L-selectin-mediated Lymphocyte-Cancer Cell Interactions under Low Fluid Shear Conditions*

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Cell migration in blood flow is mediated by engagement of specialized adhesion molecules that function under hemodynamic shear conditions, and many of the effectors of these adhesive interactions, such as the selectins and their ligands, are well defined. However, in contrast, our knowledge of the adhesion molecules operant under lymphatic fluid flow conditions is incomplete. Among human malignancies, head and neck squamous cancer cells display a marked predilection for locoregional lymph node metastasis. Based on this distinct tropism, we hypothesized that these cells express adhesion molecules that promote their binding to lymphoid tissue under lymphatic fluid shear stress. Accordingly, we investigated adhesive interactions between these and other cancer cells and the principal resident cells of lymphoid organs, lymphocytes. Parallel plate flow chamber studies under defined shear conditions, together with biochemical analyses, showed that human head and neck squamous cancer cells express heretofore unrecognized L-selectin ligand(s) that mediate binding to lymphocyte L-selectin at conspicuously low shear stress levels of 0.07–0.08 dynes/cm2, consistent with lymphatic flow. The binding of head and neck squamous cancer cells to L-selectin displays canonical biochemical features, such as requirements for sialylation, sulfation, and N-glycosylation, but displays a novel operational shear threshold differing from all other L-selectin ligands, including those expressed on colon cancer and leukemic cells (e.g. HCELL). These data define a novel class of L-selectin ligands and expand the scope of function for L-selectin within circulatory systems to now include a novel activity within shear stresses characteristic of lymphatic flow.

All cell-cell and cell-matrix interactions in nature require engagement of adhesion molecules capable of resisting prevailing forces of shear, either present within the organism or in the microenvironmental milieu. Cell lodgment and tissue residence thus require the presence of adhesion molecules capable of mediating binding within the shear stresses inherent to a particular site. Within organisms possessing circulatory systems, the shear force to be resisted in order to establish biologically relevant interactions is smallest within interstitial compartments and greatest within the hemovascular compartment. At present, our knowledge regarding the molecular basis of leukocyte hematogenous trafficking to lymph node, skin, and acute inflammatory sites is relatively robust (1, 2). Most commonly, leukocytes exit the vasculature at post-capillary venules, where shear stress ranges from 1 to 4 dynes/cm2 (1, 2). The well established “multistep” paradigm of leukocyte trafficking holds that leukocytes in flow must first make contact along the endothelial surface with adhesive interactions of sufficient strength to overcome the shear forces of blood flow. The principal effectors of the initial leukocyte adhesion to endothelium is the selectin family of adhesion molecules. The selectins (E-, P-, and L-selectins, CD62E, CD62P, and CD62L, respectively) are a family of calcium-dependent glycoproteins that are the most efficient mediators of shear-resistant interactions described to date. Each of the selectins displays optimal binding to its respective ligands under physiologic shear conditions, particularly once a shear threshold has been surpassed (3). E- and P-selectin are expressed on the vascular endothelium, and P-selectin is also expressed on platelets (4). L-selectin, however, is strictly expressed on leukocytes and is highly expressed on peripheral blood lymphocytes, particularly among naive and central memory lymphocytes (4, 5). Initial selectin-mediated interaction enables subsequent engagement of chemokine receptors and integrins that promote cellular firm adhesion, endothelial transmigration, and ultimately target site residence. It is known that similar receptor/ligand cascades, also initiated by selectin-mediated interactions, promote homing of hematopoietic stem cells to bone marrow as well as tumor cell hematogenous targeting of distant metastatic sites (2, 6–8). In contrast, however, the process of cellular trafficking within lower fluid shear stress levels such as in lymphatic compartments is poorly understood.

Tumor spread to lymph nodes is the culmination of a multistep process that includes tumor cell invasion into the lympho-
vascular compartment, tumor cell lodgment within the targeted tissue, and tumor cell growth within this new microenvironment. Recent reports have shed light on the initial steps driving the process of lymphatic metastasis (9–12) and of physiologic cellular recruitment into lymphatic vessels (13). However, the molecular basis of tumor cell lodgment within lymph nodes is uncharacterized. In head and neck squamous cell cancer (HNSCC), as opposed to most other solid tumors, spread of disease is overwhelmingly confined to regional lymph nodes, with distant metastatic disease developing only as a late feature of the most advanced cases (14). This distinctive clinical characteristic prompted us to analyze whether HNSCC cells possess unique molecular effectors that mediate adhesive interactions active within lymphovascular shear stress, such as those that may be important for tumor cell lodgment within the lymph node microenvironment. These cellular interactions occur in the setting of lymph flow and in the absence of an endothelial barrier. Thus, cells entering the lymph node via afferent lymphatic flow percolate within the nodal parenchyma, a compartment composed primarily of lymphocytes.

Measurements of shear stresses within the lymph node microenvironment have not been reported. However, measurements of flow within human lymphatic capillaries in vivo have recorded lymph median linear velocity to be 0.097 mm/s (15). Based on this value, and the similar dimensions of measured lymphatic capillaries and lymph node sinuses, mean wall physiological shear stress within the lymph node sinuses is estimated to be 0.08 dynes/cm², 10-fold lower than hematogenous shear stress levels. Within this backdrop of motion, we hypothesized that cells entering the lymph node interstitial microenvironment that express adhesion receptors specialized to function within this shear stress would have an advantage in establishing residence. Thus, we examined whether HNSCC cells are specialized to bind lymphocytes under low fluid shear. Here we show that HNSCC cells interact with L-selectin expressed on lymphocytes under conditions of shear stress, but not under static conditions. This interaction is maximal within low shear stress levels characteristic of lymph flow and is mediated by a novel class of L-selectin ligands expressed on primary HNSCC cells. This adhesive phenotype is characteristic of HNSCC but is not displayed by other solid tumor cells. Our findings unveil a previously unrecognized role for L-selectin in mediating lymphocyte-adhesive interactions under low shear stress (<1 dyne/cm²) and provide new perspectives on shear stress-based biology as may occur within the lymphatic system.

**EXPERIMENTAL PROCEDURES**

**Cells and Reagents**—Head and neck cancer cell lines JHU-SCC-011, JHU-SCC-013, and JHU-SCC-019 were a gift from Dr. James Rocco (Boston). These cell lines were developed from primary head and neck squamous cell carcinomas of the upper aerodigestive tract (16, 17). KG1a, LS174T, MCF-7, and MDA-MB-231 cells were purchased from ATCC (Manassas, VA). Cells were maintained in RPMI 1640 medium containing glutamine supplemented with 10% fetal bovine serum (FBS) under standard cell culture conditions. Human lymphocytes were purified from blood by isolation of the mononuclear cell fraction from peripheral blood (PBMCs). PBMCs were prepared by density gradient separation (Histopaque 1077, Sigma) of peripheral blood obtained from healthy donors with full consent as per an Institutional Human Subjects Internal Review Board-approved protocol. Isolated PBMC were >90% T cells (CD3+ cells) and expressed L-selectin as determined by immunofluorescent staining and flow cytometry using a Beckman-Coulter model Cytoflex FC 500 MPL (Fullerton, CA). Enzymes and chemical reagents used are as follows: α2–3-neuraminidase from *Streptococcus pneumoniae* (Calbiochem), α2–3,6,8-neuraminidase from *Vibrio cholerae* (Roche Diagnostics), O-sialylglycoprotein endopeptidase (*Cedarlane, Ontario, Canada*), heparinase II (*Sigma*), sodium chlorate (*Sigma*), tunicamycin (*Sigma*), phorbol 12-myristate 13-acetate (*Sigma*), and bromelain (*Sigma*). Recombinant human L-selectin/human Ig Fc chimera was purchased from R & D Systems. Antibodies used in the study were as follows: rat anti-human CLA, HECA-452, IgM (Pharmingen); MECA-79, rat IgM (kind gift from Dr. Phillip R. Streeter, Oregon Health Sciences University); murine anti-human PSGL-1, KPL-1, IgG1 (Pharmingen); mouse IgG1,κ isotype control (Pharmingen); rat IgM isotype control (Pharmingen); murine anti-human CD44, 2C5, IgG2a (R & D Systems, Minneapolis, MN); anti-L-selectin antibody LAM1-116, mlgG2a (Research Diagnostics Inc., Concord, MA); murine anti-human cytokeratin antibody, MAB3412, IgG1 (Chemicon, Temecula, CA); Texas red-conjugated goat anti-mouse IgG antibody (Pierce); fluorescein isothiocyanate isothiocyanate-conjugated goat anti-mouse Ig and fluorescein isothiocyanate-isothiocyanate-conjugated goat anti-rat IgM (Pharmingen); rat anti-human CD44, Hermes-1, IgG2a was a gift of Dr. Brenda Sandmaier (Fred Hutchinson Cancer Research Center; Seattle, WA). Purified and phycoerythrin-conjugated murine anti-human CD34, QBEND10, IgG1, and phycoerythrin-conjugated mouse IgG1,κ isotype control were from Coulter-Immunotech (Miami, FL). Alkaline phosphatase-conjugated anti-rat IgM and anti-mouse Ig were from Southern Biotechnology Associates, Birmingham, AL.

**Shear-dependent Binding Assays**—Cell–cell interactions were studied by growing cells in standard tissue culture-treated 6-well plates. A circular parallel plate flow chamber apparatus (GlycoTech, Gaithersburg, MD) with internal flow chamber dimensions of 2 × 0.5 × 0.025 cm was mounted over the unfixed live cell monolayer (HNSCC cells) or fixed cell monolayer (KG1a and LS174T cells) within a well and sealed using low wall suction as described previously (18, 19). The system was then equilibrated with Hanks’ balanced salt solution (Invitrogen) supplemented with 10 mM HEPES, pH 7.4, and 2 mM CaCl₂ (binding buffer). Purified lymphocytes were washed and resuspended in binding buffer at a concentration of 4.0 × 10⁶ cells/ml. Unfixed lymphocytes were drawn into the chamber under defined flow conditions by a precision syringe pump (Harvard Apparatus, Cambridge, MA), and interactions were observed in real time under stable shear force conditions using an inverted phase contrast microscope. For experiments titrat-
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ing shear stresses, the lymphocytes were first introduced into the chamber at 0.5 ml/min and then flow was momentarily suspended allowing for cells to settle, followed by an incremental increase in flow rate. Runs were recorded onto videotape using a standard CCD camera attached to a VHS recorder. Interactions were scored by review of the videotape. Primary attachments were quantified by determining the number of lymphocytes that interact with the cancer cell monolayer for more than two video frames (0.07 s) during the first 3 min of flow. Secondary interactions from cells that rolled into the field of observation from upstream regions were excluded. Rolling was defined as lateral translation of a bound cell greater than five cell diameters below the hydrodynamic velocity. Firm adhesion was defined as no lateral translation in the setting of hydrodynamic velocity and interaction resistant to shear force. Accumulation was defined as the sum of rolling and firm adhesion interactions and was scored after 4 min of flow. L-selectin specificity of interactions was established by pretreating lymphocytes with anti-L-selectin antibody LAM1-116 or anti-L-selectin polyclonal serum (50 ng/ml, on ice, 30 min) or PMA (10 ng/ml, 37 °C, 1 h).

Alternatively, unfixed cancer cells in suspension were also introduced in the flow chamber over L-selectin chimera adsorbed to the tissue culture plate. Cells were grown under standard culture conditions and harvested by incubation with EDTA (5 mM EDTA, 37 °C, 15 min) and washed with binding buffer. Cells were resuspended to a final concentration of 1.0 × 10^6 cells/ml. Chimera plates were prepared by overnight incubation with L-selectin chimera (0.5 µg/ml in PBS) at 4 °C and blocked with heat-inactivated FBS for 24 h. Attachment was assayed by introducing tumor cells into the parallel flow chamber at 0.5 ml/min (1.4 dynes/cm²) following by decreasing flow to the desired level. In experiments shown in Fig. 2A, flow was incrementally decreased every 30 s. In other experiments, flow was immediately decreased to the desired level once the cell bolus was visualized in the field. Accumulation was scored at the end of 30 s of flow at a determined shear stress. The specificity of L-selectin interaction was assessed by pretreating the cancer cells with bromelain (0.1%, 37 °C, 1 h), EDTA (10 mM EDTA, 30 min), neuraminidase (0.1 unit/ml at 37 °C, 1 h), O-sialoglycoprotein endopeptidase (24 or 240 µg/ml at 37 °C, 2 h), heparinase (0.05 or 5 units/ml, 37 °C, 4 h), sodium chlorate (10 mM, 37 °C, 20 h), or tunicamycin (15 µg/ml, 37 °C, 20 h). L-selectin specificity of interactions was established by pretreating the L-selectin chimera plates with anti-L-selectin antibody LAM1-116 or anti-L-selectin polyclonal serum (50 µg/ml) prior to the infusion of the cancer cell suspensions. Mouse isotype antibody was used as negative controls for blocking. Detachment was assayed by first establishing binding at 0.07 dynes/cm² and then incrementally raising flow rate. Adhesive interactions were scored 30 s after establishing a discrete shear stress level. Rolling velocity was calculated as the distance traveled by the centroid of an attached cell divided by the period of observation (5 s).

All experiments were performed on different days with the use of freshly prepared reagents, i.e. newly isolated lymphocytes, newly isolated cancer cell lines, or newly created L-selectin chimera spots. Wall shear stress (T) values were calculated according to the formula \( T = \frac{Qb}{a} \), where \( \mu \) is the coefficient of viscosity of the solution in the chamber (poise); \( Q \) is the volumetric flow rate (cm^3/s); \( b \) is the channel width (0.5 cm); and \( a \) is the half-channel height (0.0127 cm). A value of 0.009 poise (for water) was used for the viscosity of flow buffer at 25 °C.

**Primary Tumor-based Assays**—Fresh primary tumor specimens were dissected from the center of the tumor mass (i.e. not bordering on normal mucosa) and were minced and incubated in RPMI containing 60 µg/ml collagenase (Sigma C9697) at 37 °C for 3–6 h. Undigested tumor pieces were allowed to briefly settle to the bottom of the tube, and cells in suspension were removed with the supernatant. Cells were then pelleted at 200 × g for 10 min and resuspended in 3–6 ml of Hanks’ balanced salt solution (Invitrogen) supplemented with 10 mM HEPES, pH 7.4, and 2 mM CaCl₂ (binding buffer). Cells were examined for low shear-dependent L-selectin binding as described above using the adherent L-selectin chimera assay. Adhered human immunoglobulin (Miles Laboratories, West Haven, CT) (0.5 µg/ml in PBS) or FBS was used as negative control for binding. Cells were introduced into the parallel plate chamber at 0.07 dynes/cm² for 5 min, followed by a 1-ml wash with binding buffer introduced at 0.3 dynes/cm², and fixation with 1 ml of 4% phosphate-buffered paraformaldehyde introduced at 0.3 dynes/cm². Fixed cells were stored at 4 °C until stained. Plates containing previously fixed cells were treated with PBS, 0.5% Triton X-100 for 5 min, followed by a PBS rinse. Plates were blocked for 15 min in 1× PBS, 5% normal horse serum, 1% bovine serum albumin and then incubated with anticytokinin antibody for 30 min. Following PBS rinses, the plates were incubated with a 1:200 dilution of Texas red-conjugated goat anti-mouse antibody for 30 min at 25 °C. Following a PBS rinse, the plates were incubated with 0.4 µg/ml 4′,6-diamidino-2-phenylindole (Pierce) in PBS for 5 min and rinsed, and coverslips were mounted with Vectashield (Vector Laboratories, Burlingame, CA). Cells were examined, and digital pictures were taken using a Nikon TE2000-U microscope with Photometrics Cool SNAP EZ digital camera (Roper Scientific, Tucson, AZ) and the NIS Elements BR software (Nikon, Melville, NY).

**Selectin Ligand Expression**—The expression of CD34, PSGL-1, HECA-452, and MECA-79 was assessed by standard flow cytometry. Expression of HCELL was assessed by immunoprecipitating CD44 with the Hermes-1 antibody followed by Western blot evaluation of the immunoprecipitate with the HECA-452 antibody as described previously (19). Briefly, membrane protein lysates of the KG1a and HNSCC cell lines were prepared in 2% Nonidet P-40, Buffer A (1% SDS, 150 mM NaCl, 0.5 mM Tris, pH 10.4, 1 mM EDTA, 20 µg/ml phenylmethylsulfonyl fluoride) and incubated with immunoprecipitating antibodies or with appropriate isotype controls overnight at 4 °C. The antibody/lysat mixture was then immunoprecipitated by the addition of protein G-agarose beads (Invitrogen) followed by extensive washes with immunoprecipitating buffer. The pellet was then treated with 6 × reducing gel loading buffer, boiled, and the supernatant recovered. The samples were then resolved by standard SDS-PAGE using 7.5% Tris-HCl pre-cast gels (Bio-Rad). The gels were subsequently transferred onto polyvinylidene difluoride blot membrane (Millipore, Billerica, MA) and...
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L-selectin—In initial studies, we sought to determine whether binding was affected by treatment of lymphocytes with PMA, a lymphocyte-activating agent known to cause L-selectin shedding and concomitant activation of integrin binding activity (25–27). Treatment of lymphocytes with PMA for 30 min at 37 °C resulted in a marked decrease in L-selectin expression (<5% of treated cells were L-selectin+ by flow cytometry). Using the parallel plate chamber, adhesion of PMA-activated lymphocytes with the tumor cell monolayer at 0.07 dynes/cm² was reduced by 80% (Fig. 1B), indicating a dominant effect of L-selectin receptor-ligand interaction(s) on lymphocyte-HNSCC adhesion. We then incubated lymphocytes with blocking anti-L-selectin antibodies LAM1-116 or polyclonal anti-L-selectin serum to directly test whether L-selectin mediated shear stress-resistant interactions between lymphocytes and HNSCC cells at 0.07 dynes/cm². Antibody blockade of L-selectin using LAM1-116 or polyclonal anti-L-selectin serum reduced binding by >50% (Fig. 1B). Collectively, these data show that L-selectin is an important effector of lymphocyte binding to HNSCC cells.

To further elucidate the discrete contribution of L-selectin in capturing HNSCC cells under flow conditions, unfixed HNSCC cells were perfused at defined shear stresses over human L-selectin/Fc chimera adhered to a solid support. To assess the upper limit of shear that can support initial tethering interactions, we initiated flow at 1.4 dynes/cm² and then subsequently lowered shear stress in defined increments every 30 s. As shown in Fig. 2A, significant adhesion of HNSCC cells to immobilized L-selectin was observed at an upper limit of 0.14 dynes/cm², with maximal binding seen as shear was lowered to 0.07–0.03 dynes/cm². No additional cells bound as shear was decreased from 0.03 to 0.02 dynes/cm² (Fig. 2A), indicating that initial L-selectin-mediated capture of HNSCC is optimal at shear ~0.03 dynes/cm². Interestingly, although there was a clear threshold shear requirement to initiate tethering, the previously bound HNSCC did not detach from the L-selectin/Fc chimera at 0.02 dynes/cm², the lowest level of shear tested. In addition, HNSCC attachment was fully abrogated by treatment with EDTA as well as by proteolytic treatment of the HNSCC cells with bromelain. Altogether, these findings show that the interaction is calcium-dependent, as is required for all known L-selectin-selectin ligand interactions, and suggests that a glycoprotein(s), rather than a glycolipid(s), serves as the relevant low shear L-selectin ligand(s) on the HNSCC cells. To ensure that the observed binding activity was indeed mediated by the L-selectin moiety of the chimera, plates with adhered chimera were incubated with the blocking antibody LAM1-116 or anti-L-selectin polyclonal serum prior to the infusion of the cancer cell suspensions. Incubation with function blocking anti-L-selectin antibodies resulted in complete abrogation of binding activity. Incubation with isotype control antibody had no effect (Fig. 2B). To determine whether L-selectin ligand(s) displayed by HNSCC required sialic acid, glycosylation, or sulfation for binding to L-selectin (characteristic features of L-selectin ligands expressed on cell membranes (28)), we treated HNSCC with α2–3-neuraminidase or α2–3,6,8-neuraminidase, O-sialoglycoprotein endopeptidase, tunicamycin, or sodium chloride, respectively. Treatment with sialidases, tunicamycin, or sodium chloride led to a marked reduction in L-selectin chimera binding at 0.07 dynes/cm² (Fig. 2C). Treatment with
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O-sialoglycoprotein endopeptidase or heparinase did not alter binding activity. These results suggest the novel ligand(s) here reported require sialylation, sulfation, and N-linked glycosylation for maximal activity.

L-selectin-Selectin Ligand Interactions at Low Fluid Shear Display Elements of Rolling and Firm Adhesion—Our initial real time video characterization of this novel, low shear-based L-selectin binding activity suggested the interaction combined elements of rolling and firm adhesion. We thus further characterized the dynamics of binding by measuring interval displacement velocities exhibited by the JHU-SCC-019 cells interacting with adhered L-selectin chimera at 0.07 dynes/cm². Interval velocity was calculated for individual cells by measuring the distance traveled by each cell within uniform successive intervals of 0.64 s. These measurements revealed a remarkably reproducible exponential decrease in velocity that asymptotically approached 0 μm/sec or firm adhesion (Fig. 3A).

We further evaluated the shear resistance of the established L-selectin binding interaction by performing a detachment assay. In this assay, cancer cells were first introduced under low shear stress (0.07 dynes/cm²) over immobilized L-selectin chimera and allowed to bind. Shear stress was then incrementally increased while scoring the percentage of initial cells that remained bound at the end of each shear interval. These experiments revealed that, once the L-selectin interaction with the cancer cell was established, it was resistant to shear stresses as high as 1.4 dynes/cm², 30-fold higher shear stress than required to establish the interaction (Fig. 3B). Notably, the cells displayed no forward displacement at the higher shear stresses, confirming firm adhesion. In contrast, KG1a cells and LS174T cells continuously rolled over adherent L-selectin chimera at 1 dyne/cm², with an average rolling velocity of 18.0 μm/s (S.E. = 1.2 μm/s) and 30.8 μm/s (S.E. = 3.2 μm/s), respectively. The L-selectin-HNSCC cancer cell interaction thus displays elements of rolling and, once established, firm adhesion (Fig. 3, A and B).

HNSCC Cells Express Novel L-selectin Glycoprotein Ligands Active under Conditions of Low Fluid Shear—By flow cytometry analysis, HNSCC cells displayed no HECA-452-reactive determinants (Fig. 4A). We evaluated HNSCC cells for expression of the three known human L-selectin glycoprotein ligands, PSGL-1, CD34, and a specialized sialofucosylated glycoform of CD44 known as HCELL (19, 29, 30). Flow cytometry and Western blot analysis showed that although the HNSCC cells express CD44, they lack HCELL, and they do not express PSGL-1 or CD34 (Fig. 4). Given the requirement for ligand sulfation for L-selectin binding (Fig. 2C), we examined HNSCC cells for the presence of sulfation-dependent MECA-79 antigens (28). By flow cytometry, a subpopulation of cells displayed MECA79 reactivity (see Fig. 4B). However, preincubating HNSCC cells with MECA-79 (at up to 50 μg/ml concentration) did not block binding of HNSCC cells to adherent L-selectin chimera at 0.07 dynes/cm², whereas it did block lymphocyte binding to murine high endo-

![Image](49x80)
L-selectin-mediated Binding at Low Shear Stress Is a Distinctive Feature of HNSCC Cells

We next sought to determine whether L-selectin-mediated adherence of lymphocytes under low shear stress was a distinct feature of HNSCC cells compared with other solid malignancies. To this end, we examined JHU-SCC-013 (HNSCC), JHU-SCC-019 (HNSCC), LS174T (colon), MCF-7 (breast), and MDA-MB-231 (breast) cells for low shear L-selectin binding using the chimera-based assay. The HNSCC cells displayed significantly higher binding activity than any of the other cell lines at 0.07 dyne/cm² (Fig. 5A). At 1 dyne/cm², only LS174T cells adhered to L-selectin chimera as distinctly rolling interactions without firm adhesion (Fig. 5B).

To determine whether the low shear stress-dependent L-selectin ligand activity observed on HNSCC cell lines is a phenotypic feature of cells within primary HNSCC tumor cells, we isolated cells from seven freshly resected native HNSCC tumor specimens for analysis. The cell suspension, derived from the center of the tumor mass, was introduced in flow over adherent L-selectin chimera surface that were not interacting to measured hydrodynamic velocity. Once initial interaction was established between a cancer cell and the adhered L-selectin chimera (tethering), cells exhibited rolling as well as transition into firm adhesion. Measurements for the noninteracting cells were made over shorter time periods given the rapid transit of these free-flowing cells through the field of view.

**FIGURE 2. HNSCC cells in flow express novel L-selectin ligands.** A, JHU-SCC-019 cells were introduced in flow at 0.5 ml/min into the parallel plate flow chamber containing adhered L-selectin/Fc chimera. Flow was then decreased incrementally to the desired levels as indicated in the figure. Cells in flow interact with adsorbed L-selectin (L-sel) chimera at low shear stress. The interaction is sensitive to bromelain treatment of cells and to chelation of divalent cations with EDTA. No chimera control confirms specificity of the assay. * represents statistically significant difference from EDTA-treated, bromelain-treated, and no chimera controls. B, interaction between JHU-SCC-019 cells and the L-selectin/Fc chimera is blocked by pretreating the solid surface containing the adherent chimera with blocking anti-L-selectin antibodies. * represents statistically significant difference from isotype control. C, JHU-SCC-019 cell binding to L-selectin chimera is sensitive to treatment with broad neuraminidase (α2–3,6,8) and α2–3-specific neuraminidase, as well as to treatment with sodium chlorate and tunicamycin suggesting the presence of important sialylated, sulfated, and N-linked glycosylated determinants. Error bars represent the mean ± S.E. * represents statistically significant difference from untreated. Differences between sialidase treatments, sodium chlorate treatment, and tunicamycin treatment are not statistically significant. OSGE, O-sialoglycoprotein endopeptidase.
dissect cancer cells from other non-cancer cells. Cytokeratin-positive cells from all seven primary tumors tested adhered to L-selectin chimera at 0.07 dynes/cm² (Fig. 6). Binding was specific for L-selectin, as cytokeratin-positive tumor cells did not bind under flow conditions to surface treated with fetal bovine serum (FBS control) or to surface treated with human immunoglobulin (Ig control). Under flow, a variable number of cytokeratin-negative cells bound to adhered chimera, but such cells also bound to human Ig-treated surfaces, suggesting these interactions may be mediated by immunoglobulin receptors. No binding of cytokeratin-negative cells was observed on surface treated with FBS.

DISCUSSION

The biology of selectin-selectin ligand interactions has been conventionally analyzed in the setting of hemodynamic fluid shear, as these molecules are important effectors of leukocyte homing in blood flow. In this context, selectins are primary mediators of a multireceptor cascade consisting of “tethering” contact of flowing cells onto the endothelium (selectins), cell rolling on endothelium (selectins), activation (chemokine receptors), and firm adhesion (integrins) followed by trans-endothelial migration of the engaged cells into the target interstitial space. Tethering is prerequisite for rolling, and, in general, rolling precedes firm adhesion. Rolling is represented by the rate of forward displacement of a cell stably attached to a substrate, measured as rolling velocity, below the hydrodynamic velocity of the fluid stream near the wall, which is estimated by measuring the velocity of cells in flow that fail to engage in adhesive interaction (3). In contrast, firm adhesion represents stable interaction without forward displacement. Notably for L-selectin, engagement to its respective ligand(s) is critically dependent on a threshold level of shear, below which the adhesive interaction is not stable (3). Prior studies showed that the shear stress range for L-selectin receptor-ligand interactions observed in vitro is equivalent to that found physiologically in the vasculature at post-capillary venules (3, 32–34); these studies have shown that L-selectin receptor-ligand interactions strictly function in pro-

FIGURE 4. HNSCC cells express novel L-selectin ligands. A, expression of the three known L-selectin glycoprotein ligands, CD34, CD44 (HCELL), and PSGL1, as well as the carbohydrate determinants HECA-452 and MECA-79, was assessed on unfixed, native JHU-SCC-019 cells using flow cytometry. Staining with isotype antibody was used as negative control. No expression of CD34 or PSGL1 was detected; however, CD44 was found to be highly expressed. HECA-452 was not expressed, whereas MECA-79-reactivity was observed on a subpopulation of cells. B, to examine whether the specific sialofucosylated variant of CD44, HCELL, was expressed, CD44 was immunoprecipitated (Ippt.) from protein extracts made form JHU-SCC-019 cells and subjected to standard SDS-PAGE/Western blotting. The blot was then probed with the HECA-452 antibody that defines the sialofucosylated epitope characteristic of HCELL. No expression was detected on JHU-SCC-019 cells, whereas parallel experiments using KG1a cells, known to express HCELL, demonstrated robust expression.

FIGURE 5. Shear-dependent L-selectin ligand activity varies by type of solid tumor. A, unfixed head and neck cancer, colon cancer, and breast cancer cell lines were assessed for low shear-dependent L-selectin binding by introducing the cancer cells in flow over adhered L-selectin/Fc chimera at 0.07 dynes/cm². Head and neck cancer cells exhibited greater L-selectin ligand activity under conditions of lymphodynamic fluid shear. Error bars represent the mean ± S.E. * represents statistically significant difference from HNSCC cells. B, the same cell lines were examined for L-selectin ligand activity at 1 dyne/cm². Only LS174T cells (colon) bound to adhered L-selectin chimera (rolling). Error bars represent the mean ± S.E. * represents statistically significant difference from LS174T cells.
Low shear stress-dependent L-selectin ligand activity is expressed on primary head and neck cancer cells. Primary cancer cells were obtained from seven tumors collected at the time of resection and introduced in flow over adherent L-selectin/Fc chimera at 0.07 dynes/cm². Once binding was established, interacting cells were fixed in situ by infusion of 4% paraformaldehyde. Fixed cells were stained with a cyto-keratin antibody to identify tumor cells. Nuclei were counterstained with 4',6-diamidino-2-phenylindole. A, all primary tumors (i–vii) examined displayed L-selectin binding activity. Error bars represent the mean ± S.E. B, representative image of bound and fixed JHU-SCC-019 cells visualized by immunofluorescent staining for cytokeratin. C and D, representative picture of bound and fixed primary tumor cells (tumor 1) visualized by immunofluorescent staining for cytokeratin. Scale bars represent 100 μm.
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therapeutic modalities for a cancer whose survival statistics have remained unchanged for several decades.

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