Mechanokinetics of receptor–ligand interactions in cell adhesion

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Abstract Receptor–ligand interactions in blood flow are crucial to initiate such biological processes as inflammatory cascade, platelet thrombosis, as well as tumor metastasis. To mediate cell adhesion, the interacting receptors and ligands must be anchored onto two apposing surfaces of two cells or a cell and a substratum, i.e., two-dimensional (2D) binding, which is different from the binding of a soluble ligand in fluid phase to a receptor, i.e., three-dimensional (3D) binding. While numerous works have been focused on 3D kinetics of receptor–ligand interactions in the immune system, 2D kinetics and its regulations have been less understood, since no theoretical framework or experimental assays were established until 1993. Not only does the molecular structure dominate 2D binding kinetics, but the shear force in blood flow also regulates cell adhesion mediated by interacting receptors and ligands. Here, we provide an overview of current progress in 2D binding and regulations, mainly from our group. Relevant issues of theoretical frameworks, experimental measurements, kinetic rates and binding affinities, and force regulations are discussed.

Keywords Receptor–ligand interactions · Selectins · β2 integrins · 2D binding kinetics

1 Introduction

Cell adhesion is a fundamental biological process that is mediated by specific interactions between adhesion receptors and their ligands on other cell surfaces or in the extracellular matrix [1,2]. Such adhesions are important to processes such as inflammatory cascade, platelet thrombosis, as well as tumor metastasis [3]. As a first example, the interactions between selectins and glycoconjugates mediate neutrophil (PMN) tethering to and rolling on vascular surfaces at sites of inflammation or injury [4–6] (Fig. 1). The selectin family of adhesion molecules has three known members: P-, E-, and L-selectin. Their common structure is an N-terminal, C-type lectin (Lec) domain, followed by an epidermal growth factor (EGF)-like module, multiple copies of consensus repeat (CR) units characteristic of complement binding proteins, a transmembrane segment, and a short cytoplasmic domain [6]. P-selectin glycoprotein ligand 1 (PSGL-1), as a major selectin ligand, consists of a homodimer cross-linked by disulfate bonds and binds to selectins by its N-terminal peptide, which includes three tyrosine sulfates and the core-2 O-glycan [7–11]. As a second example, the interactions between β2-integrin of lymphocyte function-associated antigen-1 (LFA-1) or macrophage-1 antigen (Mac-1), expressed on PMNs, and intercellular adhesion molecule 1 (ICAM-1), expressed on endothelial cells, dominate the slow rolling, firm adhesion, and intravascular crawling of PMNs under blood flow [12–14] (Fig. 1). LFA-1 and Mac-1 share a common β2 subunit non-covalently associated with a respective α subunit. The β2 subunit is composed of an I-like domain, a hybrid domain, a plexin/semaphorin/integrin domain, four integrin epidermal growth factor-like domains, a transmembrane domain, and a cytoplasmic tail, while the α subunit includes an I-domain, a β-propeller, a thigm domain, two
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...calf domains, a transmembrane domain, and a cytoplasmic tail [15,16]. Inside-out signaling triggered by selectins and chemokines, or outside-in signaling triggered by ligand binding, induces β2 integrins to undergo a dramatic transition from bent low-affinity (LA) to extended intermediate- (IA) or high-affinity (HA) conformation [17–20]. ICAM-1, a member of the super IgG family, consists of five IgG-like domains (D1–D5) and binds to Mac-1 via the D3 domain and LFA-1 via the D1 domain, respectively [21,22]. As a third example, circulating immunoglobulin G (IgG) binds to foreign particles or damaged tissue through dual antigen-binding fragments (Fab). Their conserved Fc fragment is available for binding to Fcγ receptors (FcγRs) on the immune cell surface, which triggers a wide variety of immune responses [23,24].

Interactions of cellular adhesive molecules are determined by their intrinsic kinetics (reaction rates and binding affinity), since kinetic parameters govern how likely and how fast the adhesion occurs, and how strong and how long the bond remains bound. In a typical 2D binding, at least two specific aspects arise as compared to 3D binding [25]. One is the coupling of kinetics and mechanics, so-called mechanochemical coupling, of receptor–ligand interactions. In blood flow, cell adhesion is regulated by hemodynamic forces which are translated into external forces on interacting molecules. Bond formation and dissociation of interacting adhesive molecules provide the physical linkages between cells. Not only does applied force regulate the lifetime of molecular interactions, just as pressure affects the chemical rates [26,27], but the force loading rate also affects the bond strength of molecular interactions [28,29]. For example, the formation of immunological synapse between a T-cell and an antigen-presenting cell is governed by both 2D binding kinetics of interacting molecule pairs (i.e., T-cell receptor vs. major histocompatibility molecules, integrin vs. ICAM-1 ligand) and mechanics of the cell membrane (i.e., stiffness, rigidity) [30].

Another aspect is the stochastic nature of receptor–ligand interactions. In contrast to 3D binding, where thousands of interacting receptors and ligands are involved and molecular fluctuation of individual molecules is averaged out by their statistical behaviors, molecular bonds in 2D binding are rarely formed inside the contact area. This infrequent occurrence of adhesion introduces the stochastic nature of individual molecules. For example, there are only single or a few bonds of P-selectin-ligand bonds involved in leukocyte–endothelium interactions [31]. Moreover, the number of bonds varies each time. Taken together, these aspects suggest that new theoretical models and experimental measurements are required to understand quantitatively 2D kinetics of receptor–ligand interactions.

In this review, we provided an overview of the current progress in the quantitative understanding of 2D kinetics and forced dissociation of receptor–ligand bindings. Four issues were discussed, including theoretical bases, experimental measurements, 2D kinetics and affinities, and force regulations. These summaries provide new insights into understanding the receptor–ligand interactions in immune responses.

2 Theoretical framework for receptor–ligand kinetics

Consider a second-order forward and first-order reverse reaction,

\[ R + L \overset{k_f}{\rightarrow} B, \]

\[ \overset{k_r}{\rightarrow} \]

where \( R, L, \) and \( B \) denotes, respectively, the receptor, ligand, and bond. In 3D binding, kinetics of a soluble ligand binding to a receptor follows a simple, deterministic kinetic equation,

\[ \frac{d[B]}{dt} = k_f [R][L] - k_r [B]. \]

where \( [R], [L], \) and \( [B] \) denote the concentrations of receptor, ligand, and bond, respectively (in units of molar concentration or M), and \( k_f \) (in unit of M\(^{-1}\) s\(^{-1}\)) and \( k_r \) (in units of s\(^{-1}\)) are the forward and reverse rates, respectively. \( K_a(= k_f / k_r) \) is the binding affinity (in units of M\(^{-1}\)) when the reaction reaches an equilibrium state.

Two-dimensional binding of receptor–ligand interactions in cell–cell or cell–substrate adhesions is a stochastic process regulated by applied forces. On one hand, the stochastic nature of such a binding can be described using a probabili-
tic model. The basic idea is to define the probability of bonds, instead of the concentration of bonds, since the adhesion is no longer a deterministic process. A probabilistic modeling was developed based upon a small system kinetics first proposed by McQuarrie [32], and the adhesion probability, $P_a$, at contact time $t$ follows [33–36],

$$P_a = 1 - \exp \left\{ -A_c m_t m_l K_a^0 \left[ 1 - \exp \left( -k_0^0 t \right) \right] \right\}, \tag{3}$$

where $K_a^0$ (in units of $\mu m^2$) and $k_0^0$ are, respectively, the zero-force binding affinity and reverse rate, $m_t$ and $m_l$ are, respectively, the site densities of receptor and ligand (in units of $\mu m^{-2}$), and $A_c$ is the contact area (in units of $\mu m^2$). Two-dimensional kinetics parameters of $A_c m_l K_a^0$ (if $m_t$ is known) or $A_c K_a^0$ (if both $m_t$ and $m_l$ are known) and $k_0^0$ can be predicted by fitting the experimental measurements of binding curves ($P_a \sim t$ curves) to the model (Eq. 3), and 2D forward rate $k_t$ (in units of $\mu m^2 s^{-1}$) can be obtained by the definition ($= K_a^0 \times k_0^0$). Note that non-specific binding should be subtracted out of total binding before fitting the curve using Eq. 3.

On the other hand, force regulates the formation and dissociation of bonds in blood flow. Two parameters are used to quantify the effect: one is the bond rupture force or bond strength, and another is bond lifetime. The bond rupture force can be experimentally converted from 3D binding affinity through structural analyses and multiscale (atomic-scale molecular dynamics, Monte-Carlo, and lattice) simulations [49], although this review does not focus on the proteins of interest reported here.

3 Experimental approaches for quantifying 2D kinetics and forced dissociation

Until 1993, 2D kinetics measurements were not available experimentally [50]. From then on, many experimental assays have been developed by coordinating the biological experiments and mechanical measurements. These include micropipette aspiration [25,31,36,51], optical tweezers [37,52,53], biological force probes [45,54–56], atomic force microscopy (AFM) [38,39,43,57–61], flow chamber [62–65], microcantilever needle [28], centrifugation [66], rosetting [67], cone-plate viscometer [68,69], surface force apparatus [70,71], and fluorescence recovery after photobleaching (FRAP) [72,73]. Here, two assays of micropipette aspiration and atomic force microscopy are exemplified to demonstrate how they work.

In a micropipette aspiration assay, two cells (generally a human red blood cell (RBC) and a nucleated cell, respectively, expressing or coated with a receptor and the counterpart ligand) are aspirated by two micropipettes with diameters of $\sim 1.5–3 \mu m$ via a suction pressure of 1–4 mmH2O [31,36,51] (Fig. 2a). Adhesion between the RBC and the nucleated cell is staged by placing them in controlled contact via micromanipulation (Fig. 2b). The presence of adhesion and the adhesion force at the end of a given contact period are detected mechanically by observing microscopically the deflection of the flexible RBC membrane upon retracting it away from the nucleated cell (Fig. 2c, d). This contact-retraction cycle is repeated 100 times to estimate the adhesion probability, $P_a$, at that contact duration, $t$. About 100 pairs of cells are used to obtain several $P_a$ vs. $t$ curves that correspond to different receptor and ligand densities, $m_t$ and $m_l$. Equation 3 is used to estimate the zero-force reverse rate, $k_0^0$ and effective binding affinity, $A_c m_l K_a^0$ (if $m_t$ was known) or $A_c K_a^0$ (if both $m_t$ and $m_l$ were known).

In an atomic force microscopy assay, a receptor or ligand is coated directly or captured via a capturing monoclonal antibody (mAb) onto the AFM cantilever tip. Purified counterpart ligands or receptors are incorporated in lipid vesicles and

![Fig. 2](image_url) A test cycle of micropipette aspiration assay in four phases of approach (a), contact (b), withdrawal (c), and detachment (d). Here, a red blood cell (dark cell) serves as a force transducer to determine an adhesive event on membrane deflection.
suitable for real-time measurement of protein conformational change and intracellular signal transduction [75–78]. Nevertheless, this fluorescence-based assay is more complicated than those mechanical-based ones discussed above, and also asks for FRET sensor design, fluorophore conjugation, and additional force sensor calibration. To our knowledge, there are few FRET works in the field of 2D selectin/integrin–ligands binding so far. It is expected that the data collected from both assays will be compared in the future.

4 Progress update of 2D receptor–ligand binding and forced dissociation

4.1 Modeling of cell aggregation and cell adhesion mediated by receptor–ligand interactions

Under physiological blood flow, blood cells not only form homotypic or heterotypic aggregates, they are also able to adhere to the endothelium. Since aggregation and adhesion between two cells are first driven to make contact by shear flow and then are cross-linked by underlying receptor–ligand bonds, a two-body collision theory is coupled with a probabilistic model of small system kinetics and a mechananochemical coupling theory of forced bond dissociation. We first proposed a theoretical model to predict the shear-induced formation and break-up of doublets in three sequential phases: (1) formation upon two-body collision between singlets; (2) evolution of bonds at low shear rate; and (3) break-up of bonds at high shear rate. The predictions upon the model are found to be in good agreement with measurements and enable us to estimate the binding affinity and kinetic rates for three types of doublets cross-linked by two receptor–ligand systems [69]. Then the model was modified by adding a term of time-dependent forward rate to quantify the aggregation of PMNs and tumor cells [68]. Next, the model was further developed to account for the transition from zero-bond singlet to n-bond doublet by adding a term of first bond forming probability from those geometrically available cell–cell collisions. Finally, we are currently integrating cell aggregation in the free stream with cell adhesion in proximity to the endothelium, and we have proposed a unified framework to describe the cell dynamics within the blood vessel.

In addition, we discussed the single-bond hypothesis in the probabilistic model of small system binding kinetics and concluded that no single criterion is sufficient to support the single bond observation, but a cumulative body of evidence may provide reasonable confidence when a point attachment is assumed to be a quantum unit [85]. We are also developing a non-simplified model of competitive binding of receptor and ligand interactions when at least one type of molecules is both in solution and presented on cell surface, which is
Table 1 Summary of 2D kinetics measurements of selectins and integrins

| Receptor | Ligand    | Assay | $r_f$(pN s$^{-1}$) | $A_c K^O_d$(μm$^4$) | $k^O_d$(s$^{-1}$) | $A_c k^O_d$(μm$^4$s$^{-1}$) | $a$(nm) | Reference |
|----------|-----------|-------|-------------------|----------------------|----------------|-----------------------------|--------|-----------|
| sP-selectin | mPSGL-1   | AFM   | 12800–256000      | 0.02                 | 0.2            | 0.09–0.23 μm$^3$s$^{-1}$    | 0.25   | [59]      |
| sP-selectin | mPSGL-1   | AFM   | 100–10000         | 0.2                  | 0.14           | 0.14–0.63                    | 0.23   | [60]      |
| sP-selectin | sPSGL-1   | BFP   | 300–30000         | 0.37                 | 0.14           | 0.10–2.41                    | 0.07–0.72 | [61] |
| sP-selectin | mPSGL-1   | OT    | 25–600            | 1.39–4.3             | 0.14–0.63      | 0.10–2.41                    | 0.07–0.72 | [61] |
| mP-selectin | mPSGL-1   | AFM   | 10–1000000        | 0.08–33.6            | 0.14–0.63      | 0.10–2.41                    | 0.07–0.72 | [61] |
| sP-selectin | mPSGL-1   | AFM   | 130–45600         | 0.91–48.7            | 0.14–0.63      | 0.10–2.41                    | 0.07–0.72 | [61] |
| mP-selectin | mPSGL-1   | OT    | 1–20              | N/D                 | N/D            | N/D                         | 0.14–0.63 | [61] |
| mP-selectin | mPSGL-1   | OT    | 21–188            | 0.37                 | 0.14           | 0.10–2.41                    | 0.07–0.72 | [61] |
| sP-selectin | mPSGL-1   | MAT   | $m_1 A_c K^O_d = 0.08–0.25 \mu m^2$ | 0.9–1.1             | 0.09–0.23 μm$^3$s$^{-1}$ | 0.14–0.63                    | 0.07–0.72 | [61] |
| sP-selectin | mPSGL-1   | Rosett | $A_c K^O_d N = 4.66 \times 10^{-3} \mu m^4$ | 0.6–1.8             | 0.14           | 0.10–2.41                    | 0.07–0.72 | [61] |
| sP-selectin | mPSGL-1   | MAT   | 5.54 \times 10^{-5}–4.55 \times 10^{-3} | 0.6–1.8             | 0.14           | 0.10–2.41                    | 0.07–0.72 | [61] |
| sP-selectin | mPSGL-1   | OT    | 16.4 \times 10^{-8} | 0.15–0.75           | 0.14           | 0.10–2.41                    | 0.07–0.72 | [61] |
| sP-selectin | mPSGL-1   | MAT   | 1.79 \times 10^{-4}–1.52 \times 10^{-3} | 1.0–2.41            | 0.14           | 0.10–2.41                    | 0.07–0.72 | [61] |
| sP-selectin | mPSGL-1   | MAT   | 1.35–5.03         | 0.15–0.75           | 0.14           | 0.10–2.41                    | 0.07–0.72 | [61] |
| mL-selectin | mPSGL-1   | MAT   | $m_1 A_c K^O_d = 0.08–0.25 \mu m^2$ | 0.9–1.1             | 0.09–0.23 μm$^3$s$^{-1}$ | 0.14–0.63                    | 0.07–0.72 | [61] |
| mL-selectin | mPSGL-1   | MAT   | 0.72 \times 10^{-5}–3.98 \times 10^{-5} | 0.60–1.21           | 0.09-0.23 μm$^3$s$^{-1}$ | 0.14–0.63                    | 0.07–0.72 | [61] |
| mL-selectin | mPSGL-1   | MAT   | 0.026–0.103       | 0.5–6.9             | 0.09-0.23 μm$^3$s$^{-1}$ | 0.14–0.63                    | 0.07–0.72 | [61] |
| mL-selectin | mPSGL-1   | MAT   | 0.44–0.92         | 0.09-0.23 μm$^3$s$^{-1}$ | 0.14–0.63      | 0.10–2.41                    | 0.07–0.72 | [61] |
| mL-selectin | mPSGL-1   | MAT   | 0.9             | 0.09-0.23 μm$^3$s$^{-1}$ | 0.14–0.63      | 0.10–2.41                    | 0.07–0.72 | [61] |
| mLFA-1    | sICAM-1   | AFM   | 20–10000          | 0.17–4              | 0.14           | 0.10–2.41                    | 0.07–0.72 | [61] |
| mLFA-1    | sICAM-1   | AFM   | 10000–50000       | 40–57               | 0.09-0.23 μm$^3$s$^{-1}$ | 0.14–0.63                    | 0.07–0.72 | [61] |
| mLFA-1    | sICAM-1   | MAT   | 5.46 \times 10^{-6}–8.57 \times 10^{-5} | 0.19–2.01           | 0.09-0.23 μm$^3$s$^{-1}$ | 0.14–0.63                    | 0.07–0.72 | [61] |
| mLFA-1    | sICAM-1   | BFP   | 10–1000           | 0.008–2             | 0.14           | 0.10–2.41                    | 0.07–0.72 | [61] |
| mLFA-1    | sICAM-1   | BFP   | 34–8180           | 0.0001–0.1          | 0.14           | 0.10–2.41                    | 0.07–0.72 | [61] |
| mLFA-1    | sICAM-1   | OT    | 3.29 \times 10^{-6}–9.13 \times 10^{-6} | 0.55–1.24           | 0.09-0.23 μm$^3$s$^{-1}$ | 0.14–0.63                    | 0.07–0.72 | [61] |
| mLFA-1    | sICAM-1   | OT    | 0.99 \times 10^{-5}–8.39 \times 10^{-5} | 0.35–1.09           | 0.14           | 0.10–2.41                    | 0.07–0.72 | [61] |
| mMac-1    | sICAM-1   | AFM   | 100–100000        | 0.49–2.38           | 0.14           | 0.10–2.41                    | 0.07–0.72 | [61] |
| mMac-1    | sICAM-1   | OT    | 0.09 \times 10^{-6}–3.12 \times 10^{-6} | 0.20–1.12           | 0.14           | 0.10–2.41                    | 0.07–0.72 | [61] |
| sMac-1    | sICAM-1   | OT    | 0.94 \times 10^{-6}–2.26 \times 10^{-6} | 0.43–0.60           | 0.14           | 0.10–2.41                    | 0.07–0.72 | [61] |

AFM: atomic force microscopy, MAT: micropipette aspiration technique, BFP: biomembrane force probe, OT: optical trap, s: soluble, m: membrane
used to quantify the cell adhesion due to the shedding of cellular adhesive molecules off the surface.

4.2 Binding kinetics of receptor–ligand interactions

4.2.1 Selectin–ligand bindings

P-selectin P-selectin is expressed on platelets and activated endothelial cells. Molecular presentation and surface microtopology of the receptors are crucial to their binding kinetics. The randomized orientation or the shortened extension of P-selectin to its ligand- and antibody-binding epitope above the cell membrane lowers the 2D binding affinity to the PSGL-1 ligand by reducing the forward rates, but not the reverse rates [31]. Meanwhile, stiffening the carrier alone or in cooperation with surface-roughing lowers the 2D affinity of P-selectin–PSGL-1 interactions by reducing the forward rate, but not the reverse rate, whereas softening the carrier and roughing the surface have opposing effects on the 2D kinetics [36]. In contrast, the soluble antibody binds with a similar 3D affinity to surface-anchored P-selectins or PSGL-1 constructs regardless of their orientation, length, and carrier stiffness and microtopology [31,36]. A classic rosetting assay is modified to estimate the binding affinity of P-selectin to PSGL-1 (= 4.66 x 10^{-3} μm^2) by assuming that the size of the rosettes is Poisson distributed [67], which is well consistent with those measured in other biophysical assays. To visualize consecutive binding-unbinding transitions and then to quantify the association kinetics of P-selectin to PSGL-1, a thermal fluctuation assay is developed and a 2D forward rate is estimated as higher for the long construct than the short one [79], as observed Ref. [31]. We find also that bond formation is reduced by enhancing the diffusivity of the selectin-coupled carrier and appears to be temporal history-dependent [79].

L-selectin L-selectin is expressed on leukocytes. The binding affinity of L-selectin to PSGL-1 in an equilibrium state is also estimated using a modified rosetting assay (=0.94 x 10^{-3} μm^2) [67]. IL-8-induced L-selectin shedding reduced the number of membrane-anchored L-selectins and enhanced their affinity to PSGL-1 by greatly lowering its reverse rate, but slightly reducing the forward rate, suggesting that two opposite impacts control the rolling dynamics of activated PMNs: reducing the molecular presentation to enhance the rolling and lowering the kinetic rates to reduce the rolling [74]. At least one reason for such an observation stems from mowing the microvilli and smoothing of the surface of cells by cytokine-mediated L-selectin shedding [74], resulting in enhanced accessibility of residual L-selectin to its ligand [36]. Similar to those performed for P-selectin, the association kinetics of L-selectin and its dependence on carrier diffusivity are also determined using the thermal fluctuation assay, which is robust in determining 2D association kinetics and sensitive to the parameters such as sampling rate, sliding window size, and threshold in parametric analysis [79,86].

E-selectin E-selectin is expressed on activated endothelial cells. E-selectin constructs coated on red cells are found to bind in a similar forward rate, but different reverse rate to the ligands expressed on HL-60 or Colo-205 cells bearing carbohydrate ligands, since HL-60 cells only express sLe^a, but Colo-205 cells express both sLe^a and sLe^x [51]. Again, the randomized orientation of E-selectin to its ligand-binding domain above the cell membrane lowered 2D affinity to the PSGL-1 ligand by reducing the forward rates, but not the reverse rates [31].

Selectin ligands. PSGL-1, serving as a major ligand to P- and L-selectin, presents three sulfated tyrosines and an O-glycan to the interface of the selectin Lec domain. Binding kinetics measurements and molecular dynamics simulations of double or triple tyrosine substitutions indicate that these tyrosines regulate the accessibility of PSGL-1 to P- or L-selectin and that at least one of them is required for PSGL-1 binding to P- or L-selectin, which results structurally from the significant conformational change of PSGL-1 peptide, but not its binding site of O-glycan [80].

4.2.2 Integrin–ligand bindings

Two β2-integrins members, Mac-1 (αMβ2) and LFA-1 (αLβ2), are constitutively expressed on PMNs and mediate the PMN recruitment cascade by binding to ICAM-1 expressed on activated endothelial cells under blood flow. Ligand binding of β2-integrin is regulated by or induces conformational changes in the inserted (I) domain. A surprising discrepancy was revealed between the 2D and 3D reverse rate measurements of LFA-1, since 2D reverse rates of the locked open and locked closed I domains only differ a few fold, which is in sharp contrast to the 3D reverse rate of ~100-fold difference [81]. Even with similar structures, LFA-1 and Mac-1 play distinct roles in PMN recruitment, that is, the slow rolling and firm adhesion of leukocytes rely on LFA-1 while cell crawling is dependent on Mac-1. Such distinct functions are assumed to be governed by the differences in their binding affinities and kinetic rates. The difference in binding affinity between Mac-1 and LFA-1 is forward-rate dominated with a slightly or moderately varied reverse rate. This finding was further confirmed when both β2-integrins were activated by chemokines (sMLF or IL-8), divalent cations (Mg2^+ or Mn2^+), or disulfide bond lockage on an HA state. Structural analyses reveal that such kinetics difference is likely attributed to the distinct conformations at the interface of Mac-1 and LFA-1 and ICAM-1 [87]. PMN spreading is mediated specifically by β2-integrin–ICAM-1 interactions and bi-directionally regulated under shear flow. αL, αM and β2

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subunits contribute distinctly to PMN spreading on ICAM-1 substrates [88].

Interactions between β2-integrin on PMNs and ICAM-1 on melanoma cells initiate the bindings of melanoma cells to PMNs within the tumor microenvironment in blood flow, which in turn activate PMN-melanoma cell aggregation in a near-wall region of the vascular endothelium, thereby enhancing subsequent extravasation of melanoma cells in the microcirculations. Hydrodynamic shear regulates PMN-melanoma cell heterotypic aggregation that is dependent on shear rates rather than shear stresses. The heterotypic aggregation appears to reach a peak after ~60 s under shear and then starts to decrease afterward [68]. The cellular binding affinity of a PMN-melanoma cell pair is higher than that of a PMN-endothelial cell pair, but the effective binding affinities per molecular pair are comparable between the two cell pairs no matter whether the melanoma cell or endothelial cell are quiescent or cytokine-activated, indicating that the stronger adhesion between the PMN-melanoma cell pair is mainly attributed to the high expression of ICAM-1 on the melanoma cell. These results propose an alternative mechanism where melanoma cells adhere first with PMNs near vessel wall regions and then bind to endothelial cells via PMNs under blood flow [35].

4.3 Forced dissociation of receptor–ligand binding

4.3.1 Selectin–ligand bonds

External forces are found to affect the bond of receptor–ligand interactions. While the lifetime of the P-selectin-G1 mAb bond is reduced monotonically with applied forces (slip bond), the P-selectin–PSGL-1 bond yields an ascending phase (catch bond) followed by a descending phase with increased force [48]. There are several physical factors to regulate the forced dissociation of the receptor–ligand complex. For instance, the rupture force of P-selectin–PSGL-1 binding reaches a saturated plateau, following a transition phase, with the contact time and yields a threshold with the approach velocity, while the adhesion probability presents a biphasic feature with the retraction velocity [43]. A piecewise linear relationship between rupture force and logarithm of the loading rate is found at high (~10^4 pN s^{-1}) [43] and low (~10^{-1} pN s^{-1}) [89] for the P-selectin–PSGL-1 bond while the contribution of non-specific binding to the measured rupture force is not negligible at the higher loading rate (>89 pN s^{-1}) [84]. At a low loading rate (~20 pN s^{-1}), rupture force varies with different combinations of spring constant and retraction velocity even at the same loading rate and increases with the spring constant, indicating that the bond dissociation of P-selectin–PSGL-1 interactions is spring constant-dependent at a low loading rate and that the bond rupture force depends on both loading rate and mechanical compliance of the force transducer at a high loading rate [37]. Not only the rupture force, but also the bond lifetime is sensitive to the spring constant of the force transducer used. One finds that the bond lifetime of P-selectin–PSGL-1 interactions presents stochastic distributions at different spring constants and the catch bond nature is visualized at ≥3.0 × 10^{-2} pN nm^{-1} [90].

4.3.2 Integrin–ligand bonds

Shear resistance of β2-integrin–ICAM-1 is critical to maintain the slowing rolling, firm adhesion, and crawling of PMNs onto the endothelium under blood flow. A fast and a slow linear loading regime are separately observed in the dynamic force spectra of the LFA-1–ICAM-1 bond, which indicates a steep inner activation barrier and a wide outer activation barrier, respectively. The equilibrium dissociation constant of the LFA-1–ICAM-1 interaction is regulated by the energetics of the outer activation barrier of the complex, while the ability of the complex to resist a pulling force is determined by the divalent cation-dependent inner activation barrier [39]. The binding probability and adhesion force of Mac-1 with ICAM-1 are enhanced upon Mac-1 activation [58]. Small molecule agonist Leukadherin-1 fosters binding of Mac-1 to ICAM-1 via the formation of long membrane tethers, whereas Mn^{2+} additionally increases ICAM-1 binding via cytoskeleton-anchored bonds [91]. Long lifetimes and increased bond strength also occur when β2-integrin is activated [82,83]. Three states with distinct reverse rates are identified from lifetime distributions of LFA-1–ICAM-1 bond. Force shifts the associated fractions from the short- to intermediate- and long-lived states, producing catch bonds at low forces, but increases their reverse rates exponentially, converting catch to slip bonds at high forces [45]. One also notes that the elasticity of extracellular matrix (ECM) regulates the integrin affinity. For example, a soft ECM increases the activation level of integrins while a stiff ECM has a tendency to prevent the dissociation and internalization of bound integrins [92,93].

4.4 Structural bases of receptor–ligand interactions

4.4.1 Selectin–ligand interactions

Molecular structure and conformation governs its functions and varies with local chemical and biological environments. For example, applying external forces enable the structural collapse of the P-selectin molecule, which is mainly attributed to the burst of hydrogen bonds within the major β sheet of its EGF domain and the disruption of two hydrophobic cores of its Lec domain [94]. The intramolecular extension is also observed in the forced dissociation of P-selectin–PSGL-1 complex prior to the intermolecular sepa-
ration of fucose group of PSGL-1 from Ca\(^{2+}\) ion of P-selectin [95]. Further allosteric simulations indicate that the bent conformation of EGF-like domain of P-selectin with respect to its Lec domain is unable to switch directly to an extended conformation but presents a spontaneous allostery to a novel, relatively extended conformation starting with the separation between residues Q30 and K67 and terminating with the release of residue N87 from residue C109 [96]. More physically, two new molecular dynamics approaches, one involving the shear flow field with a controlled velocity gradient and the other presenting track dragging with a defined trajectory are developed to investigate the microstructural evolution and dissociation kinetics of P-selectin–PSGL-1 interactions under shear flow, indicating that the shear flow alone induces the destruction of Lec/EGF domains within the P-selectin construct before the complex dissociates and that the cooperation of shear flow and tensile stretch mediates the intramolecular destruction of EGF domain and the breaking of hydrogen bond clusters at the P-selectin Lec/EGF interface [97].

### 4.4.2 Integrin–ligand interactions

Molecular dynamics simulations are also employed to elucidate the conformational stability of \(\alpha\) subunit I domains of LFA-1 and Mac-1 in different affinity states and relevant I domain-ICAM-1 interaction features. Compared with low LA Mac-1, the LA LFA-1 I domain is unstable in the presence or absence of ICAM-1 ligand, stemming from diverse orientations of its \(\alpha7\)-helix with different motifs of a zipper-like, hydrophobic junction between \(\alpha1\)- and \(\alpha7\)-helices. Meanwhile, spontaneous transition of the LFA-1 I domain from a LA state to an IA state is first visualized. LA Mac-1 I domain is not favorable for ICAM-1 binding but HA Mac-1 and HA/LA LFA-1 I domain is able to bind to the ICAM-1 ligand readily in free MD simulations, implying that the binding pocket of LA Mac-1 could not open spontaneously, presumably because S144 residue prevents the cation in MIDAS from interacting with D229 residue of ICAM-1 D3 domain [98]. Distinct interface conformations are found between Mac-1 and LFA-1 in different affinity states, which determines the accessibility and availability of MIDAS cation to D3 or D1 domain of ICAM-1 ligand and vary the forward rate value of the two molecules [87].

### 5 Summary

Quantifying 2D kinetics and forced dissociation of receptor–ligand interactions in cell adhesions is crucial to further the understandings in immune responses. A probabilistic model of a small system is developed to predict 2D kinetic rates and binding affinities, while a mechaehemical coupling model is introduced to analyze forced regulation of receptor–ligand interactions. The state-of-the-art techniques including micropipette aspiration and atomic force microscopy are widely used to measure the receptor–ligand binding kinetics and regulation of applied forces. Structural variation, surface environment, and membrane microtopology and stiffness affect the kinetic rates and affinities. Applied forces regulate the bond strength and lifetime in multiple phases.

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