Editorial

Congenital Disorders of Glycosylation (CDG): A Rapidly Expanding Group of Neurometabolic Disorders

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The recently delineated Congenital Disorders of Glycosylation (CDG), formerly known as Carbohydrate-Deficient Glycoprotein syndromes, are a group of inherited multisystem diseases due to defective N-glycosylation of proteins. Protein glycosylation is a frequent posttranslational modification and many secreted and membranous proteins have covalently bound carbohydrate units. These serve a variety of functions, allowing correct folding of the proteins, their protection against proteases and modulation of their biological activities [17]. The biosynthesis of N-linked oligosaccharides is elaborate and numerous glycosyltransferases are involved, attaching sugars to a growing lipid-linked oligosaccharide chain, stepwise. The donors are either nucleotide-activated or dolichol phosphate-linked sugars for these transfer reactions. Finally, the completed structure (Gluc3Man9GlcNAc2) is transposed en bloc within the endoplasmatic reticulum (ER). It is then attached to an asparagine residue of a nascent protein via the oligosaccharyltransferase complex. In the Golgi compartment, several glycosylases can remodel the oligosaccharide chain into a more complex structure by removal of mannose residues and the addition of N-acetylglucosamine, galactose, fucose and sialic acid residues [13].

Considering the ubiquitous presence of glycosylation and the multiple functions of the glycans, it is obvious that defects in the biosynthesis of N-glycoproteins will be detrimental and result in a broad spectrum of clinical signs and symptoms. In 1980, Jaeken and colleagues described the first patients retrospectively diagnosed as CDG-Ia, with multisystem involvement including developmental delay, dysmorphic features and endocrine and biochemical abnormalities [7]. From the early 80s, isoelectric focussing (IEF) of serum transferrin has been a powerful tool to search for defects of N-glycosylation. When a protein is hypoglycosylated, the number of negatively charged sialic acids of serum transferrin is diminished, which results in a cathodal shift in IEF. This test, however, cannot distinguish between different types of CDG and additional studies must be performed to reach a definitive diagnosis. In the last few years, many new patients were identified with an abnormal transferrin pattern. Up to now six different defects were delineated. Much of the recent progress was achieved by taking advantage of what is known in yeast: the early steps in N-glycosylation are highly conserved between eukaryotes, and yeast mutants have proven to be a powerful tool in the characterisation of novel types of CDG-I.

In contrast to other metabolic disorders, in which defects are restricted to one organelle, the enzyme defects in CDG are found in the cytosol, the ER and the Golgi compartment. The decreased activity of two cytosolic enzymes, phosphomannomutase 2 (in CDG-Ia) [14, 16] and phosphomannose isomerase (in CDG-Ib) [9] result in a depletion of the GDP-mannose pool, required for the biosynthesis of the lipid-linked oligosaccharide chain (LLO). Deficiencies of the ER located α-1,3 glycosyltransferase (in CDG-Ic) [1, 5, 11] and the α-1,3 mannosyltransferase (in CDG-Ib) [12] impair the assembly of the LLO. Mutations in a gene coding for a subunit of the dolichol-phosphate mannose synthase (in CDG-Ie) [6, 10] result in a shortage of the donor molecule phosphate mannose.

The new classification (Table 1) follows the recommendations of a group of experts in the field at the “First International Workshop on CDGs” (November 1999 in Leuven, Belgium). This nomenclature no longer primarily refers to the transferrin pattern in IEF, but rather divides CDG in two groups, on the basis of the localisation of the enzymatic defect in the biosynthetic pathway of N-glycans. Deficiencies which affect the assembly of the dolichylpyrophosphate-linked oligosaccharide and/or its transfer to the asparagine residues on the nascent polypeptide are classified as CDG-I. CDG-II comprises the defects in the subsequent processing of the protein-bound glycans. The latter group includes CDG-IIa which results from a reduced N-acetylglucosaminyltransferase II activity [15].

Although all disorders grouped as CDG-I result from a defect in assembly or transport of the LLO, the corresponding patients display distinctive symptoms. Most patients with CDG-I present with mild to severe psychomotor retardation and a variable non-neurological involvement with inverted nipples, abnormal fat distri...
Table 1  New classification of Congenital Disorders of Glycosylation

<table>
<thead>
<tr>
<th>Type</th>
<th>Old synonym</th>
<th>Defect and localization</th>
<th>Defective gene</th>
<th>Updated synonym</th>
</tr>
</thead>
<tbody>
<tr>
<td>Type I</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Type Ia</td>
<td>Phosphomannomutase (c)</td>
<td>Defects in the synthesis of the dolichol-pyrophosphate-linked oligosaccharide chain and/or its transfer onto the protein</td>
<td>PMM2</td>
<td>CDG-Ia</td>
</tr>
<tr>
<td>Type Ib</td>
<td>Phosphomannose Isomerase (c)</td>
<td></td>
<td>PMI</td>
<td>CDG-Ib</td>
</tr>
<tr>
<td>Type Ic/V</td>
<td>α-1, 3 Glucosyltransferase (ER)</td>
<td></td>
<td>ALG6</td>
<td>CDG-Ic</td>
</tr>
<tr>
<td>Type IV</td>
<td>α-1, 3 Mannosyltransferase (ER)</td>
<td>Dolichol-P-Man Synthase 1 (ER)</td>
<td>ALG3</td>
<td>CDG-Id</td>
</tr>
<tr>
<td>Type Ie</td>
<td>PMI</td>
<td></td>
<td>DPM 1</td>
<td>CDG-Ie</td>
</tr>
<tr>
<td>Type x</td>
<td>All defects in the synthesis of N-glycoproteins, incompletely characterized</td>
<td></td>
<td>MGAT2</td>
<td>CDG-IIa</td>
</tr>
</tbody>
</table>

c = cytosol, ER = endoplasmatic reticulum, GA = Golgi apparatus

These are exciting times for people involved in CDG. Within months, CDG-Ic, CDG-Id and CDG-Ie were added to the list of known disorders. In the near future, it seems certain that more disorders will emerge, in particular in CDG-II. The complexity of CDG-II will exceed that of CDG-I because of the complexity of the processing of the N- and O-glycan in the Golgi. Camillo Golgi (1843 – 1926) initiated basic research work on this tangled compartment. He was awarded the Nobel Prize in 1906 for his silver osmium staining technique (the ”reazione nera”) [2]. 25 years had been necessary to discover the ”apparato reticolo interno”. Now, a hundred years later, we are still far away from fully understanding this complex compartment, but progress in CDG research might give some more light and insight to these most interesting structures.

References


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17 Varki A. Biological roles of oligosaccharides: all theories are correct. Glycobiology 1993; 3: 97 – 130

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