Expression of the developmental I antigen by a cloned human cDNA encoding a member of a β-1,6-N-acetylglucosaminyltransferase gene family

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The blood group i/I antigens were the first identified alloantigens that display a dramatic change during human development. The i and I antigens are determined by linear and branched poly-N-acetyllactosaminoglycans, respectively. In human erythrocytes during embryonic development, the fetal (i) antigen is replaced by the adult (I) antigen as a result of the appearance of a β-1,6-N-acetylglucosaminyltransferase, the I-branching enzyme. Here, we report the cDNA cloning and expression of this branching enzyme that converts linear into branched poly-N-acetylglactosaminoglycans, thus introducing the I antigen in transfected cells. The cDNA sequence predicts a protein with type II membrane topology as has been found for all other mammalian glycosyltransferases cloned to date. The Chinese hamster ovary cells that stably express the isolated cDNA acquire I-branched structures as evidenced by the structural analysis of glycopeptides from these cells. Comparison of the amino acid sequence with those of other glycosyltransferases revealed that this I-branching enzyme and another β-1,6-N-acetylglucosaminyltransferase that forms a branch in O-glycans are strongly homologous in the center of their putative catalytic domains. Moreover, the genes encoding these two β-1,6-N-acetylglucosaminyltransferases were found to be located at the same locus on chromosome 9, band q21. These results indicate that the I-branching enzyme represents a member of a β-1,6-N-acetylglucosaminyltransferase gene family of which expression is controlled by developmental programs.

[Key Words: Developmental I antigen; fetal/adult erythrocytes; β-1,6-N-acetylglucosaminyltransferase; oligosaccharide branching; poly-N-acetyllactosamines; expression cloning]

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Glycoconjugates are major components of the outer surface of mammalian cells and their carbohydrate structures change dramatically during the mammalian developmental process. Specific sets of carbohydrates are characteristic for different stages of differentiation, and very often these carbohydrates are recognized by specific antibodies, thus providing differentiation antigens [Feizi 1985; Fukuda 1985]. In the mature organism, expression of distinct carbohydrates is eventually restricted to specific cell types, and aberrations in these cell-surface carbohydrates are very often associated with malignant transformation [Hakomori 1984]. The functional significance of the alterations in cell-surface carbohydrates during cell differentiation and in malignancy is not completely understood at present, although several reports suggest that these molecules are involved in the modulation of adhesive processes.

It has been generally accepted that each glycosyltransferase catalyzes only one enzymatic reaction to form a specific linkage, with one notable exception for the Lewis fucosyltransferase, which can synthesize both α1,3 and α1,4 linkages [Prieels et al. 1981; Kukowska-Latallo et al. 1990]. Such formation of a specific linkage is usually associated with the formation of specific oligosaccharides in conjunction with other glycosyltransferases. Therefore, it can be anticipated that the presence of specific oligosaccharides on the cell-surface is a result of the coordinate expression of the glycosyltransferase genes responsible for their synthesis. Although in recent years cDNAs have been obtained for approximately a dozen different glycosyltransferases [Paulson and Colley 1989; Schachter 1991; Joziasse 1992], little is known about their regulation during development and in malignancy.

The blood group i/I antigens were the first alloantigens identified that display a dramatic change during human development [Wiener et al. 1956]. The i antigen is

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expressed on erythrocytes of the fetus and neonate, whereas it is replaced by the I antigen on erythrocytes in the majority of adults (Marsh 1961). During mouse embryogenesis, it was shown that the I antigen is expressed throughout the preimplantation period, whereas the i antigen is first detected in the 5-day embryo. This expression of the i antigen is more pronounced in the primary endoderm, and the increase in the i antigen is associated with a decrease in the I antigen (Kapadia et al. 1981; Knowles et al. 1982). The determinants that define the i/I antigens have been characterized, and it has been shown that they are carried by linear and branched polylactosaminoglycans, respectively (Feizi et al. 1979; Watanabe et al. 1979). Polylactosaminoglycans are composed of repeats of N-acetyllactosamine [Galβ1 → 4GlcNAcβ1 → 3] (Fukuda 1985), and the conversion of the i into the I antigen is the result of the expression of a β-1,6-N-acetylgalactosaminyltransferase, the I-branching enzyme [IGnT (Fukuda et al. 1979; Piller et al. 1988)] for O-glycans. The expression of the latter two enzymes appears to be differentially regulated during differentiation (Piller et al. 1988) and in malignancy (Yamashita et al. 1984; Pierce and Arango 1986; Brockhausen et al. 1991; Saitoh et al. 1991; Yousefi et al. 1991). Because these β-1,6-N-acetylgalactosaminyltransferases are likely to regulate carbohydrate–protein interactions during development and in malignancy by regulating the amount of poly-N-acetyllactosamine and their terminal structures, it will be essential to determine their gene structures and to define the mechanisms for the regulation of their expression.

In this report we describe the transient expression cloning of cDNA encoding the I-branching enzyme using Chinese hamster ovary (CHO) cells that stably express the polyoma virus large T antigen (Hefner and Dennis 1991). Expression of the polyoma virus large T antigen allows the replication of a plasmid vector harboring the polyoma virus origin of replication in these CHO cells. Recently, this approach led us to obtain cDNA encoding C2GnT, the enzyme responsible for the formation of the core 2 branch [Galβ1 → 3GlcNAcβ1 → 6]GlcNAc in O-glycans (Bierhuizen and Fukuda 1992). The nucleotide and deduced amino acid sequences of the newly isolated IGnT were found to have limited but clear homology with the corresponding sequence of C2GnT. In addition, we found that the two genes encoding these two different β-1,6-N-acetylgalactosaminyltransferases are located at the same locus on chromosome 9, demonstrating a direct relationship between the two genes.

**Results**

**Expression cloning and sequence of cDNA**

It has been shown that CHO cells express the linear i antigen (Sasaki et al. 1987; Smith et al. 1990). Recently, we have established a CHO cell line (CHO–Py·leu) that stably expresses the polyoma virus large T-antigen, enabling transient expression cloning by using vectors that have the polyoma virus replication origin (Bierhuizen and Fukuda 1992). This CHO cell line neither reacted with anti-I antibodies (Step, Ma) in a panning assay nor stained with anti-I antibodies using immunofluorescence techniques. Because PA-1 human teratocarcinoma cells express a large amount of I-branched structures in their polylactosaminoglycans (Fukuda et al. 1985), a cDNA expression library from poly[A] + RNA of PA-1 cells was prepared in the mammalian expression vector pcDNA1 and screened for cDNA that directed the expression of the I antigen. After transfection, cells expressing the I antigen were enriched by panning using anti-I antibodies, and plasmid DNA was recovered from adherent cells by the Hirt procedure (Hirt 1967). After DpnI treatment to remove plasmids that were not replicated in...
transfected cells, plasmid DNA was amplified in the host bacteria and used for a second round of screening by the same procedure. After the second enrichment, the transformants prepared by the Hirt procedure were divided into small pools, and plasmid DNA was prepared again from each plate. Plasmid DNA was then transfected separately into the CHO-Py·leu cells, and the transfectants were screened by immunofluorescence using anti-I antibodies. One of the plasmid pools was selected, and subsequent rounds of sibling selection with sequentially smaller, active pools identified a single plasmid (pcDNAI-IGnT) that directed the expression of the I antigen at the cell-surface.

The cDNA insert of 1807 bp in size contains a single open reading frame in the sense orientation with respect to the pcDNAI promoter (Fig. 2). This reading frame predicts a protein of 400 amino acids in length, with a molecular mass of 45,860. Hydropathy analysis predicts that this protein has a type II transmembrane topology as has been shown for all mammalian glycosyltransferases cloned to date (Paulson and Colley 1989; Schachter 1991; Joziasse 1992). In this topology a very short cytoplasmic amino-terminal segment of 6 amino-acid residues is followed by a 19-amino-acid transmembrane domain that is flanked by basic amino acid residues. The carboxy-terminal sequence presumably consisting of stem and catalytic domains is large and most likely resides in the lumen of the Golgi complex. Because a consensus sequence for polyadenylation could not be found in the 3'-flanking sequence, it is likely that during construction of the library cDNA synthesis started at an A-rich sequence (nucleotides 1537–1547; see Fig. 2), rather than at the poly[A] tail.

This cloned cDNA hybridized to a single prominent 4.4-kb transcript in poly[A]+ RNA from PA-1 cells (Fig. 3A, lane 3), whereas it was not detected in poly[A]+ transfected cells, plasmid DNA was amplified in the host bacteria and used for a second round of screening by the same procedure. After the second enrichment, the transformants prepared by the Hirt procedure were divided into small pools, and plasmid DNA was prepared again from each plate. Plasmid DNA was then transfected separately into the CHO-Py·leu cells, and the transfectants were screened by immunofluorescence using anti-I antibodies. One of the plasmid pools was selected, and subsequent rounds of sibling selection with sequentially smaller, active pools identified a single plasmid (pcDNAI-IGnT) that directed the expression of the I antigen at the cell-surface.

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RNA isolated from CHO-Py · leu or HL-60 cells under the high stringent conditions for washings (Fig. 3A, lanes 1,2). This result is consistent with the reported presence of the I antigen in PA-1 cells [Fukuda et al. 1985] and its absence in CHO [Sasaki et al. 1987; Smith et al. 1990] and HL-60 cells [Mizoguchi et al. 1984; Lee et al. 1990].

Expression of the I antigen directed by the cloned cDNA

As shown in Figure 3B, CHO-Py · leu cells, transfected with pcDNAI-IGNT, express the I antigen recognized by human anti-I antibody [Ma]. The cells transfected with pcDNAI itself showed no staining at all. To confirm that the isolated cDNA encodes for IGNT, CHO cells were stably transfected with both pcDNAI-IGNT and pSV2neo and, as a control, with pSV2neo alone. After selection with G418, clonal cell lines were isolated by limiting dilution. The cells obtained will be referred to as CHO-neo · IGNT and CHO-neo, respectively. CHO-neo · IGNT cells, as well as the control CHO-neo cells, were metabolically labeled with [3H]galactose, and glycopeptides were prepared from the labeled cells by pronase digestion and analyzed by G-50 gel filtration.

Figure 4A demonstrates that CHO-neo · IGNT cells produced more glycopeptides with larger molecular mass than CHO-neo cells. When these glycopeptides were digested with endo-β-galactosidase [Fukuda and Matsumura 1976], the glycopeptides from CHO-neo · IGNT cells were more resistant to the enzyme treatment and yielded much less disaccharide, GlcNAcβ1 → 3Gal (Fig. 4A). This disaccharide can be produced only from a linear poly-N-acetyllactosamine chain that contains at least three N-acetyllactosamine repeats [Fukuda et al. 1978, 1979]. On the other hand, the branched galactose present in the I antigen is resistant to endo-β-galactosidase treatment [Fukuda et al. 1978]. These results are therefore in good agreement with the presence of the I antigenic structure in CHO-neo · IGNT cells. Finally, methylation analysis of the [3H]-galactose-labeled glycopeptides demonstrated the presence of galactose substituted at positions 3 and 6 [2,4-di-O-methylgalactose] in CHO-neo · IGNT cells, which was absent in the control CHO-neo cells [peak in Fig. 4B]. Taken together, these results clearly demonstrate that the CHO-neo · IGNT cells acquired the GlcNAcβ1 → 6 linkage as a result of the expression of IGNT.

IGNT and C2GnT are homologous to each other

When this newly isolated sequence was tested for homology, no significant similarity was found with any other sequences in our protein database. In addition, sequence comparison with glycosyltransferases cloned by others, including β-1,2-N-acetylglucosaminyltransferase I [Kumar et al. 1990; Sarkar et al. 1991] and β-1,4-
N-acetylglucosaminyltransferase III [Nishikawa et al. 1992], did not reveal any similarity.

We have recently cloned a cDNA encoding C2GnT [Bierhuizen and Fukuda 1992]. This enzyme adds a GlcNAcβ1→6 residue to Galβ1→3GalNAc but not to Galβ1→4GlcNAc. When the sequence of Ignt was compared with C2GnT, limited but distinct homology was found in both cDNA and deduced amino acid sequences. The amino acid sequences of the two enzymes are significantly homologous in the presumed catalytic domain [Fig. 5A]. Comparison of the cDNA sequence of the two enzymes demonstrated homology in the carboxy-terminal half of the catalytic domain as well [Fig. 5B,C]. However, the homology in the amino acid sequences of this portion is not as long as the aforementioned. The location of the homologous region with respect to the protein sequence is schematically represented in Figure 6. Taken together, these results suggest that the two β-1,6-N-acetylglucosaminyltransferases are related to each other.

The genes encoding the two β-1,6-N-acetylglucosaminyltransferases are related to each other

To further understand how these two proteins are related, chromosomal localization of these genes was determined by in situ chromosome hybridization. As shown in Figure 7, the gene encoding C2GnT was found to be localized at the chromosome 9, q21 band. Surprisingly, the gene encoding Ignt was found to be localized at the same locus (results not shown).

To further understand the relationship between Ignt and C2GnT, the genomic structures coding for the two enzymes were examined. First, the genomic sequences of the two enzymes were amplified by polymerase chain reaction [PCR] using genomic DNA as template. The 5' and 3' primers were synthesized according to the 5'- and 3'-flanking sequences of the cDNAs. The results showed that the products amplified from the genomic DNA are the same size as that expected from the amplification of the cDNA sequences (1589 bp for Ignt and 5B,C).
probes. Second, there are several genomic fragments that hybridized with one cDNA probe but not with the other IGnT, respectively (Fig. 8B). The results also reveal that hybridization using IGnT- and C2GnT-specific sequences are probes under the low stringent conditions. In these experiments, we tried to detect as many genomic sequences as probes under the low stringent conditions. In particular, a HindIII fragment of ~16 kb in size and a XbaI fragment of ~12 kb in size were detected by the IGnT probe but not by the C2GnT probe. In addition, XbaI digestion yielded at least four fragments that were detected by both probes despite the fact that there is no XbaI restriction site in either of the cDNAs. Considering that there is no intron in these sequences, these results strongly suggest that there is at least one more gene that is related to IGnT and/or C2GnT.

The results obtained suggest that IGnT and C2GnT belong to a family of β-1,6-N-acetylgalcosaminyltransferases, which probably arose by gene duplication and subsequent divergence.

**Discussion**

The present study describes the isolation of a cDNA clone encoding IGnT, the enzyme responsible for the formation of the GlcNAcβ1 → 3[GlcNAcβ1 → 6]Gal structure. The formation of this structure results in the I antigen, Galβ1 → 4GlcNAcβ1 → 3[Galβ1 → 4GlcNAcβ1 → 6]Gal [Fig. 1], because β1,4-galactosyltransferase is expressed almost ubiquitously in mammalian cells. For this cloning, a CHO cell line that stably expresses the polysoma large T antigen was used [Helfman and Dennis 1991; Bierhuizen and Fukuda 1992], allowing the replication of a plasmid vector harboring a polysoma replication origin such as pcDNAI.

There are four different β1 → 6-N-acetylgalosaminyl linkages: GlcNAcβ1 → 3[GlcNAcβ1 → 6]Gal, the IGnT product [Piller et al. 1984]; Galβ1 → 3[GlcNAcβ1 → 6]GlcNAc, the core 2 structure [Piller et al. 1988]; GlcNAcβ1 → 3[GlcNAcβ1 → 6]GlcNAc, the core 4 structure [Brockhausen et al. 1985]; and GlcNAcβ1 → 2[GlcNAcβ1 → 6]Man, the N-acetylgalcosaminyltransferase V product [Cummings et al. 1982]. The enzymes responsible for all of these linkages share the same unique property that Mn²⁺ is not required for their activity. Only the cDNAs encoding IGnT and C2GnT have been cloned so far. As described recently [Bierhuizen and Fukuda 1992], C2GnT acts exclusively on Galβ1 → 3GlcNAc and not on other substrates. Because another study suggests that there is a β-1,6-N-acetylgalcosaminyltransferase in tracheal epithelium that can form core 2, core 4, and 1 structures [Ropp et al. 1991], there must be at least one other glycosyltransferase related to IGnT and C2GnT. Southem blot hybridization of human genomic DNA with IGnT- and C2GnT-specific probes indicates that there is at least one additional sequence related to C2GnT and IGnT.

The present study demonstrates that the genes for IGnT and C2GnT are located at the same locus on chromosome 9. The present study also showed that the entire coding sequence for both enzymes is located in a single exon. In combination with the observed structural homology, these results point toward the possibility that the different β-1,6-N-acetylgalcosaminyltransferases belong to a gene family and that they probably evolved by gene duplication and subsequent divergence. There are precedents for the presence of gene families concerning glycosyltransferases. For α-1,3-galactosyltransferase, for example, it has been demonstrated that both the blood group A and B transferases [Yamamoto et al. 1990] and a human pseudogene related to bovine and murine

**Figure 8.** Separation of genomic sequences obtained by PCR (A) and Southern blot analysis of human genomic DNA using IGnT- and C2GnT-specific sequences as probes (B). The nucleotide sequences of IGnT and C2GnT were amplified by PCR, as described in Materials and methods, and separated by agarose–gel electrophoresis. The 5‘ and 3‘ primers for PCR were synthesized according to the 5‘- and 3‘-flanking sequences of eDNA encoding IGnT or C2GnT. The numbers at left denote the size of two molecular mass markers (in bp), which are similar in size to the PCR products. [Lanes 1,2] C2GnT; [lanes 3,4] IGnT. Lanes 1 and 3 were control experiments without template genomic DNA. [B] Genomic DNA prepared from HL-60 cells was digested separately with BamHI, EcoRI, HindIII, and XbaI (lanes 1–4, respectively), and aliquots of the digestions were separated in duplicate by agarose-gel electrophoresis. The mobility of molecular mass markers is indicated at left (in kb). Among the restriction enzymes employed, there is no restriction site in the coding sequence for BamHI, EcoRI, and XbaI.
Galβ1 → 4GlcNAc α-1,3-galactosyltransferase (Shaper et al. 1992) are located on human chromosome 9q33-34. Similarly, it has been shown that at least two α1,3 fucosyltransferases (FucTIII and FucCVI) are related to each other, and the genes encoding these two enzymes were found to reside on chromosome 19 (Weston et al. 1992). The coding sequences of these two fucosyltransferases are also located in one exon. If these genes are localized at the same locus on chromosome 19, it is likely that these two fucosyltransferases also evolved from a common precursor gene.

All of the glycosyltransferases cloned so far share a common type II transmembrane topology, consisting of a short amino-terminal cytoplasmic sequence, a signal-anchor sequence followed by a short stem region, and a large carboxy-terminal catalytic domain. Apart from this common topology, IGnT and C2GnT have no apparent homology with other glycosyltransferases, including two other N-acetylglucosaminyltransferases. When the amino acid sequences of IGnT and C2GnT are compared, a limited but distinct homology can be noticed. As shown in Figure 6, the region of the extensive homology is located around the center of the presumed catalytic domain. In addition, the sequence close to the carboxy-terminal region also has some homology in the two enzymes. As shown previously, this intraluminal portion contains the catalytic domain (Colley et al. 1989; Kukowska-Latallo et al. 1990; Bierhuizen and Fukuda 1992).

More recently, it has been demonstrated (Wen et al. 1992) that there is a homology in part of the amino acid sequences among three different sialyltransferases: Galβ1 → 3[4]GlcNAca-2,3, Galβ1 → 3GlcNAca-2,3, and Galβ1 → 4GlcNAca-2,6 sialyltransferases. The region of this homology lies in the center of the catalytic domains of these enzymes. The region of homology between the two β1,6-N-acetylglucosaminyltransferases (IGnT and C2GnT) lies in a similar place with respect to the domain structures of the glycosyltransferases (Fig. 6). However, the extent of the homology is much more significant between IGnT and C2GnT than between the three sialyltransferases. Because this homology is not observed with other N-acetylglucosaminyltransferases, it is unlikely that this homologous region represents the binding site for UDP-GlcNAc. In the β1,6-N-acetylglucosaminyltransferase gene family there are additional regions close to the carboxyl terminus where distinct homology exist [Fig. 6]. It is thus possible that these three homologous sequences are close to each other once the polypeptides are folded to form the three-dimensional structures. If this is the case, the conserved sequences are essential to form the correct framework, allowing specific amino acids to bind to the acceptor.

In relation to this, it is worthy to mention the structure formed by the carbohydrate recognition domain of C-type lectins. The three-dimensional structure of one of these proteins was elucidated recently (Weis et al. 1992), and the results strongly suggest that such conserved amino acid residues in the carbohydrate recognition domain are involved in generating the folding patterns that should be shared by different C-type lectins. Moreover, some of those residues are also involved in calcium or carbohydrate binding.

The availability of a cDNA clone encoding the I-branching enzyme now enables us to regulate the amount of the I branchings by regulating the transcription level of the I-branching enzyme. For example, it will be possible to express the I branchings in sialyl Leα-expressing cells and then test whether such branches increase the binding to E-selectin. It will also be possible to reduce or abolish the expression of the I-branching enzyme by anti-sense technology or gene knockout in transgenic mice. Such experiments will be important in future studies to determine whether I branching plays a critical role during embryonic development and differentiation.

It is noteworthy that the expression of all of the β1,6-N-acetylglucosaminyltransferases changes dramatically during development and oncogenesis. The occurrence of the I antigen is closely associated with development and maturation of erythroid cells [Marsh 1961; Fukuda et al. 1979], whereas the formation of the core 2 structure in O-glycans has been observed in a variety of biological processes such as T-cell activation (Piller et al. 1988) and immunodeficiency due to the Wiskott–Aldrich syndrome (Piller et al. 1991) and AIDS (Saitoh et al. 1991). On the other hand, an increase in the activities of N-acetylglucosaminyltransferase V and C2GnT has been associated with malignant transformation (Yamashita et al. 1984; Pierce and Arango 1986; Yousefi et al. 1991). The present study suggests that these enzymes, which change dramatically during development and oncogenesis, might have evolved from a common precursor gene. It is possible, therefore, that their expression is regulated by common regulatory genomic elements as well as intrinsic genomic elements. Thus, the genomic relationship of the different β1,6-N-acetylglucosaminyltransferases and their regulation in expression during development and oncogenesis will be of paramount interest in understanding the roles of carbohydrates in development and oncogenesis.

Materials and methods

Construction of stably transfected CHO cells expressing the polyoma virus large T antigen

The plasmid vectors pPSVEI-PyE, harboring the polyoma virus early genes (Muller et al. 1984), and pZIPneo-leu, harboring leukosialin (CD43) and neomycin cDNA, were constructed as described (Bierhuizen and Fukuda 1992). Polyoma large T antigen and human leukosialin-expressing cell lines were established by cotransfecting CHODG44 cells with pPSVEI-PyE and pZIPneo-leu and by subsequent selection for G418 resistance. Polyoma virus large T antigen-mediated replication of plasmids in these cell lines was assessed by measurement of the methylation status of the recombinant DNA (Heffernan and Dennis 1991) using pcDNA1 harboring cDNA-encoding galactosyltransferase (Aoki et al. 1992). One particular cloned CHO cell line, designated CHO-Py-leu, was used for transient expression cloning (Seed and Aruffo 1987).
Isolation of a human IgT cDNA clone

A cDNA library, pcDNAI-PA-1, was constructed from poly(A)⁺ RNA isolated from human PA-1 teratocarcinoma cells and the mammalian expression vector pcDNAI [Invitrogen, San Diego, CA]. This cDNA library was screened similarly as described [Bierhuizen and Fukuda 1992]. Briefly, plasmid DNA was transfected into CHO-Py·leu cells using lipofectin, and the transfected cells were detached at 37°C in PBS/5 mM EDTA (pH 7.4) after a 64-hr expression period. The detached cells were pooled, centrifuged, and resuspended in cold PBS, containing 10 mM EDTA, 10% fetal calf serum (pH 7.4), and human anti-I antibodies [Step] as serum in a 1:100 dilution. After a 1-hr incubation on ice, the cells were washed and pooled on dishes coated with goat anti-human IgM [Sigma, St. Louis, MO], prepared as described [Wysocki and Sato 1978]. Anti-I antibodies from two patients [Step and Ma] were kindly donated by Dr. Eloise Giblett [Blood Bank Center, Seattle, WA]. Plasmid DNA was rescued [Hirt 1967] from transfected CHO-Py·leu cells adherent to the panning dishes, digested with DpnI to remove plasmids that were not replicated in transfected cells, and transformed into the host Escherichia coli MC1061/P3 [Seed and Aruffo 1987]. Plasmid DNA was prepared again and used for an additional round of screening by the same procedure. E. coli transformants thus prepared from this second enrichment were plated to yield four pools of ~2000 colonies each. Plasmid DNA prepared from each plate was transfected separately into CHO-Py·leu cells, and the transfected cells were screened by panning as described above. One of the plasmid pools yielded relatively more attached and partially agglutinated cells. Transformants corresponding to this group were plated again to yield eight pools of ~500 colonies each, and replica plates were made. Plasmid DNA was prepared from the replica plates and transfected separately into CHO-Py·leu cells; transfecants were then screened for the expression of the I antigen by immunofluorescence, as described below. One of the plasmid pools was selected, and three subsequent rounds of sibling selection with sequentially smaller active pools identified a single plasmid, pcDNAI-IgT, that determined the expression of the I antigen at the cell-surface.

Immunofluorescence microscopy

Sixty-four hours after transfection, transfected cells were fixed with 0.05% p-formaldehyde in PBS and stained with human anti-I antibody [Ma or Step] as serum in a 1:100 dilution, followed by fluorescein-conjugated goat anti-human IgM [Sigma, St. Louis, MO]. The cells were then examined under a Zeiss Axioskop microscope as described previously [Williams and Fukuda 1990].

Sequencing

The cDNA insert of pcDNAI-IgT was sequenced by the chain-termination method [Sanger et al. 1977] using oligonucleotide primers synthesized according to the flanking sequences in the plasmid. The sequence was then extended by using oligonucleotide primers synthesized according to the sequences obtained within the cDNA insert.

Northern blot analysis

Poly(A)⁺ RNA prepared using a commercial kit [Stratagene, La Jolla, CA] was resolved by electrophoresis in a 1.2% agarose-2.2 M formaldehyde gel and then blotted onto a nylon membrane [Micro Separations, Inc., MA] [Sambrook et al. 1989]. The putative catalytic domain of IgT was amplified by PCR [Saiki et al. 1988], labeled with [³²P]dCTP by a random priming method [Feinberg and Vogelstein 1983], and used as a probe. Hybridizations were performed at 42°C in buffers containing 50% formamide for 24 hr, and blots were then washed several times in 0.1 x SSPE/0.1% SDS at 42°C for several hours [Sambrook et al. 1989] before exposure to Kodak XAR film at -70°C.

Southern blot analysis

Genomic DNA was prepared from HL-60 cells as described [Sambrook et al. 1989] and subjected to Southern blotting and hybridization as described previously [Siebert and Fukuda 1986]. Briefly, the blots were hybridized with cDNA probes in 6× SSPE, 0.5% SDS, 50 μg/ml of denatured, sheared salmon sperm DNA containing 50% formamide at 42°C for 16 hr, and then washed in 2× SSPE/0.5% SDS at room temperature for several hours. The probe used was identical to the one used for Northern blot analysis. Similarly, the putative catalytic domain of C2GnT was amplified by PCR as described previously [Bierhuizen and Fukuda 1992] and used as a probe after labeling by a random priming method.

Amplification of genomic DNA sequences by PCR

The genomic sequences encoding IgT and C2GnT were amplified by PCR using HL-60 genomic DNA as a template. The 5’ and 3’ primers for amplification of the IgT gene start at the sequence 154 nucleotides upstream from the initiation codon and 232 nucleotides after the stop codon, respectively. The 5’ and 3’ primers for amplification of the C2GnT gene start at the sequence 125 nucleotides upstream from the initiation codon and 141 nucleotides after the stop codon, respectively. Amplification of genomic DNA was repeated 35 times under the following conditions: denaturation for 1 min at 94°C, annealing for 2 min at 55°C, and polymerization for 5 min at 68°C. After amplification, the PCR products were subjected to 1.0% agarose gel electrophoresis.

Establishment of CHO cells stably expressing the 1-branching enzyme

CHODG44 cells were transfected either alone with pSV2neo or with pSV2neo and pcDNAI-IgT using a calcium phosphate technique [Graham and Van der Eb 1973] and subsequently selected for G418 resistance. Clonal cell lines were obtained by limiting dilution, and two cell lines, designated CHO-neo and CHO-neo·IgT, respectively, were selected.

Analysis of glycopeptides from CHO-neo and CHO-neo·IgT cells

The CHO cells were metabolically labeled with [³H]galactose (10 μCi/ml) in α-MEM supplemented with 10% fetal calf serum for 24 hr. The labeled cells were harvested with a rubber policeman, washed with PBS, and collected by centrifugation. The cell pellets were then extracted with 10 volumes of chloroform-methanol [2:1, vol/vol] as described [Fukuda et al. 1985]. The cell residues were digested with pronase for 24 hr at 60°C in a toluene atmosphere. The digest was then boiled for 10 min to denature the remaining enzyme. After centrifugation, the supernatants were applied to a column (1.0 × 110 cm) of Sephadex G-50 [Superfine] equilibrated with 0.1 M NH₄HCO₃. High-molecular-weight glycopeptides were pooled and desalted, and a portion of them was subjected to endo-β-galactosidase treatment [Fukuda and Matsumura 1976]. The digested glycopep-
tides and control glycopeptides were subjected to the same Sephadex G-50 gel filtration. Endo-β-galactosidase from Escherichia freundii was kindly provided by Dr. Michiko Fukuda of this institute.

**Methylation analysis**

The glycopeptides obtained from the [3H]galactose-labeled CHO-neo and CHO-neo IgNt cells were subjected to the same procedure as described above and were then methylated as described (Ciucanu and Kerek 1984). Nonradioactive glycopeptides prepared from fetuin (Sigma, St. Louis, MO) were added as carrier before methylation. The methylated glycopeptides were dissolved in chloroform, washed five times with water, and dried under a nitrogen stream. The methylated glycopeptides were then hydrolyzed in 3 N HCl for 3 hr at 80°C. After drying the hydrolysates under a nitrogen stream, the partially methylated galactoside residues were dissolved in a small volume of chloroform–methanol [1:1, vol/vol], applied to a silica gel G plate, and subjected to thin layer chromatography in acetone/water/ammonium hydroxide[250 : 3 : 1.5 (vol/vol/vol)] as described (Lee et al. 1990). After chromatography, the sample lanes were separated into 0.5-cm sections, and the radioactivity was determined by liquid scintillation counting.

**In situ chromosome hybridization**

In situ hybridization was carried out on chromosome preparations obtained from phytohemagglutinin-stimulated human lymphocytes cultured for 72 hr. The conditions for labeling probes, hybridization, and washing were as described previously [Nguyen et al. 1986]. After coating with nuclear track emulsion [Kodak NTB2], the slides were exposed for 19 days at 4°C and developed. To avoid any slipping of silver grains during the developing procedure, chromosome spreads were first stained with a buffered Giemsa solution and metaphases were photographed. R-banding was then performed by the fluorescence-photolysis–Giemsa method, and the metaphases were photographed before analysis. In general, 100–200 metaphase cells were examined for minimizing the statistical error caused by background staining.

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**Note**

The sequence data described in this paper have been submitted to the EMBL/GenBank data libraries under accession number Z19550.

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**Abstract:**

The mammalian genome contains a large number of genes coding for the enzymes involved in glycosylation. Some of these enzymes have been cloned and characterized, and their function and regulation are beginning to be understood. This review provides an overview of the enzymes involved in glycosylation and their role in the synthesis of oligosaccharides and gangliosides. It also discusses the regulation of glycosylation and the implications of this for understanding the development of disease.
Expression of the developmental I antigen by a cloned human cDNA encoding a member of a beta-1,6-N-acetylglucosaminyltransferase gene family.

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