Distinct glycosyltransferases synthesize E-selectin ligands in human vs. mouse leukocytes

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The binding of selectins to carbohydrate epitopes expressed on leukocytes is the first step in a multi-step cell adhesion cascade that controls the rate of leukocyte recruitment at sites of inflammation. The glycans that function as selectin-ligands are post-translationally synthesized by the serial action of Golgi resident enzymes called glycosyltransferases (glycoTs). Whereas much of our current knowledge regarding the role of glycoTs in constructing selectin-ligands comes from reconstituted biochemical investigations or murine models, tools to assess the impact of these enzymes on the human ligands are relatively underdeveloped. This is significant since the selectin-ligands, particularly those that bind E-selectin, vary between different leukocyte cell populations and they are also different in humans compared with mice. To address this shortcoming, a recent study by Buffone et al. (2013) outlines a systematic strategy to knockdown up to three glycoTs simultaneously in human leukocytes. The results suggest that the fucosyltransferases (FUTs) regulating selectin-ligand synthesis may be species-specific. In particular, they demonstrate that FUT9 plays a significant role during human, but not mouse, leukocyte-endothelial interactions. Overall, this article discusses the relative roles of the FUTs during human L-, E-, and P-selectin-ligand biosynthesis, and the potential that the knockdown strategy outlined here may assess the role of other glycoTs in human leukocytes also.

Role of Carbohydrates in the Leukocyte Adhesion Cascade

A multi-step mechanism governs leukocyte infiltration into inflamed tissue. It is initiated by the capture and subsequent rolling of circulating leukocytes on the activated vascular endothelium, followed by cell activation, firm adhesion, and trans-endothelial migration.1,2 Cell rolling is a crucial step in this cascade since it defines the contact time of leukocytes with chemokines bound to glycosaminoglycans expressed on the inflamed endothelium. This contact is essential for leukocyte activation so that the cells may firmly adhere on to the vessel wall prior to diapedesis. In this context, the selectins are a family of three calcium-dependent carbohydrate binding proteins or lectins (C-type) that facilitate leukocyte capture from flowing blood and subsequently mediate cell rolling interactions.1,2 Among the three members of the selectin family, P-selectin is expressed on activated platelets and endothelial cells, and mediates the initial leukocyte capture and rolling on the endothelium.3 E-selectin is exclusively expressed on the inflamed endothelium and it supports slower rolling interactions4,5 with ligation of this lectin also contributing to leukocyte activation. The leukocyte-borne L-selectin primarily supports secondary capture and this effectively enhances the number of leukocytes recruited at sites of inflammation.1,2
The selectins bind a variety of carbohydrate epitopes on the leukocyte cell surface, with the sialyl Lewis-X (sLe\(^\text{X}\)) glycan representing the prototypic selectin-ligand.\(^4,6,7\) This is a sialofucosylated tetrasaccharide (Neu5Ac\(\alpha\)2,3Gal\(\beta\)1,4(Fuc\(\alpha\)1,3)GlcNAc) containing an \(\alpha\)(2,3)-linked sialic acid and \(\alpha\)(1,3)-linked fucose. SLe\(^\text{X}\) and related glycans are post-translationally synthesized by a family of enzymes called glycosyltransferases (glycoTs) on either cell membrane glycoprotein or glycolipid scaffolds.\(^8\) Due to the high on- and off-rates of the selectin-carbohydrate molecular interaction, selectin-mediated leukocyte binding under hydrodynamic shear results in a transient cellular interaction which is characterized by the rapid formation and breakage of molecular bonds.\(^9\) While the high on-rate facilitates the initial leukocyte capture/recruitment or “cell tethering” event, the force dependent off-rate regulates the “cell rolling” velocity.

**Novel Tools to Study Glycosyltransferases in Human Leukocytes**

The glycoTs are Golgi resident enzymes that transfer monosaccharides from sugar-nucleotide donors to protein and lipid based acceptors.\(^10\) They constitute a family of \(-200\) enzymes or \(-1\%) of the human genome.\(^11\) Biochemical assays performed with cloned glycoTs and murine studies performed with knockouts that lack one or more of these enzymes are a major source of our current knowledge regarding the contributions of the individual glycoTs to selectin-ligand biosynthesis. The recent widespread application of RNA-interference methods provides an opportunity to determine the degree to which knowledge gained from these murine models translate to humans. Critical steps in the application of this technology to human leukocytes involves: (1) Development of quantitative methods to identify effective siRNA/shRNA that silence individual glycosyltransferases. This is important since well-characterized antibodies against most glycoTs are unavailable. (2) Stable expression of these constructs in human leukocytes that are typically hard-to-transfect.

In the manuscript by Buffone et al.,\(^12\) these challenges were overcome by developing a streamlined two-step methodology (Fig. 1). In the first step, glycoT fusion proteins containing a C-terminal enhanced green fluorescent protein (EGFP) were expressed in the trans-Golgi network (TGN) of either Chinese hamster ovary (CHO) or human embryonic kidney (HEK293T) cell lines. Lentiviral vectors carrying shRNA were transduced into HL-60 cells to create stable knockdowns. Cell function was measured under fluid shear using flow chamber substrates bearing either CHO cells expressing P-selectin, recombinant L-selectin, E-selectin expressing L-cells, or IL-1\(\beta\) stimulated HUVeCs.
FUTs Regulating E-Selectin Ligand Biosynthesis are Distinct in Human vs. Mouse Leukocytes

While many potential leukocyte E-selectin ligands have been proposed in literature, the quantitation of the relative importance of these candidate glycoconjugates remains an active area of investigation. This is in part due to missing information on the precise O- or N-glycan structure(s) on glycoproteins and also human gangliosides that bind E-selectin under physiologically relevant hydrodynamic shear conditions. In addition, except for PSGL-1, which is a relatively minor E-selectin ligand, there are no monoclonal antibodies that can individually block the contributions of other potential E-selectin ligands. For example, while some studies suggest a prominent role for the N-glycans of ESL-1 (E-selectin ligand-1) and CD44 in binding mouse E-selectin,27 others suggest that E-selectin-mediated cell adhesion is largely driven by O-linked glycans.16,28 In the latter investigations, granulocytes from mice lacking the core-2 GlcNActransferase C2GnT-I16 and also mice lacking T-synthase activity 28 exhibit dramatic loss of E-selectin binding function. Adding to this complication, the E-selectin ligands in different leukocyte populations may be distinct. For example, while CD43 is considered to be an E-selectin ligand on T-cells,29,30 ESL-1 is a more potent ligand on mouse granulocytes.27 In this regard, the expression pattern of scaffold proteins bearing the carbohydrate-ligands and also the level of cellular glycosyltransferase activities are critical factors that modulate E-selectin ligand expression.

Transgenic mouse studies where putative glycoTs responsible for the construction of this carbohydrate epitope were knocked out suggest a role for polypeptide α-GalNAcT ppGalNAcT-1,15 core-1 β1,3GalactosylT T-synthase,10 core-2 β1,6GlcNAcT C2GnT-I,16 β1,4GalactosylT β4GalT-I,17 α(2,3)sialylT ST3GalT-IV and VI,18,19 and α(1,3) fucosyltransferases (FUTs), FUT720 and FUT4,21 during the synthesis of such structures (Fig. 2). Sulfation of the peptide backbone by tyrosine sulfotransferases is also important for functional selectin ligand biosynthesis on PSGL-1.

Table 1. Effect of FUT silencing on leukocyte rolling density (% reduction)*

| Cell types** | P-selectin | L-selectin | E-selectin |
|--------------|------------|------------|------------|
| FUT4−       | −          | −          | 10%        |
| FUT7−       | 56%        | 50%        | 20%        |
| FUT9−       | 20%        | 20%        | 50%        |
| FUT4−7−     | 80%        | 80%        | 30%        |
| FUT7−9−     | 56%        | 50%        | 70%        |
| FUT4−7−9−   | N.D.       | N.D.       | 80%        |

*% quantifies the reduction in cell rolling density on L-, P-, and E-selectin. N.D., Not done; −, no significant reduction. **Specific FUTs silenced in HL-60 cells are designated with a superscript minus sign.

In human systems, a number of studies that characterize the O-glycans of PSGL-1 (P-selectin glycoprotein ligand-1) in both murine and human leukocytes.13 In human systems, the N-terminus of the glycoprotein PSGL-1 is the major selectin-ligand expressed on both murine and human leukocytes.13 In humans, this glycan resides at Threonine 57 (T57), 15 amino acids from the N-terminus of mature PSGL-1. Further, in order to enhance its accessibility to the selectins, PSGL-1 is localized at the tips of the leukocyte microvilli.14 While there is a heterogeneous distribution of O-glycans at T57, the functional selectin-ligand at this site is a non-extended sialyl Lewis-X glycan contained within a core-2 motif7 (Fig. 2).
activity are likely to be important parameters that define the E-selectin ligand in specific cell types. In addition to differences in the nature of the E-selectin ligands between different leukocyte sub-populations, even in a single cell type like granulocytes, the E-selectin ligands in humans are likely to be different from that in mice. In support of this, the treatment of mouse neutrophils with pronase results in a complete loss of leukocyte adhesion to all three selectins.31,32 In contrast, the addition of pronase to both primary human neutrophils and HL-60 cells abrogates cell binding to L- and P-selectin, but this treatment has little effect on cell adhesion to E-selectin (refs. 31 and 32 and our unpublished data). The possibility that the pronase-insensitive feature of human E-selectin ligands may be attributed to cell-surface glycosphingolipids or gangliosides has been tested by several investigators.33,34 These studies show that sialylated glycosphingolipids or gangliosides extracted from primary human neutrophils and HL-60 cells contain repeating N-acetyl-lactosamine units with internal fucosylation that support E-selectin binding under static and fluid flow conditions. Inhibition of ganglioside biosynthesis with a small molecular inhibitor of glucosylceramide synthase reduces E-selectin mediated cell rolling by ~50%.35 In further support of this species-specific difference: (1) L-selectin in human but not mouse leukocytes is reported to act as an E-selectin ligand.36 (2) ESL-1 has been shown to be a functional E-selectin ligand on murine but not human myeloid cells.37 (3) An N-linked sialofucosylated glycoform of CD44 called “Hematopoietic cell E and L-selectin ligand (HCELL)” is thought to represent a functional E/L-selectin ligand on human but not murine hematopoietic stem and progenitor cells (hHSPCs).38 (4) CD44 expressed on mature neutrophils and also some lymphocytes is thought to act as an E-selectin ligand in murine but not human cells.27 Buffone et al.12 addressed the hypothesis that differences in the relative roles of glycoT1s in human vs. mouse leukocytes may, at least partially, drive the synthesis of distinct E-selectin ligands in the two species. More specifically, the authors compared the relative roles of all three myoid α(1,3)FUTs, FUT4, FUT7, and FUT9, in the two species. Here, consistent with previous studies,21 predominantly FUT7 and secondarily FUT4 synthesized all the E-selectin ligands on mouse granulocytes. In contrast, FUT9 played an important role during human leukocyte binding to E-selectin with the relative roles of the three α(1,3)FUTs during HL-60 E-selectin ligand biosynthesis varying as FUT9 > FUT7 > FUT4 (Table 1). In these assays, the number of HL-60 interacting with E-selectin under flow decreased by 30% in dual knock-outs lacking FUT4 and FUT 7 (FUT 7-7- HL-60) while this was reduced by 70% in the FUT7-9- HL-60. Since the differentiation of myeloid precursor cells to mature human leukocytes is accompanied by a substantial increase in FUT9 expression,12,38 the contribution of FUT9 to primary human neutrophils may also be significant and this can impact human inflammatory ailments.

Conclusions and Future Directions

The framework established by Buffone et al.12 provides a streamlined strategy to examine the impact of the glycoT1s in regulating human leukocyte cell adhesion function. While the current studies focus on the α(1,3)FUTs, extensions of this approach can also be applied to study the impact of other members of the glycoT1 family. Further advancement of this approach to primary cells can also enable testing of these hypotheses in human leukocyte populations derived from peripheral blood and other sources. These efforts can help identify the molecular players and corresponding drug targets for diverse human inflammatory ailments.

Disclosure of Potential Conflicts of Interest
No potential conflicts of interest were disclosed.

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