Cytoplasmic anchorage of L-selectin controls leukocyte capture and rolling by increasing the mechanical stability of the selectin tether

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L-selectin is a leukocyte lectin that mediates leukocyte capture and rolling in the vasculature. The cytoplasmic domain of L-selectin has been shown to regulate leukocyte rolling. In this study, the regulatory mechanisms by which this domain controls L-selectin adhesiveness were investigated. We report that an L-selectin mutant generated by truncation of the COOH-terminal 11 residues of L-selectin tail, which impairs association with the cytoskeletal protein α-actinin, could capture leukocytes to glycoprotein L-selectin ligands under physiological shear flow. However, the conversion of initial tethers into rolling was impaired by this partial tail truncation, and was completely abolished by a further four-residue truncation of the L-selectin tail. Physical anchorage of both cell-free tail-truncated mutants within a substrate fully rescued their adhesive deficiencies. Microkinetic analysis of full-length and truncated L-selectin-mediated rolling at millisecond temporal resolution suggests that the lifetime of unstressed L-selectin tethers is unaffected by cytoplasmic tail truncation. However, cytoskeletal anchorage of L-selectin stabilizes the selectin tether by reducing the sensitivity of its dissociation rate to increasing shear forces. Low force sensitivity (reactive compliance) of tether lifetime is crucial for selectins to mediate leukocyte rolling under physiological shear stresses. This is the first demonstration that reduced reactive compliance of L-selectin tethers is regulated by cytoskeletal anchorage, in addition to intrinsic mechanical properties of the selectin–carbohydrate bond.

Introduction

L-selectin is a leukocyte member of the three-member selectin family which mediates initial capture (tethering) to and rolling of circulating leukocytes on high endothelial venules (HEVs)* of peripheral lymph nodes and inflamed endothelia (Rosen and Bertozzi, 1994; Kansas, 1996). These adhesive interactions, although reversible, are required for subsequent activation of the tethered leukocyte, which result in firm vascular adhesion (Springer, 1994). Binding of endothelial selectins (P- and E-selectin) and the leukocyte selectin, L-selectin to cell-surface carbohydrate ligands involves a unique tolerance of bonds to applied shear forces (Alon et al., 1995a; Chen et al., 1997). Ligand recognition by selectins is mediated by their lectin domains, yet the translation of ligand binding into productive adhesive interactions under physiological conditions of shear flow is tightly regulated by both extracellular and cytoplasmic selectin domains (Kansas et al., 1993; Patel et al., 1995; Setiadi et al., 1998; Dwir et al., 2000). The cytoplasmic domains of many adhesion molecules regulate their affinity and adhesive activity through interactions with the cytoskeleton or inner membrane-associated proteins (Angres et al., 1996; Goldstein and Wofsy, 1996; Kucik et al., 1996; Yauch et al., 1997; Setiadi et al., 1998). The cytoplasmic tails of the three selectins are structurally distinct although highly conserved among species for any given selectin (Kansas, 1996), suggesting specialized functions for this domain. L-selectin is constitutively associated with the actin-binding protein, α-actinin (Pavalko et al., 1995), as well as with the regulatory cytoplasmic protein calmodulin (Kahn et al., 1998). Truncation of the COOH-terminal 11 residues of the cytoplasmic tail of L-selectin (see Fig. 1 A) or disruption of the actin cytoskeleton were shown to disrupt L-selectin association with α-actinin and abolish L-selectin rolling on inflamed venules under physiological shear flow, without altering carbohydrate recognition by the selectin (Kansas et al., 1993), or microvillar localization of...
L-selectin (Pavalko et al., 1995). These early data suggested that the L-selectin tail controls L-selectin function through an α-actinin–facilitated association with the actin cytoskeleton.

At least three mechanisms could account for the role of the cytoplasmic tail in L-selectin–mediated adhesion under physiological shear flow. First, the tail may prevent L-selectin from being uprooted from the membrane by its ligand before a functional tether has formed. Second, cytoskeletal association of L-selectin could modulate L-selectin clustering and facilitate formation of multivalent tethers. Finally, cytoskeletal association could directly stabilize the L-selectin tether bond. To differentiate between these possibilities, we have taken a quantitative approach to analyze at high temporal resolution the kinetic properties of cell capture events and rolling motions mediated by full-length L-selectin and two cytoplasmic tail truncation mutants on purified L-selectin ligands. This approach provides further evidence for the functional importance of association of L-selectin with the actin cytoskeleton mediated by its cytoplasmic domain. Tail truncation increased the sensitivity of the L-selectin tether \( k_{\text{off}} \) to increased shear force, and thereby dramatically lowered the lifetime of tethers forming at physiological shear stresses and restricted leukocyte capture and rolling. Cytoskeletal association of L-selectin appears therefore critical for millisecond stabilization of L-selectin bonds at reversible adhesive contacts.

**Results**

**A tail-truncated L-selectin mutant supports reduced rolling and partial tethering to L-selectin ligands**

To delineate how the cytoskeletal interactions of L-selectin control its adhesiveness, the ability of full-length and tail-truncated (\( \Delta \text{cyto} \)) L-selectin transfectants to interact with purified L-selectin glycoprotein ligands was examined under shear flow. Full-length or truncated L-selectin (\( \Delta \text{cyto} \), lacking the COOH-terminal 11 residues of the cytoplasmic domain) were stably expressed in the murine pre-B-lymphocyte line 300.19 (Kansas et al., 1993) (Fig. 1 A). Clones of L-selectin and \( \Delta \text{cyto} \) expressed at comparable surface levels (Fig. 1 B) were perfused under low physiological shear flow on adhesive substrates coated with glycoprotein cell adhesion molecule 1 (GlyCAM-1) or with peripheral node addressin (PNAd), the major HEV L-selectin ligands (Berg et al., 1991). Although previously shown to lack significant adhesiveness in vitro (HEV frozen section assay or in rolling through rat mesentery venules, \( \Delta \text{cyto} \)-expressing cells established persistent rolling on high density GlyCAM-1 in vivo (Fig. 2 A). Furthermore, the ability of \( \Delta \text{cyto} \)-transfected cells to initiate tethers to GlyCAM-1 (i.e., either transient or followed by rolling) was not impaired even at low GlyCAM-1 densities (Fig. 2 A). However, the ability of the \( \Delta \text{cyto} \) to convert initial cell tethering to persistent rolling was markedly reduced compared with L-selectin, concomitant with faster rolling, especially at lower GlyCAM-1 densities or elevated shear stresses (Fig. 2, A and B). Furthermore, the ability of initial \( \Delta \text{cyto} \)-mediated tethers to form and convert to rolling adhesion on PNAd was markedly reduced relative to full-length L-selectin (Fig. 2, A and B). The ability of the tail-truncated mutant to support L-selectin–dependent neutrophil rolling on monolayers of substrate-bound selectin transfectants was also significantly impaired (Fig. 2 C). However, individual \( \Delta \text{cyto} \) transfectants were nearly as efficient as their full-length L-selectin counterparts in promoting singular neutrophil tethers (Fig. 2 D), albeit the duration of \( \Delta \text{cyto} \)-mediated tethers was significantly shorter than L-selectin–mediated tethers.

Notably, the ability of the \( \Delta \text{cyto} \)-expressing cells to form both transient and rolling tethers to GlyCAM-1 required an intact actin cytoskeleton. Disruption of actin microfilaments in both the full-length and \( \Delta \text{cyto} \) transfectants by cell pretreatment with cytochalasin B (CB) abolished all tethers to either GlyCAM-1 or PNAd (Fig. 2 A and unpublished data). Therefore, these results indicate that \( \Delta \text{cyto} \) retained considerable adhesive capacity, which depends on an intact actin cytoskeleton. Thus, the membrane proximal residues of the cytoplasmic domain of L-selectin (Fig. 1) mediate considerable tethering and rolling adhesions on physiological L-selectin ligands, albeit with reduced stability. Consistent with this notion, the ability of the \( \Delta \text{cyto} \) transfectant, but not of the L-selectin transfectant to sustain rolling on high density GlyCAM-1, rapidly declined at progressively increasing shear stresses (Fig. 3 A) and reduced GlyCAM-1 density (Fig. 3 B). Notably, dimerization of cell surface L-selectin by a non-blocking dimerizing mAb (Li et al., 1998) failed to augment tethering or rolling mediated by the \( \Delta \text{cyto} \) mutant (Fig. 3 B), although it augmented the capacity of full-length L-selectin to support rolling adhesion. Although the dimerizing mAb binds its antigenic determinant on both full-length and tail mutated L-selectin with similar efficiency (unpublished data), this mAb could not rescue the adhesive defects of the \( \Delta \text{cyto} \) detected on GlyCAM-1 (Fig. 3 B). Thus, the weaker adhesive activity of the tail truncated mutant could not be caused by impaired dimerization of this mutant. L-selectin occupancy during rolling also did not appear to modify in situ its adhesive activity, since L-selectin–expressing cells

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<tr>
<td>full-length L-selectin</td>
<td>356</td>
</tr>
<tr>
<td>L( \text{cyto} ) mutant</td>
<td>RRLKK</td>
</tr>
<tr>
<td>L358stop mutant</td>
<td>RRL</td>
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Figure 1. **Amino acid sequence and surface expression of full-length L-selectin and its cytoplasmic domain mutations.** (A) Amino acid sequences of the cytoplasmic domain of L-selectin and the L-selectin mutants characterized in this study. Sequences are aligned at their membrane proximal residue, R356. (B) Immunofluorescence flow cytometry of 300.19 clones stably transfected with cDNA encoding L-selectin, L\( \text{cyto} \) mutant lacking the 11 C’ residues of its cytoplasmic domain and L358stop mutant lacking the 15 C’ residues of the cytoplasmic domain. Transfectants were stained with the anti–L-selectin mAb DREG-200 (filled line) or control mAb (dotted line) followed by FITC-labeled goat anti-mouse mAb.
Carbohydrate ligand clustering rescues the adhesive defects of the LΔcyto mutant

The impaired rolling adhesion mediated by the LΔcyto-expressing cells could reflect a defect in the ability of individual LΔcyto molecules to generate sufficient avidity for ligand. In light of the failure to rescue the adhesive defects of LΔcyto-expressing cells by L-selectin dimerization, we next compared the adhesiveness of LΔcyto and L-selectin towards two nonphysiological polyvalent carbohydrate ligands for L-selectin. The first ligand, the sulfated polysaccharide fucoidin, consists of closely spaced L-selectin binding sulfated fucose repeats (Mulloy et al., 1994), and the second ligand, sialyl Lewis x (sLe x)-decorated neoglycolipid, forms clusters when immobilized on polystyrene surfaces (Galustian et al., 1997). Strikingly, the ability of LΔcyto to tether to and roll on immobilized fucoidin or on sLe x-glycolipid was indistinguishable from that of L-selectin, irrespective of the shear stress tested and even when ligands were present at subsaturating densities (Fig. 4, A–C). Disruption of actin microfilaments with CB inhibited rolling and tethering on immobilized fucoidin in both L-selectin and LΔcyto transfectants (Fig. 4 B), suggesting that the LΔcyto required cytoskeletal associations for its adhesiveness to the polysaccharide L-selectin ligand. Thus, high local valency of carbohydrate L-selectin ligand completely rescued the adhesive deficiency of the LΔcyto mutant, even under limiting ligand conditions (Fig. 4 D). Highly clustered carbohydrate ligands may therefore bind multiple LΔcyto molecules at microvillar contacts with comparable efficiency to full-length L-selectin.

Microkinetics of full-length L-selectin and LΔcyto-mediated rolling on GlyCAM-1

Displacement motions of leukocytes rolling on glycoprotein L-selectin ligands are comprised of discrete steps separated by short pauses with characteristic duration, reflective of bond stability at defined adhesive contact zones (Alon et al., 1997; Chen and Springer, 1999; Smith et al., 1999). Under conditions of ligand densities which favor the engagement of the rolling cell through singular tethers, the duration of the tether becomes progressively shorter (Chen and Springer, 1999). To gain insight into the dynamic differences between rolling tethers mediated by L-selectin or LΔcyto mutant, microkinetics of rolling motions mediated by the variants on GlyCAM-1 was next analyzed at high temporal resolution (Fig. 5 A). Pauses >0.004 s could be attributed to specific
L-selectin binding events, as they could be completely inhibited by mAb blocking, EGTA, or soluble fucoidin (Fig. 5 B and unpublished data). The duration of the vast majority of the rolling pauses mediated by L-selectin at a shear stress of 1.75 dyn/cm² could be fit into an homogenous group with a single first order dissociation rate constant of 8.9 ± 0.2 s⁻¹ for full-length L-selectin or LΔcyto mutant-mediated tethers, respectively. Data in C and D are representative of three independent experiments.
Figure 5. Microkinetic analysis of full-length L-selectin and LΔcyto mutant-mediated rolling on GlyCAM-1 reveals tail-dependent enhancement of bond formation and stabilization. (A) Characteristic scenes of digitized Quick-Time videos recorded with high speed camera of representative WT L-selectin and tail-truncated transfectants jerkily rolling on GlyCAM-1 (100 sites/μm²) at a shear stress of 1.75 dyn/cm². Cells were visualized with 40X objective. The time (in seconds) elapsed from the beginning of the recording is shown at the top of each scene. Supplemental Video 1 is available at www.jcb.org/cgi/content/full/145/DC1. (B) Instantaneous velocities of L-selectin- and LΔcyto mutant-expressing cells rolling on GlyCAM-1 (100 sites/μm²) at a shear stress of 1.75 dyn/cm². Motions of all cells were recorded by fast camera at 500 frame/s and analyzed by computerized cell tracking as described in Materials and methods. Instantaneous velocity drops were defined as rolling pauses. Motion analysis of representative full-length L-selectin– or LΔcyto mutant–expressing cells perfused alone or in the presence of EGTA is shown in the first, second, and third panels, respectively. The velocity drops in this nonadherent cell lasted 0.004 s and were considered nonspecific pauses. Motion analysis of a representative L358stop mutant–expressing cell moving on identical field in the absence or presence of EGTA is shown in the fourth and fifth panels, respectively. (C and D) Dissociation kinetics of rolling pauses mediated by L-selectin and LΔcyto mutant on GlyCAM-1 at a shear stress of 1.75 dyn/cm² (C) or at 1 dyn/cm² (D). The koff values were determined from the slope of the natural log of number of tethers plotted vs. the duration of each tether, or pause. The r values are indicated. In C and D, the data points of full-length L-selectin–mediated tethers which did not fit the first order dissociation approximation are indicated by filled symbols. Data in C and D were collected from 3-s recording segments and the kinetic results were summarized in Table I.

Table I. Microkinetics of full-length L-selectin-, LΔcyto- and L358stop-expressing 300.19 cells interacting with GlyCAM-1 100 sites/μm² at 0.5, 1, and 1.75 dyn/cm²

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<tr>
<th>Wall shear stressᵃ</th>
<th>1.75 dyn/cm²</th>
<th>1 dyn/cm²</th>
<th>0.5 dyn/cm²</th>
<th>1 dyn/cm²</th>
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<tr>
<td>Rolling fraction (%)</td>
<td>Full-length</td>
<td>LΔcyto</td>
<td>Full-length</td>
<td>LΔcyto</td>
</tr>
<tr>
<td>Tethering fraction (%)</td>
<td>ND</td>
<td>ND</td>
<td>ND</td>
<td>ND</td>
</tr>
<tr>
<td>Mean pause duration (ms)</td>
<td>20 ± 1</td>
<td>12 ± 1</td>
<td>31 ± 1</td>
<td>20 ± 1</td>
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<tr>
<td>koff (s⁻¹)</td>
<td>64.4 ± 5.4</td>
<td>138.7 ± 4.7</td>
<td>34.1 ± 2.3</td>
<td>61.1 ± 5.4</td>
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<tr>
<td>Mean step distance (μm)ᵇ</td>
<td>4 ± 0.1</td>
<td>10.2 ± 0.3</td>
<td>2.5 ± 0.1</td>
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ᵃAll measurements were performed on cell motion segments of at least 20 cells recorded at 500 frames/s.
ᵇStep distance was defined as the distance traveled by a rolling cell between successive tethers.
ᶜNot applicable, as no rolling could be detected at low shear stress.
ᵈThe 358stop mutant did not support any cell tethering to GlyCAM-1 at shear stresses >1 dyn/cm².
Tethering and rolling supported by LΔcyto are mediated by the residual cytoskeletal association of its membrane-proximal region

The susceptibility of the weak adhesive activity of LΔcyto-expressing cells to disruption of the actin cytoskeleton (Figs. 2 A and 4 B) suggested that its RRLKKG segment maintains residual cytoskeletal association sufficient to mediate leukocyte capture and weak rolling. Indeed, further truncation of the LΔcyto mutant by four additional residues, (designated L358stop) (Fig. 1 A), abolished all rolling adhesion even on high density GlyCAM-1 (Fig. 6 A). Furthermore, only a minor fraction of L358stop transfected cells could establish transient tethering to fucoidin, (Fig. 6 B), despite the fact that fucoidin completely rescued the adhesion deficiency of the LΔcyto mutant (Figs. 4 and 6 B). Moreover, the residual tethering activity of the L358stop on fucoidin was insensitive to CB treatment (Fig. 6 B), in contrast to the sensitivity of L-selectin and of LΔcyto to the cytoskeletal drug (Fig. 4 B). Microkinetic analysis of L358stop transfectant adhesion to medium density GlyCAM-1 at millisecond temporal resolution revealed that this tail mutant could not support any L-selectin specific adhesive tethers to GlyCAM-1 at a shear stresses of 1.75 dyn/cm² (Fig. 5 B). However, analysis of cell capture to GlyCAM-1 at drastically reduced shear (0.5 dyn/cm²) revealed low frequency of short-lived L-selectin specific tethers supported by the L358stop mutant (Fig. 6 C). The duration of these tethers could be fit into an homogeneous group with a single first order dissociation rate constant (koff) independent of GlyCAM-1 density (unpublished data), but threefold faster than the corresponding koff of L-selectin-mediated tethers (Fig. 6 C). Reminiscent of the tethering results observed on fucoidin (Fig. 6 B), the tethering ability of the L358stop to GlyCAM-1 was completely insensitive to CB treatment of the transfectants (tether frequency of 7.3% vs. 7.1%). Furthermore, the dissociation rate constant of L358stop-mediated tethers from GlyCAM-1 was not affected by CB treatment (koff = 44.6 s⁻¹ vs. 45.3 s⁻¹, respectively). Thus, very low residual adhesive activity of the L358stop towards GlyCAM-1 or fucoidin which did not depend on association of the mutant with the actin cytoskelon could be detected at low shear stresses, but was eliminated at higher stresses.

The response of tether koff to elevated shear forces is augmented by progressive truncation of L-selectin tail

The duration of individual transient tethers to medium density GlyCAM-1 mediated by the various mutants was next measured over a range of wall shear stresses. As observed

Figure 6. Further truncation of the LΔcyto mutant abolishes rolling and impairs tether formation and millisecond stabilization under physiological shear flow. (A) Lack of accumulation under flow of L358stop mutant–expressing 300.19 cells on high density GlyCAM-1 (400 sites/μm²). Accumulation and resistance to clearance of cells expressing full-length L-selectin and LΔcyto mutant by incremental shear stresses measured under identical conditions is shown for comparison. The assay was performed identically as in Fig. 3. The experiment shown is a representative of three independent experiments. (B) Frequency of tethering and rolling of 300.19 cells expressing L-selectin, LΔcyto mutant or L358stop mutant interacting with immobilized fucoidin at a shear stress of 1 dyn/cm². Mean values ± range of overall tether frequencies determined in several fields are shown. Mean velocities of rolling cells are shown on top of the bars. Both rolling and tethering were blocked in the presence of L-selectin blocking mAbs (unpublished data). (C) Dissociation kinetics of transient tethers mediated by L-selectin-, LΔcyto mutant– or L358stop mutant–expressing cells interacting with medium density GlyCAM-1 (100 sites/μm²) at 0.5 dyn/cm² determined from high speed camera recordings performed as in Fig. 5. Pauses ≤0.006 s were excluded from this analysis. The kinetic results are summarized in Table 1.
with L-selectin and the LΔcyto mutant, the dissociation rate constants of L358stop-mediated tethers increased with increasing shear stresses in an exponential manner (Fig. 7). This was in agreement with a relationship previously found between quantal L-selectin tether dissociation rates and applied force (Alon et al., 1997; Ramachandran et al., 1999). Strikingly, all three exponential curves established for the various tethers predict a similar koff, i.e., the koff of the unstrained tether bond in the absence of shear force for all three L-selectin variants (Fig. 7). However, the reactive compliance of L358stop tethers, i.e., the increase in koff of the stressed tether with applied shear force, was twofold higher than the reactive compliance of L-selectin tethers. As the reactive compliance exponentially decreases tether bond stability, this difference in reactive compliance was sufficient to destabilize tethers mediated by the L358stop on GlyCAM-1 to an extent they became essentially undetectable at physiological shear stresses >1 dyn/cm² (koff > 300 s⁻¹; Figs. 5 B and 7). In contrast, the reactive compliance of LΔcyto tethers was 1.4-fold higher than that of full-length L-selectin tethers (Fig. 7). This milder adhesive deficiency abrogated rolling adhesions at elevated shear forces, while cell capture ability of this mutant was still largely conserved (Figs. 2, 3, 5, and 6). Consequently, even at high shear stresses, the koff of tethers mediated by the LΔcyto mutant was only twofold faster than the koff of L-selectin tethers (Fig. 7).

**Anchorage of cell-free L-selectin tail mutants rescues their adhesive defects**

We previously described an adhesion assay to quantify the adhesive properties of cell-free L-selectin variants under shear flow (Dwir et al., 2000). In this assay, L-selectin mutants are captured in a functional state on a substrate coated with nonfunction blocking L-selectin specific mAbs, and their lectin-dependent adhesive capacities to ligands expressed on perfused neutrophils are compared under physiological shear flow. Selectin mutants differing in their intrinsic adhesiveness exhibited large adhesive differences in this assay (Dwir et al., 2000). However, when the full-length and the two tail-truncated L-selectin mutants were captured at similar densities on a substrate coated with the anti-SCR domain specific mAb, LAM1-118 (Fig. 8, or other SCR mAbs (unpublished data), essentially identical capacities to support neutrophil tethering or rolling were detected for all L-selectin mutants. Thus, anchorage of both tail truncated L-selectin mutants to a solid surface appears to have fully rescued their adhesive defects.

**Discussion**

The ability of selectin tethers to successfully support persistent rolling depends on a proper balance of tether formation and breakage at the cell front and rear, respectively. Leukocyte rolling mediated by L-selectin is supported by single microvillus tethers per adhesive contact zone lasting between 10⁻² and 10⁰ s (Alon et al., 1997; Chen and Springer, 1999). We find that L-selectin tethers shorter than 20 ms fail to support persistent rolling adhesion, suggesting that the probability of successive tether formation collapses when tether duration drops below this critical duration. Microkinetic analysis of L-selectin tethers, the building units of leukocyte rolling, mediated by structural tail variants of L-selectin on the prototypic L-selectin ligand, GlyCAM-1, provided key insights into how the cytoplasmic domain of L-selectin controls its ability to mediate leukocyte rolling. The first order dissociation kinetics of tethers formed by all structural L-selectin variants was found to approach a constant value near zero shear stress, suggesting that the koff of unstrained L-selectin tethers is independent of the cytoplasmic...
mic tail (Fig. 7). Nevertheless, the reactive compliance of tethers, i.e., the sensitivity of the tethers to shear force, was found to be considerably lower for full-length L-selectin than for its respective LΔcyto or L358stop mutants. Low compliance of L-selectin tethers to applied shear forces results in prolonged tether duration at elevated shear stresses and permits L-selectin to support both leukocyte capture and rolling under a wide range of shear stresses. In contrast, the LΔcyto, and to a greater degree the L358stop mutant, exhibit impaired abilities to support leukocyte rolling or capture. The analysis of L-selectin tether kinetics, performed here for the first time on tail mutants of L-selectin at millisecond temporal resolution, strongly suggests that translation of bonds to adhesive tethers involves millisecond stabilization events, critically regulated by the cytoplasmic tail of L-selectin and an intact cytoskeleton.

What could be the molecular basis for the impaired stabilization of tail-mutated L-selectin–mediated tethers? The exceptional ability of L-selectin to mediate cell capture under highly disruptive shear flow, even in its cell-free state (Alon et al., 1998; Dwir et al., 2000; Greenberg et al., 2000) has been attributed to intrinsically low reactive compliance of its carbohydrate bonds (Alon et al., 1997; Ramachandran et al., 1999). The present results suggest that, in addition to intrinsic molecular properties of L-selectin bonding with ligand, the association of L-selectin with the actin cytoskeleton reduces the effective reactive compliance of its cellular tethers under flow. How could cytoskeletal association of L-selectin contribute to dynamic stabilization of its tethers? Our observation that immobilization of cell-free tail-truncated L-selectin mutants fully rescues their adhesive defects, and the earlier finding that tethers mediated by cell-free L-selectin retain kinetic properties of tethers mediated by leukocyte-expressed L-selectin (Alon et al., 1998; Greenberg et al., 2000), suggests that mere anchorage of the selectin to a solid support can confer on L-selectin tethers their characteristic low reactive compliance. It is therefore possible that restriction of L-selectin’s lateral mobility within the adhesive contact zone is both required and sufficient for the L-selectin bond to undergo rapid stabilization at microvillus contact sites (von Andrian et al., 1995).

Recent measurements on single L-selectin carbohydrate bonds ruptured by a biomembrane force microsphere probe (Evans et al., 2001) reveals a \( k_{\text{off}} \) value of unstressed L-selectin: carbohydrate bond (\( k_{\text{off}} = 3 \text{ s}^{-1} \)) similar to that of unstressed tethers measured previously (\( k_{\text{off}} = 7–9 \text{ s}^{-1} \) ) (Alon et al., 1997, 1998; Ramachandran et al., 1999; Dwir et al., 2000), as well as those measured here at temporal resolution comparable to that in the biomembrane force probe experiments (Evans et al., 2001). However, under forces predicted to be applied to individual leukocyte microvilli at physiological shear stresses, i.e., > 100 pN (Alon et al., 1997), the bond between a single L-selectin and carbohydrate ligand forms and dissociates within a single millisecond period (Evans et al., 2001). Bond rupture by a force probe escaping instantaneously from the contact zone does not allow any re-binding of the dissociated receptor to ligand on the counter surface effects (Evans et al., 2001). Thus, dissociation kinetics of the receptor ligand pair studied in this system may in fact reflect the inherent reactive compliance of single L-selectin: ligand bonds dissociating in the absence of any stabilizing effects (Evans et al., 2001). Notably, single L-selectin bonds exhibit sharp response to applied force, i.e., a 1,000-fold increase in bond \( k_{\text{off}} \) between zero force and force applied on a single microvillus at a shear stress of 1 dyn/cm² (Evans et al., 2001). Remarkably, this high force sensitivity (reactive compliance) is reminiscent of the high response of tethers mediated by the L358stop mutant in our experimental system (Fig. 7). Thus, it appears that the \( k_{\text{off}} \) and the force sensitivity of the L358stop tethers are similar to the corresponding values measured for singular L-selectin bonds ruptured by force probes (Evans et al., 2001). Therefore, it is possible that once dissociated from the ligand, tail-truncated L-selectin variants are unable to instantaneously rebind it and thereby stabilize the tether at its original contact (Fig. 9 A).

We hypothesize that constitutive association of L-selectin with the actin cytoskeleton anchors the selectin within the plasma membrane, reduces its effective escape from the original ligand binding site, and thus facilitates ligand re-binding and tether stabilization. The escape radius of full-length L-selectin is predicted to be 7–8-fold smaller than that of the tailless mutant based on differences in lateral diffusion coefficients reported for anchored and unanchored proteins of similar dimensions to L-selectin (10⁻¹¹ and 5×10⁻¹⁰ cm²/s, respectively) (Chan et al., 1991; Kucik et al., 1996). The degree to which ligand re-binding by L-selectin may take place would therefore be considerably higher for L-selectin than for the L358stop mutant, with an intermediate value for the tail truncated LΔcyto mutant. Preliminary mathematical simulations of L-selectin re-binding kinetics suggests that this event takes place over a range of 10 μs (unpublished data). This mechanism of ligand re-binding would become more pronounced if the density of glycoprotein ligands for L-selectin at the microvillar site of contact is low, such that the probability of L-selectin re-binding to a neighboring ligand molecule on the adhesive surface is low (Fig. 9). Indeed, at GlyCAM-1 densities ≤ 100 sites/μm², the contact area encountered by a single tip of a microvillus contains, on average, only one GlyCAM-1 molecule (Bruehl et al., 1996). In contrast, when a highly clustered ligand like fucoidin or sLe⁻ decorated lipid serves as the leukocyte tethering ligands, this contact contains 10–100 carbohydrate sites (Alon et al., 1995b). The adhesive defects of the tail mutants could be therefore partially or fully rescued on these ligands through re-binding to a neighboring ligand molecule within the microvillus contact.

Restriction of L-selectin’s lateral mobility within its contact zone may be required but insufficient for rapid bond stabilization. Anchorage of surface proteins to the cytoskeleton has also been suggested as means to protect these molecules from being uprooted from the membrane by receptors on counter surfaces (Evans et al., 1991; Shao and Hochmuth, 1999). However, uprooting of L-selectin from the plasma membrane of neutrophils takes 0.5–1 s (Shao and Hochmuth, 1999), much more slowly than L-selectin occupancy by ligand at reversible adhesive contacts. Even after disruption of cytoskeletal associations of membrane glycoproteins, uprooting of unanchored glycoproteins from the plasma membrane takes 0.3–0.5 s (Shao and Hochmuth, 1999), two orders of magnitude longer than the duration of
Cytoplasmic regulation of L-selectin adhesion | Dwir et al. 153

tethers mediated by unanchored L358stop mutant (Fig. 6). Bond stabilization through the L-selectin tail could also conceivably involve enhanced L-selectin clustering in microvilli. However, constitutive L-selectin clustering is apparently unaffected by tail truncation (Pavalko et al., 1995). Moreover, artificially induced dimerization of L-selectin ectodomains could not rescue the adhesive defects of either the LΔcyto or L358stop mutant on any ligand system tested (Fig. 3 B and unpublished data). Thus, the adhesive defect caused by tail truncation is unlikely to be the result of impaired dimerization of the LΔcyto or L358stop mutants increased as the density of GlyCAM-1 decreased, the contribution of L-selectin dimerization to L-selectin adhesiveness in fact diminished as ligand density was reduced (Dwir et al., in preparation). Even when ligand was present at density below that sensitive to L-selectin dimerization, tail-dependent association of L-selectin still augmented selectin adhesiveness. Therefore, tether stabilization through the L-selectin tail appears to primarily depend on intrinsic stabilization of singular bonds, even though at high ligand densities, individual microvillar contacts may stabilize through more than one bond (Chen and Springer, 1999) (Fig. 9 A, right panel). Rapid stabilization of the first microvillar contact would favor subsequent formation of mul-

Figure 9.  
(A) A postulated sequence of bond formation, dissociation and rebinding events potentially mediated by L-selectin or tail-truncated mutant at a single microvillus contact site bearing GlyCAM-1 under shear flow \( \geq 1 \text{ dyn/cm}^2 \). At 100 sites/\( \mu \text{m}^2 \), GlyCAM-1 supports rolling through a single GlyCAM-1 molecule at each microvillus contact site. The bond between L-selectin and GlyCAM-1 forms and dissociates within a millisecond irrespective of the selectin tail (left panels). Once dissociated from the ligand, full-length L-selectin or tail-truncated mutant is either restricted or mobile within the microvillus membrane, respectively (center panels). Reduced mobility of L-selectin limits its escape from the original ligand and increases the probability of its rebinding to the same GlyCAM-1 molecule (upper row, third panel). No rebinding takes place for the highly mobile mutant (lower row, third panel). Successful ligand rebinding by anchored L-selectin retards the leukocyte sufficiently to allow subsequent stabilization steps. The neighboring L-selectin (shown as a faded image) is too distant from the GlyCAM-1 site to participate in this interaction, unless GlyCAM-1 density is higher (right panel). For simplicity, only a single microvillus contact site is depicted. (B) Engagement of multiple microvillar sites facilitate cell capture and rolling.
Multiple microvilli tethers at the cell-substrate contact zone (Fig. 9 B). Upon temporary capture, leukocytes are flattened by downward forces that increase their contact area to several \( \mu m^2 \) within a few milliseconds (Evans and Leung, 1989; Evans et al., 2001), which potentially contain several microvillar sites (Chen and Springer, 1999). Multimicrovillar contacts would be important for dispersing the force applied on each selectin bond, slow down its dissociation, and further prolong the lifetime of the tether. Such avidity enhancement may underlie the shear-dependent increase in the number of L-selectin bonds formed at adhesive contact zones during leukocyte capture and rolling (Finger et al., 1996; Lawrence et al., 1997; Chen and Springer, 1999).

L-selectin is the only selectin constitutively associated with the actin cytoskeleton (Pavalko et al., 1995). P-selectin regulates its adhesiveness through constitutive clustering on the cell surface via an association with \( \alpha \)-adaptin, a component of clathrin-coated pits (Setiadi et al., 1998; Ramachandran et al., 2001). Interestingly, interference with the P-selectin \( \alpha \)-adaptin association does not abolish the selectin’s ability to support leukocyte capture and subsequent rolling, although it impairs the mechanical strength of P-selectin tethers (Setiadi et al., 1998; Ramachandran et al., 2001). It is striking that although L-selectin is the shortest selectin, it exhibits the highest efficiency of cell capture at elevated shear stresses among the three selectins (Arbones et al., 1994; Bosse and Vestweber, 1994; Alon et al., 1997) and the highest shear tolerance (Alon et al., 1997; Ramachandran et al., 1999). Thus, the ligand-rebinding mechanism postulated here may provide specialized means for cytoskeletonally anchored L-selectin to mediate cell capture over a wider range of shear conditions than P- or E-selectin. The residual cytoplasmic associations of the L\( \Delta \)cyto mutant gave rise to intermediate tether stabilization levels between those of the L358\( \text{stop} \) mutant and those of full-length L-selectin. Although the RRLKKG segment retained by the L\( \Delta \)cyto mutant was insufficient for L-selectin association with \( \alpha \)-actinin under in vitro conditions of detergent cell lysis (Pavalko et al., 1995), this segment appears to mediate considerable anchorage to the cytoskeleton based on the much higher tether stabilization generated by this mutant relative to the L358\( \text{stop} \) mutant lacking the LKKG segment. These four COOH-terminal cytoplasmic residues may confer residual cytoskeletal anchorage through binding of the regulatory protein calmodulin, also implicated in L-selectin shedding (Kahn et al., 1998). However, PMA-induced or spontaneous shedding of L-selectin were fully conserved upon L-selectin tail truncation (unpublished data), suggesting that differences in L-selectin shedding were unlikely to account for the adhesive differences among L-selectin and its tail mutants. Indeed, inhibition of L-selectin shedding did not rescue the adhesive defects of the tail-truncated mutants (unpublished data).

In conclusion, our study supports the hypothesis that extremely rapid bond stabilization at microvillus-endothelium contact sites is facilitated by cytoskeletal association that anchors L-selectin within the ligand contact site. Our results suggest for the first time that the low force sensitivity of L-selectin tethers may be regulated by a post ligand diffusion-controlled stabilization mechanism in addition to intrinsically kinetic and mechanical properties of the selectin–carbohydrate bond. The notion that cytoskeletal anchorage of L-selectin can dramatically prolong tether lifetime introduces a new regulatory mechanism for L-selectin adhesiveness under shear flow. This anchorage may vary considerably among different cell types as well as in different states of leukocyte activation (Spertini et al., 1991; Roberts et al., 1999). Hyperthermic conditions have been shown to enhance cytoplasmic tail–dependent association of L-selectin with the actin cytoskeleton and to increase L-selectin-dependent lymphocyte binding to HEV (Evans et al., 1999). Further insights into how cytoskeletal L-selectin anchorage is regulated in different leukocytes under homeostatic and inflammatory conditions will allow better understanding of its indispensable functions in immune cell recruitment to lymphoid and extralymphoid organs.

### Materials and methods

#### Antibodies and reagents

The anti–L-selectin mAbs, LAM1–101, and LAM1–118, directed against the SCR domain of L-selectin (Steeber et al., 1997; Li et al., 1998) were a gift of Dr. T.J.F. Tedder (Duke University, Durham, NC). The anti–L-selectin mAb, DREG–200 (Kishimoto et al., 1990), was provided by Dr. T.K. Kishimoto (Boehringer). All mAbs were used as purified Ig. Rabbit anti-murine IgG Fc and FITC-conjugated goat anti–mouse IgG were obtained from Zymed Laboratories, Inc. GlyCAM–1, purified from mouse serum by immunofinity chromatography (Laskey et al., 1992), was a gift from Dr. S.D. Rosen (University of California at San Francisco, San Francisco, CA). PNAd, purified from human tonsil lysates by MECA–79 mAb affinity chromatography (Berg et al., 1991), was a gift from Dr. E.L. Berg (Protein Design Labs). The glycoprotein mixture was stored in 1% octyl glucoside/PBS solution at 4°C. The chemically synthesized neoglycolipid 3’sle[glc]-phosphatidylethanolamine dipalmitoyl (C16:0) was a gift from Dr. L.L. Kiessling (University of Wisconsin, Madison, WI). Fucoidin, a plant-derived sulfated polysaccharide that saturably blocks the lectin domain of L-selectin (Rosen and Bertozzi, 1994), BSA (fraction V), protein A, Ca\(^{2+}\)–, and Mg\(^{2+}\)–free HBSS, Ficoll–Hypaque 1077, cytochalasin B (CB), n-octyl–B–glucopyranoside (octyl glucoside), and poly-L-lysine were all obtained from Sigma–Aldrich. HSA (Fraction V) was obtained from Calbiochem.

#### Cells

The stable expression of full-length L-selectin or L\( \Delta \)cyto in the mouse pre–B-cell line 300.19 was described elsewhere (Kahn et al., 1993). L358\( \text{stop} \) (Fig. 1) was stably expressed in 300.19 cells at comparable levels to the other two transfectants. Clones expressing identical levels of native and mutated L-selectin were maintained as described by Dwir et al. (2000). Peripheral blood granulocytes were isolated from anticoagulated blood after dextran sedimentation and density separation over Ficoll–Hypaque as described by Alon et al. (1997).

#### Laminar flow assays

Preparation of ligand-coated substrates has been described previously (Dwir et al., 2000). GlyCAM–1 site densities were assessed using 125I–labeled CAM–402 (Dwir et al., 2000). Cell-free L-selectin and L-selectin tail mutants were derived from lysates of the various transfected 300.19 cells, as described (Dwir et al., 2000). Polystyrene plates were overlaid with the various selectin variants using a specific capture approach designed to immobilize the selectin variants in a functional state by the use of SCR–specific mAbs. Specificity of the adsorbed L-selectin variants was confirmed by complete blockade of neutrophil adhesion in the presence of the L-selectin ligand, fucoidin.

Transfected cells (2 \( \times \) 10^6/ml) were perfused at room temperature in binding medium (HBSS/10 mM Hepes, pH 7.4, supplemented with 1 mg/ml BSA and 2 mM CaCl\(_2\)) through the flow chamber at desired flow rates as described (Dwir et al., 2000). Cellular interactions were visualized at two fields of view (each one 0.17 mm^2 in area) using a 10\( \times \) objective of an inverted phase contrast microscope (Diaphot 300; Nikon Inc.). Cells were videotaped at 0.02-s resolution with a LIS–700 CCD camera (Applitrek) and a time-lapse SVHS Video recorder (AG–6730; Panasonic). Tethering events were defined as adhesive interactions of those freely flowing cells.
moving closest to the lower wall of the flow chamber coated with the test substrate. Two types of initial cell tethers to the substrate were determined: transient tethers, in which cells attached briefly to the substrate, and rolling tethers, in which tethered cells remained rolling on the substrate, i.e., moving at a mean velocity < 0.25 the hydrodynamic velocity for at least 3 s after initial tethering. Transient tethers to low density ligands was determined as described (Alon et al., 1997).

For inhibition studies, cells were perfused in Ca²⁺−free binding medium supplemented with EGTA, or preincubated in binding medium for 5 min at 4°C with 10 μg/ml of either the L-selectin blocking mAb, DREG-200, or with 50 μg/ml of fucoidin. Dimerization of surface L-selectin on the various transfectants was induced by incubating L-selectin-expressing cells (2 × 10⁷/ml) with the L-selectin dimerizing mAb LAM1-118 (Li et al., 1998), or the control L-selectin SCR-specific mAb, LAM1-101, in binding medium at 25°C for 15 min. To determine the adhesiveness of the various transfectants towards L-selectin ligands expressed on neutrophils, fresh human neutrophils (10⁷/ml) were perfused over either scattered substrate-bound transfected cells or monolayers of closely spaced transfectants immobilized on polystyrene (at 100 ng/ml).

**Computerized microkinetic analysis**

The imaging system developed for quantitative analysis of instantaneous velocities of cell movement, WSCAN-Array-3 (Galai), was described elsewhere (Dwir et al., 2000). Individual transfectants rolling on GlyCAM-1 were tracked for 3 s at 0.02 s resolution and their mean displacement velocities were derived. Two categories of rolling cells, jerky and smooth rolling, were identified based on their mean displacement velocities. Microscopy of individual cells exhibiting jerky rolling on low density ligand was analyzed at high temporal resolution on video segments of rolling cells recorded with a high-speed camera (Motion Counter Analyzer, FAST-CAM-SUPER 500; Eastman Kodak). Cell movement was recorded at a rate of 500 frames/s and digital movies were recorded at a rate of 50 frames/s on SVHS Video recorder. Analogue movies (Fig. 5) were used as a template for the WSCAN-Array-3 software. Motion analysis compared the forward displacement of a moving cell at 0.02 s intervals, corresponding to 0.002 s intervals between successive frames of the original digital movie. Pauses shorter than 0.006 s could not be eliminated by blocking L-selectin. The majority of these pauses were 2 ms long. As they were observed also in freely flowing cells, they were considered as imaging noise. At a shear stress of 1.75 dyn/cm², specific pauses were defined as cell displacements of <0.9 μm during a period of at least 0.006 s. At 0.75–1 or 0.4–0.6 dyn/cm², pauses were defined as displacements of <0.6 μm or <0.4 μm, during this period, respectively. The natural log of the number of pauses with a given duration after pause initiation was plotted against pause duration. First-order dissociation plot yielded a straight line with the slope equal to -koff. The dependence of koff on applied force was assumed to follow the Bell equation (1978): koff = koff⁰ exp (-Fb/kbT), where koff⁰ is the dissociation rate constant in the absence of applied force, Fb is the reactive compliance of the tether, koff is the force on the bond, kb is the Boltzmann’s constant and T is the absolute temperature. The force on the bond was calculated to be 180 pN/1 dyn/cm² wall shear stress using a diameter of 12 μm and assigning a bond angle of 50°. The Bell equation was fit to the koff vs. Fb data using the exponential curve-fitting function in EXCEL.

**Online supplemental material**

Video 1 shows a playback of motions of WT and tail-truncated L-selectin transfectants with substrate bound GlyCAM-1 under physiological shear flow, recorded with a high speed camera at 500 frames/s. The actual time elapsed from the beginning of each recording is indicated at the top of each scene.

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**Cytosplastic regulation of L-selectin adhesion** | Dwir et al. 155

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