Differential regulation of P-selectin ligand expression in naive versus memory CD4⁺ T cells: evidence for epigenetic regulation of involved glycosyltransferase genes

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Lymphocytes are targeted to inflamed sites by specific “homing” and chemokine receptors. Most of them, including ligands for P- and E-selectin, are absent from naive CD4⁺ T cells and become induced after activation and differentiation in effector/memory cells. Polared effector cells are characterized by the rapid production of distinct cytokines upon restimulation. Their cytokine memory is in part controlled by epigenetic imprinting during differentiation. Here we ask whether a similar mechanism could regulate selectin ligand expression, mediating entry into inflamed sites, notably within the skin. We report that acquisition of selectin ligands by naive but not memory CD4⁺ cells depends on progression through the Gₛ/S phase of the cell cycle—a phase susceptible to modification of the chromatin structure. Cell-cycle arrest prevented transcriptional activation of glycosyltransferases involved in the generation of selectin ligands, suggesting that progression through the cell cycle is required to unlock their genes. Artificial DNA demethylation strongly increased the frequency of selectin ligand-expressing cells, suggesting that DNA methylation keeps transferase genes inaccessible in naive T cells. Due to these findings we propose that selectin-dependent inflammation-seeking properties are imprinted by epigenetic modifications upon T-cell differentiation into effector cells. (Blood. 2004;104:3243-3248)

Introduction

Activation of naive T lymphocytes by specific antigen under appropriate conditions leads to functional differentiation into subsets of effector cells.¹ This differentiation is accompanied by changes in homing receptor expression resulting in memory subsets displaying distinct homing properties. While naive T cells recirculate between lymphoid organs, fractions of effector/memory T cells gain access to peripheral inflamed sites.²,³ In contrast to naive T cells, these effector cells express receptors for inflammatory chemokines and ligands for E- and P-selectin but lack L-selectin expression.⁴,⁵

E- and P-selectin ligands are fucosylated oligosaccharides decorating distinct carrier proteins such as P-selectin glycoprotein binding protein-1 (PSGL-1) and enable trafficking of leukocytes and T cells into inflamed areas, particularly within the skin.⁶-¹⁰ The oligosaccharide epitopes become induced upon differentiation into effector/memory cells on a fraction of cells.⁶,¹¹ A related E-selectin binding epitope involved in skin-specific homing of T cells has been termed “cutaneous lymphocyte antigen” (CLA).¹²

The generation of selectin ligands depends on the coordinated action of a set of glycosyltransferases, which determines both tissue-specific and differentiation-dependent generation of selectin-binding epitopes.⁸ Particularly in T lymphocytes, expression of fucosyltransferase (FucT) VII and core-2 glycosaminyltransferase (C2 GlcNAcT) appears to correlate with the expression of selectin ligands, and deletion of these enzymes abrogates the inducible expression on T cells.¹³,¹⁴

Secondary immune responses are more efficient than primary responses due to the presence of effector cells, which can rapidly recall their effector functions. Those cells evolved from naive T cells, which after activation by an appropriate peptide–major histocompatibility complex (peptide–MHC) underwent clonal expansion and programmed differentiation. This differentiation is associated with the commitment to produce particular effector cytokines such as interferon-γ (IFN-γ) (“Th1” cells) or interleukin-4 (IL-4) (“Th2” cells), which are recalled upon subsequent stimulation.¹⁵,¹⁶

Permanent differentiation of cell lineages is often associated with chromatin modification controlling gene expression in a heritable, clonal manner. Key mediators of epigenetic changes are posttranslational modifications of DNA itself or of proteins, which are closely attached to the DNA.¹⁷,¹⁸

Changes in the pattern of DNA methylation and histone acetylation of regulatory regions of the IL-4 and IFN-γ gene suggest that similar mechanisms are involved in the stabilization of the functional polarization of effector lymphocytes.¹⁹,²⁰ The cell cycle, providing a window for chromatin remodeling, is a major checkpoint for the acquisition of effector functions. Thus, arrest of naive T cells after T-cell receptor (TCR) engagement prevented
functional differentiation without affecting the levels of lineage-determining transcription factors. As a result, the model emerged that master transcription factors interfere with the action of specific DNA methyltransferases like DNA methyltransferase 1, which are recruited to replication forks at S phase where they establish symmetric CpG methylation of hemimethylated substrates, effectively duplicating the DNA methylation pattern on newly synthesized DNA strands.

The rapid recruitment of effector/memory cells to sites of inflammation is a prerequisite for an effective immune response. It implies that subsets of effector/memory cells retain inflammation-seeking homing properties for longer times or rapidly recall this property. We hypothesized that epigenetic mechanisms regulating gene accessibility could be involved in the fixation of such an inflammation-seeking homing pattern. Therefore, we determined whether induction of selectin ligands after activation of naive T cells is controlled by the cell cycle, and show that up-regulation and induction of the involved glycosyltransferases are indeed strictly dependent on entry into the cell cycle in naive but not memory T cells. This indicates that the respective gene loci are closed in naive T cells and become permanently accessible after priming in a part of T cells. DNA methylation appears to be involved in the control of accessibility of the genes, because expression of the selectin ligands is strongly enhanced by treatment with methylation inhibitors. These findings provide first evidence for the involvement of epigenetic mechanisms resulting in a stable imprinting of homing properties on effector T cells.

Materials and methods

Mice

DO11.10 mice (a kind gift from D.Y. Loh, Washington University School of Medicine, St Louis, MO) carrying a transgenic TCR specific for the ovalbumine (OVA)–peptide 323-339 and BALB/c mice were bred under specific pathogen-free conditions in the Bundesinstitut für Risikobewertung, Berlin, Germany. All animal experiments were performed in accordance with institutional, state, and federal guidelines.

Isolation and cell purification

CD4+ T cells were purified from pooled peripheral and mesenteric lymph nodes of DO11.10 mice either by panning using anti-CD8 (53-672), anti-CD25 (PC/6), anti–Mac-1 (M1/70), and anti-FcR II/III (2.4G2) antibodies or by direct isolation of CD4+ cells by anti-CD4–fluorescein isothiocyanate (anti-CD4-FITC) (NK1.5) and anti-FITC multisort-magnetic cell separation (MACS) beads (Miltenyi Biotec, Bergisch Gladbach, Germany) to a purity of at least 98%. Naive CD4+ CD62L+ cells were positively selected with anti-CD62L microbeads (Miltenyi Biotec) to a purity of at least 98%. Antigen-presenting cells (APCs) were prepared by depletion of CD90+ cells from spleen cells or purified CD4 negative fraction using anti-CD90 MACS microbeads (Miltenyi Biotec). APCs were irradiated (30 Gy) before culture. For cultures containing naive and memory cells, CD4+ cells were isolated from peripheral lymph nodes with anti-CD4 (L3T4) microbeads (Miltenyi Biotec).

CFSE labeling

Labeling of naive CD4+ CD62L+ cells with 5-(and 6-)carboxyfluorescein diacetate succinimidyl ester (CFSE; Molecular Probes, Eugene, OR) was performed as described. In brief, washed cells were resuspended at 1×10^6/mL in phosphate-buffered saline (PBS), CFSE (5 μM final concentration) was added, and cells were incubated for 3 minutes at room temperature. The reaction was stopped by washing with RPMI 1640 containing 10% fetal calf serum (FCS).

Cell culture

Cell culture was set up with 2×10^6 cells per milliliter in complete RPMI 1640 containing 10% FCS and 10 μM 2-mercaptoethanol (2-ME) (Life Technologies, Bethesda, MD). CD4+ CD62L+ T cells from OVA TCR transgenic mice and APCs were cultured at a ratio of 1:4 in the presence of OVA323-339 peptide (Biochemistry Department, Charité Universitätsmedizin Berlin, Germany) at 0.5 μM. Cell cultures containing total CD4+ cells were stimulated with 0.1 μM OVA323-339 peptide, supporting better survival of memory cells. Cell cultures were supplemented with recombinant murine IL-12 (R&D Systems, Minneapolis, MN) at 5 ng/mL, IFN-γ (R&D Systems) at 20 ng/mL, and neutralizing anti–IL-4 antibodies at 5.5 μg/mL (11B11) as indicated. For mRNA detection, 1×10^6/mL naive T cells were activated with plate-bound anti-CD3 (1 μg/mL) and soluble anti-CD28 (1 μg/mL).

Inhibitors

Progression of cells through the cell cycle was inhibited by addition of either 300 μM 1-mimosine (ICN, Santa Ana, CA), 2 μg/mL aphidicolin (Sigma Chemical, St Louis, MO), 200 nM paclitaxel (ICN), or mycophenolic acid at 1 μM (Sigma Chemical) to the cell culture.

To study the possible role of DNA methylation, 5-aza-2-deoxycytidine (Sigma Chemical, St Louis, MO) and 5-aza-2-deoxycytidine (Sigma Chemical) as an inhibitor of cytosine methylation, was added to the cell culture at concentrations of 5 μM for 48 hours.

Cytometric analysis and cell sorting

For cytometric analysis dead cells were excluded by staining with 4,6 diamidino-2-phenylindole (Sigma Chemical). OVA TCR transgenic T cells were identified by the clonotype-specific monoclonal antibody KJ1-26.1. P-selectin binding ligands were detected by P-selectin–human immunoglobulin G (IgG) chimeric protein (provided by Dr D. Vestweber, Max-Planck-Institut für Vaskuläre Biologie, Münster, Germany) and phycocerythrin (PE)–conjugated anti–human IgG antibody F(ab’)2 (DiaNova, Hamburg, Germany) as secondary reagent, as previously described. Naive and effector/memory cells were identified by the expression of CD45RB (clone 16A, Pharmingen, San Diego, CA). The state of activation was determined by fluorescence-conjugated monoclonal antibody to CD25 (Pharmingen). Cytometric analysis was performed using an LSR or FACSCalibur and CellQuest research software (BD Biosciences). Cell sorting was done using a MoFlo fluorescence-activated cell sorter (FACS; Cytomation, Fort Collins, CO).

**Figure 1.** Proliferation enhances P-lig expression on activated CD4+ lymphocytes. Naive CD4+ T cells from DO11.10 mice were CFSE labeled and stimulated with OVA323-339 peptide and APCs plus IL-12, IFN-γ, and aL-4. On day 3, individual generations (arrows, nos.) of CD4+ T cells were identified by anti-CD4–fluorescein isothiocyanate (anti-CD4–FITC) and anti-CD25–biotin (PK136) as secondary reagent, as previously described. Naive and effector/memory cells were identified by the expression of CD45RB (clone 16A, Pharmingen, San Diego, CA). The state of activation was determined by fluorescence-conjugated monoclonal antibody to CD25 (Pharmingen). Cytometric analysis was performed using an LSR or FACSCalibur and CellQuest research software (BD Biosciences). Cell sorting was done using a MoFlo fluorescence-activated cell sorter (FACS; Cytomation, Fort Collins, CO).
Results

Proliferation enhances P-lig expression on CD4+ lymphocytes

In vivo studies have shown that after activation of naive T cells the frequency of cells expressing P-selectin ligands (P-lig) increases with proliferation.26 We found similar results after activation of naive CD4+ cells in vitro under appropriate conditions—that is, in the presence of IL-12, IFN-γ, and anti-IL-4: among T cells that had not divided, only 5% to 15% expressed P-lig while completion of the first mitosis and further divisions significantly increased the likelihood of P-lig expression on day 3 after activation (Figure 1).

Proliferation enhances mRNA expression of FucT VII and C2 GlcNAcT

To verify if glycosyltransferases involved in the generation of selectin ligands are the target of cell-cycle–dependent regulation, we analyzed mRNA expression of FucT VII and C2 GlcNAcT. Therefore, we activated CFSE-labeled naive T cells from OVA TCR transgenic D011.10 mice for 3 days by OVA223-239 peptide and APCs in the presence of IL-12, IFN-γ, and αIL-4. Then, transgenic T cells were stained by the clonotype-specific antibody KJ1-26.1 and sorted into different generations according to their CFSE content (Figure 2A). Expression of FucT VII and C2 GlcNAcT mRNA was determined by quantitative PCR. Cells that had undergone one or more divisions showed a dramatically increased FucT VII mRNA and C2 GlcNAcT mRNA expression compared with naive T cells and, to a lesser degree, with cells from the same culture that had not divided yet (Figure 2B-C).

Arresting the cells in G0/S phase of the cell cycle abrogates P-lig induction, which is accompanied by inhibition of FucT VII and C2 GlcNAcT mRNA induction

A small but significant fraction of nondivided cells expressed P-lig; these cells might have entered but not yet completed the cell cycle. To clarify this, cell-cycle inhibitors were used to stop mitosis at different phases of the cell cycle. The presence of inhibitors during
activation strongly suppressed the induction of P-lig as shown in Figure 3A-B. Arrest in an early phase of the cell cycle (G1 phase by mycophenolic acid and G1/S phase by L-mimosine) almost completely abrogated the induction of P-lig in naive cells (2% to 4% P-lig–positive cells, Figure 3A), and arrest in later phases (S or G2/M phase) by aphidicolin or paclitaxel allowed P-lig expression at levels comparable to untreated cells that had not completed cell division (Figure 3B). Furthermore, we confirmed the absence of functional P-selectin ligands on L-mimosine–treated cells in a nonstatic adhesion assay and found that adhesion of G1/S–arrested T cells to P-selectin was reduced to only 5% (± 0.75%) of that of untreated Th1 cells (data not shown).

Induction of E-selectin ligands appears to be regulated in parallel as shown in Figure 3A: undivided T cells showed a low frequency of E-lig–positive cells, and L-mimosine treatment abrogated induction of E-lig.

Therefore, entry into the S phase seems to be indeed a critical determinant in the regulation of the relevant genes. In contrast, expression of activation markers such as CD25 is observed rapidly after activation and is not blocked by inhibition of mitosis (Figure 3C). The same applies to CD69, L-selectin down-regulation, and IL-2 secretion 1 or 2 days after stimulation.22 This confirms unimpaired activation in the inhibitor-treated cultures and the peculiar type of gene regulation involved in P-lig induction.

The impact of the cell cycle on the mRNA expression of fucosyltransferase VII (FucT VII) and core-2 glycosaminyltransferase (C2 GlcNAcT) was determined 3 days after activation of naive T cells in the presence or absence of L-mimosine. Naive T cells were activated by plate-bound anti-CD3 and soluble anti-CD28 plus IL-12, IFN-γ, and anti-IL-4 for 3 days. The relative expression of FucT VII (A) and C2 GlcNAcT (B) mRNA to housekeeping gene HPRT was determined for naive T cells and cells activated in the presence or absence of L-mimosine. The mean (± SD) from 4 individual experiments is shown.

Most memory/effecter cells can express P-selectin ligands independent of cell cycling

If imprinting by chromatin remodeling is involved in glycosyltransferase regulation, at least subfractions of memory cells primed under P-lig–permissive conditions should carry the ligand-generating genes either permanently active or in an accessible, cell cycle–independent state. To investigate this prediction, CFSE-labeled CD4+ T cells isolated from peripheral lymph nodes of DO11.10 mice were activated in the presence or absence of L-mimosine, arresting the cells in G1/S phase of the cell cycle. For identification of naive and effector/memory cells, we used a combination of CFSE labeling and CD45RB staining, which traces the proliferation-associated down-regulation of CD45RB and allows discrimination of naive and effector/memory cells for up to 3 days after activation (Figure 5A). In line with previous observations, only negligible numbers of naive, CD45RBhigh cells expressed P-lig when arrested in G1/S phase on day 3 after activation (Figure 5B). In contrast, memory cells contained about 10% P-lig–expressing cells after isolation ex vivo, and an additional fraction of 20% to 35% up-regulated the ligands independent of cell cycling. To exclude the possibility that selective survival of P-lig–positive memory/effecter cells results in enhanced frequencies of P-lig–positive cells under these conditions, we sorted P-lig–positive and P-lig–negative CD45RBhigh CD4+ cells and activated the cells as described in “Materials and methods.” Comparable survival rates of 20% and 25% at day 3, respectively, excluded a preferential enrichment of P-lig–positive compared with P-lig–negative memory T cells (data not shown).

Thus, more than 50% of memory cells able to express P-lig, according to nontreated culture, did so independent of cell cycling, indicating that they carry the P-lig–generating loci in an open conformation.

Artificial demethylation enhances induction of selectin ligands

The above data do not provide clues as to the mechanisms that stabilize accessibility of the transferase genes. Demethylation of
deoxycytidine (5-aza-2-deoxycytidine, a cytosine methylation inhibitor, enhances the induction of selectin ligands under the least permissive in vitro conditions to levels observed under optimal inductive conditions. This effect of DNA demethylation and the requirement to complete early phases of the cell cycle that are vulnerable to chromatin remodeling strongly suggest that induction of selectin ligands is epigenetically modulated in lymphocytes during differentiation. Therefore, our data indicate that methylation keeps P-lig–generating genes inaccessible in naive cells and that chromatin remodeling has to occur with primary induction of selectin ligands under permissive conditions.

**Discussion**

Expression of appropriate homing receptors on effector cells allows their accumulation at the site of infection, which, in concert with the polarized effector functions, ensures a rapid and efficient immune response during secondary infection. Here, we provide first evidence that an inflammation-seeking homing potential might be imprinted by epigenetic gene modification, a mechanism shown to be involved in the fixation of functional differentiation of effector cells.

First, we demonstrate that the cell cycle is a major checkpoint for the regulation of P-lig expression in lymphocytes. In general, proliferation enhances the frequency of cells expressing P-selectin ligands as shown here and upon in vivo activation.26 The observed frequency of memory cells expressing P-lig without the need for mitosis resembles that among CD4+ effector cells found in vivo during acute inflammation28 and most likely represents the fraction of memory cells initially primed to express P-lig. In conclusion, the findings of this study reveal a striking similarity between induction of the cytokine memory and generation of effector cell subsets displaying selective, inflammation-seeking trafficking properties. Dependency on cell cycling, a role of DNA methylation, and the capacity to recall a functionally polarized state are hallmarks of epigenetic imprinting. Thus, epigenetic modifications might provide the basis for a long-term inflammation-seeking homing pattern of effector/memory cells, ensuring a rapid secondary immune response within peripheral extralymphoid sites.

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**References**


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