma membranes and bind hundreds of cell-surface proteins, performing a plethora of cellular functions. Most of these diseases are autosomal recessive with the notable exception of X-linked PIGA deficiency.

**Clinical manifestations**

Given the ubiquitous presence of glycosylation pathways, virtually any organ system may be involved in CDG, although most cases involve neurologic abnormalities. Some CDG present with ichthyosis, including MPDU1-CDG, DOLK-CDG, SRD5A3-CDG (Figure 1), and PIGL-CDG (18,19). Almost all CDG present with multi-systemic disease within the first few years of life, except some affect only a single organ system (i.e., retina in DHDDS-CDG; neuromuscular junction in ALG2-CDG, ALG14-CDG, CFPT1-CDG; brain in ST3GAL3-CDG, TUSC3-CDG; skin or skeletal muscle in POGLUT1-CDG, POFTU1-CDG; cartilage in EXT1/EXT2-CDG; liver in TMEM199-CDG; red blood cells in SEC23B-CDG). The age of onset and severity may range from neonatal lethal to nearly asymptomatic adulthood, and any permutation in between.
The most commonly reported constellation of symptoms includes developmental delay, failure to thrive, hypotonia, neurologic abnormalities, hypoglycemia, and variable liver, eye, skin, gastrointestinal, immunologic, skeletal, and coagulation abnormalities (19).

The complete phenotype for many CDG subtypes remains to be fully delineated due to the rarity of reported cases. Therefore, CDG should be considered in any setting of multi-systemic disease, especially in cases with a neurologic component or nonspecific developmental delay with unclear etiology.

Although the pathophysiology of multitude symptoms remains to be elucidated, the relationship between certain glycosylation pathways and specific clinical symptoms has been clarified. For instance, failure to thrive seen in many types of CDG is attributable to the hypoglycosylation and impaired formation of several glycoproteins within the insulin growth pathway including IGF-1, ALS, and IGFBP-3 (20). As we better understand this group of complex disorders, CDG are increasingly recognized in individuals with elusive diagnoses. The organ system involvements of different CDG are summarized in Table S1.

We will discuss the clinical features of most common forms and forms with targeted treatments of CDG below.

**N-linked protein glycosylation defects**

As the most commonly diagnosed CDG, the phenotype of
Figure 2 Clinical portraits of patients with PMM2-CDG from childhood to adulthood (left to right). Esotropia, long philtrum, large, hypoplastic or dysplastic ears, hypotonic facies, and downslanting palpebral fissures are seen (top row). Profile view shows the prominent chin, which is more pronounced with age (bottom row). Courtesy of NIH, Lynne Wolfe, CRNP, Donna Krasnewich, MD, PhD.

N-linked glycosylation disorders is often heralded as the classic presentation. However, the phenotypic spectrum of CDG is quite diverse, and many CDG may not present with stereotypical symptoms associated with PMM2-CDG.

**PMM2-CDG (CDG-Ia, PMM2 deficiency)**

PMM2-CDG is the most common CDG, with over 700 reported cases worldwide. It is characterized by multisystemic severe disease in infancy, neurologic disease and developmental delay in childhood, and/or stable intellectual disability in adulthood (21,22).

In infancy, PMM2-CDG presents with neurologic abnormalities typically shortly after birth, namely strabismus and abnormal eye movements, cerebellar hypoplasia, hypotonia, psychomotor retardation, ataxia, hypotonia, and hyporeflexia. Infants may also have hepatic disease, nephrotic syndrome and renal cysts, pericardial effusion and hypertrophic cardiomyopathy, failure to thrive, and multi-organ failure resulting in death within the first year of life in up to 20% of affected individuals (21,23-28).

A constellation of dysmorphic features has been described in patients with PMM2-CDG (Figures 2,3). These include a hypoplastic cerebellum, facial dysmorphisms (i.e., large, dysplastic ears), inverted nipples, and abnormal distribution of adipose tissue over the buttocks or suprapubic region which may resolve with age (14,21,29-32). Patients have been described to have an outgoing and happy demeanor.
Presentation is highly variable, although strabismus can be seen in more than 70% of affected patients (21,23,33-35). Inverted nipples and abnormal fat pads are seen in about 25–50% of patients (36).

In childhood, affected individuals may develop retinitis pigmentosa, stroke-like episodes and seizures, speech and motor delays, and peripheral neuropathy. Constitutionally, patients commonly have failure to thrive due to feeding and GI abnormalities and global developmental delay. Elevated liver transaminases without clinical consequence may be observed, which typically normalize by 5 years of age with occasional fluctuations with illness (21,24). Liver biopsies are rarely indicated in CDG unless hepatic fibrosis is suspected (1). Clinical hypothyroidism is rare, but patients with CDG should have their thyroid hormones and free T4 measured, which may show low thyroid binding globulin (TBG) and transient elevations in thyroid stimulating hormone (TSH) (37). Liver and bile duct malformations have not been reported in PMM2-CDG patients.

Adults with PMM2-CDG may live until their 7th or 8th decades with stable cognitive delay, peripheral neuropathy, and progressive thoracic and spinal kyphoscoliosis with osteopenia or osteoporosis (34). Cerebellar ataxia is an increasingly recognized symptom along with multisystemic involvement (38-40). Endocrine abnormalities including hyperprolactinemia, growth hormone release with hyperglycemia, insulin resistance, and hyperinsulinemic hypoglycemia (41,42). In affected women, hypogonadotrophic hypogonadism may lead to absent secondary sexual development or absent ovaries (41,43,44). Patients may be at increased risk of thrombosis due to decreased serum coagulation factors including factors IV, IX, and XI, antithrombin III, protein C, and protein S (29).

**MPI-CDG (CDG-Ib, mannosephosphate isomerase deficiency)**

MPI-CDG is unique because affected patients have little to no neurologic involvement and some manifestations of the disease are treatable by oral mannose (2). Symptoms are mainly hepatic-intestinal without dysmorphic features or cognitive delays. Patients typically present with recurrent
vomiting, significant hypoglycemia, failure to thrive, potentially life-threatening protein-wasting enteropathy, liver fibrotic changes, and bile duct dilatation (45-51). Patients are at increased risk for thrombotic events due to low serum concentrations of protein C and S, and anti-thrombin III.

**ALG6-CDG (glucosyltransferase 1 deficiency)**

ALG6-CDG is the second most common N-glycosylation defect characterized by similar but milder phenotype than PMM2-CDG. Patients with ALG6-CDG have failure to thrive, developmental delay, hypotonia, seizures, strabismus, ataxia, coagulopathy, and facial dysmorphisms (i.e., low-set ears, hypertelorism, and macroGLOSSIA). Similar to MPI-CDG, they may also have protein-losing enteropathy. Additionally, affected patients may have skeletal abnormalities including brachydactyly and finger malformations and scoliosis. Affected patients typically do not typically have retinitis pigmentosa or cerebellar hypoplasia (52).

**O-linked glycosylation defects and combined N- and O-linked glycosylation defects**

Due to the substantial presence of O-glycans in mucin-containing proteins including glycosaminoglycans (GAGs) and epithelial surfaces (53), disorders of GAG synthesis typically lead to skeletal dysplasias or connective tissue disease. Affected patients may present with musculoskeletal, skin, and joint abnormalities (e.g., joint laxity, multiple exostoses, chondro/osteosarcomas) in addition to neurologic symptoms (54-56). For example, N-acetylgalactosaminyltransferase 3 (GALNT3) O-glycosylates the phosphaturic hormone, FGF23, preventing proteolytic cleavage and allowing its intact secretion. GALNT3-deficiency leads to familial tumoral calcinosis, characterized by hyperphosphatemia and ectopic calcifications (57,58).

**Lipid glycosylation and GPI anchor biosynthesis defects**

Glycosphingolipids and their sialylated derivatives, gangliosides, are mainly expressed by neurons. Defects in the breakdown of gangliosides lead to accumulation and the well-characterized lysosomal storage diseases. On the opposite end, defects in ganglioside biosynthesis such as ST3GAL5-CDG and B4GALNT1-CDG are exceedingly rare and lead to severe neurodegenerative disease. Patients may present with spastic paraplegia, severe intellectual delay, epilepsy, and non-neurologic symptoms including skeletal dysplasia, dysmorphic features, and abnormal skin pigmentation (59,60).

Mutations in many genes within the GPI anchor biosynthetic pathway cause a variety of multiple congenital anomalies, intellectual disability, and epilepsy. The best-characterized GPI biosynthesis defect, X-linked PIGA deficiency, presents with infantile spasms with hypersarrhythmia, hypotonia, multiple brain abnormalities, and facial dysmorphisms. Patients may also have variable skin, liver, heart, and renal disease (61-69). Some mutations within PIGA cause the phenotypically distinct disease paroxysmal nocturnal hemoglobinuria (PNH), an acquired disorder of bone marrow failure (70,71).

**Diagnosis**

When a CDG is clinically suspected, the first step is to order biochemical CDG testing in plasma or serum, including CDT and N-glycan testing. Serum CDT and N-glycan analysis can only detect N-glycosylation defects, therefore they would not be useful in differentiating isolated O-glycosylation or GPI anchor defects. Transferrin isof orm analysis was originally obtained by isoelectric focusing of transferrin, as failure of N-glycan synthesis causes partial deficiency of sialic acid, which alters the charge on serum transferrin and subsequently its cathodal migration on an electrophoretic field. However, mass spectrometry-based analysis of transferrin and N-glycan have now largely replaced isoelectric focusing by identifying specific changes in oligosaccharides by mass and charge (72).
or monosialotransferrin bands, and indicates defects in N-glycan processing in the Golgi apparatus (73).

If a type I serum transferrin CDT pattern is detected, PMM2 deficiency or MPI deficiency should be at the forefront of differentials, as PMM2-CDG is the most common CDG and MPI-CDG is treatable and potentially fatal if left untreated. To differentiate between diagnoses, N-glycan profiling, molecular sequencing, or enzymatic testing should be undertaken. The diagnosis of PMM2-CDG or MPI-CDG is confirmed through molecular testing showing biallelic pathogenic variants in PMM2 or MPI, followed by PMM or MPI enzyme activity in leukocytes or fibroblasts if the pathogenicity of genetic variants are uncertain. N-glycan analysis or molecular analysis would differentiate the majority of ALG-CDG from PMM2 or MPI-CDG (15).

A type II serum transferrin CDT pattern indicates Golgi defects such as N-acetylglucosaminyltransferase (GnT) II deficiency (CDG type IIA, MGAT2-CDG). Apolipoprotein CIII (Apo-CIII) isoform analysis is a complementary test for a type II CDT profile, as it measures mucin type O-glycosylation defects in the Golgi apparatus. There is limited sensitivity for CDT or Apo-CIII in detecting type II CDG. Thus N-glycan and O-glycan profiling and molecular panel or exome sequencing should be undertaken when these clinical tests are available. Transferrin glycosylation patterns may normalize sporadically; therefore, repeated testing may be indicated in patients with high index of suspicion. False positives may be obtained in patients with acute crisis of hereditary fructose intolerance, galactosemia, acute liver disease, and some bacterial infections. None of the biochemical CDG tests can screen for all CDGs, thus even in the presence of normal screening results, molecular gene panel testing or exome sequencing may be performed for strong clinical suspicion. Conversely, biochemical and functional confirmation of molecular genetic findings are also essential as the majority of patients with CDG carry at least one mild and often novel missense mutation.

O-linked glycosylation defects and combined N- and O-linked glycosylation defects

Diagnosis relies on molecular sequencing, as transferrin isoform analysis would not detect isolated O-glycosylation defects. Combined N- and O-linked glycosylation defects can be detected by CDT, ApoCIII analysis, and plasma N-glycan and O-glycan analysis.

Lipid glycosylation and GPI anchor biosynthesis defects

Flow cytometry of blood granulocytes measures cell surface expression of GPI-anchored proteins such as CD16 and CD24. Flow cytometry analysis of white blood cells or red blood cells for certain GPI-anchored cell surface proteins are available clinically as a test for PNH due to acquired mutations in the PIGA gene. The PNH test may reveal abnormalities in other GPI anchor deficiencies, but diagnosis mostly relies on molecular analysis.

Molecular analysis

The highest diagnostic yield for CDG is next-generation based gene sequencing panel or clinical exome sequencing (CES). Genetic sequencing proofreads the nucleotide sequence, or “letter” spelling, of genes to determine whether there is a change that affects the function of the gene. The human genome consists of 3 million nucleotides but only 1–2% of these, called exons, are translated into a functional protein product. The remaining non-coding DNA interspersed between exons that do not get translated are called introns (74). CES examines almost all known exons of the approximately 20,000 genes in the human genome, which accounts for a minority of the genetic material in chromosomes but are most likely to contain disease-causing (pathogenic) variants. CES can also include mitochondrial DNA (mtDNA) sequencing, which interrogates small, extranuclear, circular DNA located in the mitochondria that is exclusively maternally-inherited.

The possible results for CES include positive, negative, and variants of unknown significance. A positive result means known disease-causing (i.e., pathogenic) variants are identified, after which the diagnosis, natural history, prognosis, recurrence risk, and treatment options can be discussed. A negative result means no detectable pathogenic variants were identified. Variants of unknown significance (a.k.a., VUS) means that although genetic changes were identified, there is not enough information about the specific genetic change to know definitively if it is disease-causing. Variations in each individual’s DNA are expected, thus having parental samples tested concurrently for comparison can assist laboratory and clinical interpretation of results. The diagnostic yield of CES is estimated at 30–35% and is increasing over time as gene discovery and knowledge about the human genome continue to progress (75-77). CES is increasingly ordered as the first-line broad
genetic testing of choice given its quick turnaround time and low relative cost for amount of genetic information analyzed. The limitations of CES include the lack of 100% sensitivity, the inability to detect certain types of genetic changes (e.g., deletions, duplications, trinucleotide repeats, deep intronic mutations, or methylation defects), and the fact that a diagnosis may not provide additional information about the disease or change management.

In the reporting of CES, incidentally-detected pathogenic variants in genes associated with well-known genetic conditions may be reported as secondary findings (78). This list of recommended diseases is curated by the American College of Medical Genetics (ACMG). The Genetic Information Nondiscrimination Act (GINA) is an important consideration when deciding whether to opt in or out of learning incidental findings (79). GINA protects individuals from the misuse of genetic information in health insurance and employment, but not life insurance. GINA protects the following genetic information: family medical history, carrier testing, prenatal genetic testing, susceptibility and predictive testing, and analysis of tumors or other assessments of gene, mutations, or chromosomal changes.

**Management**

The management of CDG depends largely upon the individual’s specific symptoms. Recurrent symptoms in patients with CDG include failure to thrive, global developmental delay, vomiting, stroke-like episodes, and skeletal abnormalities. Clinical or sub-clinical coagulopathy, endocrinopathy, hepatopathy, and cardiac defects are also commonly seen. Baseline laboratory tests to establish the extent of disease and routine monitoring are recommended, especially for PMM2-CDG. These include liver function tests, serum albumin, thyroid function tests including free T4, protein C, protein S, antithrombin III, factor IX, urinalysis, and serum gonadotropins and growth hormone.

Recommended imaging includes echocardiogram, renal ultrasound, bone age, ophthalmologic examination for evaluation of lens, retina, ocular mobility, and intraocular pressure. Unless otherwise indicated, routine vaccinations are recommended for adults and children affected with CDG. Antibody titers should be obtained after vaccination as patients may have a suboptimal immunogenic response. Prophylactic repletion of coagulation factors prior to any surgical procedures may be necessary if deficiencies exist at baseline.

Clinical genetics evaluation should be undertaken to discuss the hereditary aspects of CDG, as well as establishing a medical home for these complex patients. The medical home is typically the biochemical genetics service, although genetics, neurodevelopment, or neurology departments have also served in this capacity if a dedicated biochemical genetics service is unavailable. Specialist referral for gastroenterology, hematology, endocrinology, nutrition support, speech, occupational, physical, and feeding therapies, orthopedics, and rehabilitation medicine are often necessary.

**Targeted therapies and prognosis**

Treatment for the majority of CDG types is largely supportive, with a few exceptions. MPI-CDG is the most effectively treatable of all CDG. Oral mannose is converted to mannose-6-phosphate by intracellular hexokinases, therefore bypassing the enzymatic block and producing the deficient substrate. Mannose supplementation typically starts at 1 g/kg body weight per day, divided into 4–6 doses a day. While the potentially life-threatening protein-wasting enteropathy is especially responsive to mannose treatment, the liver disease in MPI-CDG may continue to progress. Clinical symptoms improve rapidly and transferrin CDT normalize over months, although liver disease may continue to progress with treatment (45,80,81).

Caution should be exercised in supplementing mannose during pregnancy, as mannose administration in pregnant hypomorphic phosphomannose isomerase mouse models resulted in embryonic lethality and blindness in their pups (82). In addition, intravenous mannose has been associated with decreased consciousness and seizures, which resolved with glucose administration (83).

The treatment for PMM2-CDG is largely supportive and based on symptomatology. However, upcoming clinical trials on mannose-1-phosphate substrate replacement therapy are currently under development.

For other CDG, various oral simple sugars have been investigated with the aim of theoretically improving hypoglycosylation. Fucose has been tried for SLC35C1-CDG and galactose for PGM1-CDG and SLC35A2-CDG with mixed results (84). D-galactose at 1.0–2.5 g/kg/day (max 50 grams) has been demonstrated to improve the hypoglycemia, coagulopathy, and endocrinopathy in PGM1-CDG (85,86). Galactose has also been shown to improve the endocrinopathy and coagulopathy in TMEM165-CDG (87) and SLC39A8-CDG. Considerable clinical improvement was also reported in SLC39A8-CDG patients.
on 15–20 mg/kg/day of MnSO$_4$ (88). Clinical trials are ongoing to investigate the utility of N-acetylmannosamine (ManNAc) in GNE-CDG (89), and several pre-clinical trials are underway for other CDG (90).

Despite medical strides, significant mortality exists for children with CDG within the first year of life from multi-organ failure or severe infection (91). Infants with CDG may present with fulminant multi-organ disease, intractable seizures, or severe hypoalbuminemia progressing to anasarca. Some patients respond to aggressive diuresis and albumin replacement while others are refractive to treatment. Sodium butyrate has been demonstrated to improve seizure control in CAD-CDG and PIGM-CDG (92). Ketogenic diet has also been shown to decrease seizure frequency in some cases of PIGA-CDG (93). During stroke-like episodes, intravenous hydration and maintenance of normal blood glucose may be helpful while underlying vascular thrombotic or bleeding etiology is ruled out.

With the advent of genome-editing techniques and better understanding of the mechanism of diseases encompassed by the diagnostic umbrella of CDG, the future of targeted therapeutic development remains promising.

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**References**


66. Krawitz PM, Murakami Y, Hecht J, et al. Mutations in...
PGO, a member of the GPI-anchor-synthesis pathway, cause hyperphosphatasia with mental retardation. Am J Hum Genet 2012;91:146-51.


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