

Structural basis of the leukocyte integrin Mac-1 I-domain interactions with the platelet glycoprotein Ib

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Key Points

- The integrin Mac-1 I-domain interaction with platelet receptor GPIb α has been mapped by NMR.
- The GPIb α LRR capping α -helix coordinates directly to the Mac-1 MIDAS Mg²⁺ ion via an acidic residue.

Cell-surface receptor interactions between leukocyte integrin macrophage-1 antigen (Mac-1, also known as CR3, α M β 2, CD11b/CD18) and platelet glycoprotein Ib α (GPIb α) are critical to vascular inflammation. To define the key residues at the binding interface, we used nuclear magnetic resonance (NMR) to assign the spectra of the mouse Mac-1 I-domain and mapped the residues contacting the mouse GPIb α N-terminal domain (GPIb α N) to the locality of the integrin metal ion-dependant adhesion site (MIDAS) surface. We next determined the crystal structures of the mouse GPIb α N and Mac-1 I-domain to 2 Å and 2.5 Å resolution, respectively. The mouse Mac-1 I-domain crystal structure reveals an active conformation that is stabilized by a crystal contact from the α 7-helix with a glutamate side chain completing the octahedral coordination sphere of the MIDAS Mg²⁺ ion. The amino acid sequence of the α 7-helix and disposition of the glutamic acid matches the C-terminal capping region α -helix of GPIb α effectively acting as a ligand mimetic. Using these crystal structures in combination with NMR measurements and docking analysis, we developed a model whereby an acidic residue from the GPIb α leucine-rich repeat (LRR) capping α -helix coordinates directly to the Mac-1 MIDAS Mg²⁺ ion. The Mac-1:GPIb α N complex involves additional interactions consolidated by an elongated pocket flanking the GPIb α N LRR capping α -helix. The GPIb α N α -helix has an HxxxE motif, which is equivalent by homology to RxxxD from the human GPIb α N. Subsequent mutagenesis of residues at this interface, coupled with surface plasmon resonance studies, confirmed the importance of GPIb α N residues H218, E222, and the Mac-1 MIDAS residue T209 to formation of the complex.

Introduction

The integrin receptor macrophage-1 antigen (Mac-1, also known as CR3, integrin α M β 2, or CD11b/CD18) is expressed on the surface of myeloid leukocytes and mediates numerous responses of these cells critical to innate immunity.¹ The Mac-1 receptor contributes to the recruitment, firm adhesion, and transendothelial migration of leukocytes at sites of vascular injury and facilitates tissue inflammation.² Biochemical and cell-based studies have characterized the interactions of the Mac-1 integrin with diverse ligands, including plasma protein fibrinogen,³ complement protein fragment iC3b,⁴ high-molecular-weight kininogen,⁵ and the cell-surface receptors platelet glycoprotein Ib α (GPIb α)² and intercellular adhesion molecule 1 (ICAM-1).⁶ The Mac-1 integrin exists in an inactive conformation and, when activated by a variety of stimuli, undergoes a structural change to an active form capable of binding to ligands with high affinity.^{7,8} Concurrently, allosteric changes occur on ligand binding that result in “outside-in” cell signaling.⁹

Mac-1-deficient mice show delayed thrombosis, but largely unimpaired hemostasis.¹⁰ Blocking the Mac-1:GP1b α interaction prevents neutrophil extracellular trap formation,¹¹ and an antibody targeting Mac-1 was shown to block inflammation.¹² Inhibiting the Mac-1 I-domain with the small molecule allosteric regulator leukaderin-1 has demonstrated efficacy in animal models of inflammatory disease.^{13,14}

For the majority of Mac-1 interactions described, the inserted α M-subunit, or “I-domain,” is the principal ligand-binding domain. The Mac-1 I-domain contains a Mg²⁺ binding site on its surface at the top of the β sheet called the metal ion-dependant adhesion site (MIDAS), which is capable of accepting the side chain of an acidic residue (aspartate or glutamate) from coordinating ligands.^{1,7} A crystal structure has defined the interaction between Mac-1 and iC3b,⁴ but the complex between Mac-1 and GPIb α and the principal ligand fibrinogen³ are poorly understood at the molecular level. Here, nuclear magnetic resonance (NMR) assignment of the mouse Mac-1 I-domain, together with X-ray structural analysis of both the mouse GP1b α N and Mac-1 I-domain, has enabled us to map the binding surface and model the interaction site for the GP1b α leucine-rich repeat (LRR) N-terminal domain (GP1b α N) with the MIDAS face of Mac-1.

Materials and methods

Protein expression, purification, and characterization

A complementary DNA fragment encoding mouse GP1b α N residue E1 to T266 (mature protein sequence numbering) was cloned into the pMT-puro vector for expression using the DES system (Invitrogen). At the N terminus, the signal sequence corresponds to a *Drosophila* homolog of the immunoglobulin binding chaperone protein secretion signal, and a polyhistidine tag sequence TRTGHHHHHH was added at the C terminus. Site-specific mutations H218A and E222A were introduced using the QuikChange method (Stratagene) and confirmed by DNA sequencing. The mouse amino acid sequence numbering is defined by alignment with the mature human GP1b α sequence, such that H218 corresponds to H234 in the UniProt numbering of entry GP1BA_MOUSE, which includes the signal sequence. *Drosophila* S2 cells were grown in Dulbecco's modified Eagle medium supplemented with 10% fetal bovine serum at 28°C and transfection was performed in each case using calcium phosphate. Cells were grown for an additional 48 hours before selection with puromycin to establish stable cell lines. Serum-free Express Five (Invitrogen) insect culture media was collected containing secreted proteins. Initial capture of the crude product was performed using Ni-sepharose affinity chromatography, followed by gel filtration and ion exchange chromatography (supplemental Figure 1). Protein identity and successful removal of the signal peptide was confirmed by trypsin digest mass spectrometry, analyzed using the Mascot Database (ProteinID, University of York) shown in supplemental Figures 1 and 2.

The mouse Mac-1 I-domain gene fragment (P129 to A318; mature protein sequence numbering) was cloned into the pGEX-4T-1 vector, for expression as an N-terminal GST-fusion using *Escherichia coli*. All mutants were produced using Q5 site-directed mutagenesis. XL1-Blue and BL21 cells were transformed using calcium chloride for storage and expression, respectively. BL21 cells were grown in LB media (for surface plasmon resonance [SPR] and mass spectrometry) or M9 minimal media (for NMR spectroscopy) supplemented with ¹⁵N ammonium chloride and/or ¹³C glucose as required for

isotope labeling. GST-affinity chromatography and incubation with thrombin resulted in the separation of the tag (residual Gly-Ser remains at the N terminus) and further purification using cationic exchange chromatography. Mass spectrometry, gel filtration studies, NMR line width analysis, and diffusion measurements indicate that the recombinant Mac-1 I-domain is monomeric in solution (supplemental Figures 3 and 4).

NMR spectroscopy

NMR data were recorded at 25°C on a Bruker 800 MHz Avance III spectrometer with a QCI cryoprobe using Topspin 3.1 software. Backbone resonances (¹³C/¹⁵N) were assigned using HNCO, HN(C α)CO, C β C α NH, and C β C α (CO)NH experiments. Backbone assignment (>146/182 residues) was facilitated by a number of locally perturbing single-point mutations (to Ala/Gly) and through selective unlabeled experiments (Glu and Lys, in particular). ¹⁵N-TROSY 2D spectra were collected in titration experiments at a Mac-1 concentration of 120 μ M in 25 mM potassium phosphate buffer (pH 7.0), 25 mM NaCl, 5% D₂O, and 0.02% NaN₃ using 1.5 equivalents of MgCl₂ to ensure metal complexation. ¹⁵N-TROSY spectra were collected at 0.2 molar ratio intervals up to a 2:1 excess of GPIb α N. Chemical shift perturbations (CSPs) were calculated using a weighting of ¹H and ¹⁵N shifts according to: CSP = $\sqrt{[1/2 \times (\delta_H^2 + 0.14 \times \delta_N^2)]}$. For structural modeling, the standard deviation (SD) from the mean CSP was calculated and mean + SD and mean + 2 \times SD used for surface mapping.

Quantitation of Mac-1 I-domain ligand binding

SPR was performed using a BIAcore 3000 (GE Healthcare). Ligands of recombinant mouse GPIb α N, GPIb α N variants H218A and E222A, and fibrinogen (σ) were amine-coupled to a CM5 sensor chip (GE Healthcare); a reference cell was prepared by blank amine-coupling. Recombinant Mac-1 I-domain wild-type and T209A variant samples at protein concentrations ranging from 3.125 μ M to 200 μ M, in running buffer (10 mM *N*-2-hydroxyethylpiperazine-*N*'-2-ethanesulfonic acid, pH 7.4, 140 mM NaCl, 1 mM MgCl₂) were injected at a flow rate of 50 μ L/min. Kinetic studies of the Mac-1 I-domain wild-type and T209A constructs were conducted as a dilution series, in the presence and absence of Mg²⁺ against the immobilized GPIb α N. Each of the series contains a minimum of 10 concentrations (11.25-200 μ M) and was repeated a minimum of 2 times. The Mg²⁺ ion dependence of the GPIb α N interaction was tested against a wild-type Mac-1 I-domain concentration series (6-100 μ M) in which the running buffer contained 5 mM EDTA to sequester metal ions. Binding to the immobilized GPIb α N was also evaluated using a dilution series of the antibody Xia.B2 (emfret ANALYTICS) with protein concentration of 3 to 50 nM. The CM5 chip was regenerated with 2 M NaCl after each experiment. Binding curves were generated by subtracting the appropriate controls and the resulting curves were analyzed using BIAevaluation software or a steady-state equilibrium approach using a Hill plot generated with Prism 6 (GraphPad Software Inc.).

Protein crystallization and crystal structure determination

Recombinant mouse GP1b α N was concentrated to 12 mg/mL and subjected to sparse matrix screens of Morpheus, Proplex, Index, PACT, JCSG+, and MIDAS using the sitting drop vapor diffusion method. Crystals of GP1b α N grew in Morpheus condition D2 containing 0.2 M mixture of 1,6-hexanediol; 1-butanol; 1,2-propanediol;

2-propanol; 1,4-butanediol; 1,3-propanediol, 0.1 M imidazole; MES monohydrate (acid) pH 6.5, 20% v/v ethylene glycol; and 10% w/v PEG 8000. Crystals of the mouse Mac-1 I-domain (6 mg/mL) grew in the JCSG+ screen condition D7, 0.2 M lithium sulfate, 0.1 M tris(hydroxymethyl)aminomethane, pH 8.5, 40% PEG 400. Crystals were cryocooled in liquid nitrogen and X-ray diffraction data were collected using the ID30B beamline at the ESRF (GP1b α N) and Diamond beamline I24 (Mac-1 I-domain). Data were processed with XDS and molecular replacement was performed with PHASER using a human GP1b α N¹⁵ and Mac-1 I-domain structure⁷ (Table 1). Partial model building was performed using BUCCANEER and completed manually using COOT and refined with REFMAC5 (deposited PDB code: 6EJX).

Molecular docking calculations

Three-dimensional models of the Mac-1 I:GP1b α N docked complex structures were calculated using HADDOCK¹⁶ with template Mac-1 I-domain (PDB code: 1IDO) and GP1b α N (PDB code: 1M0Z) crystal structures used. Active residues were defined based on the Mac-1 I-domain NMR titration experiments: E244, G143, and N147 plus the MIDAS Mg²⁺ ion. Active residues for GP1b α N were D222 and R218.

Results

NMR assignment and structural analysis of the Mac-1 I-domain

A bacterially expressed mouse Mac-1 I-domain construct (residues 129-318) was isotopically (¹³C/¹⁵N) labeled and the NMR spectra assigned using heteronuclear 3-dimensional NMR experiments. The ¹H/¹⁵N heteronuclear single quantum coherence is highly dispersed with sharp resonances indicative of well-ordered secondary structure in both the Mg²⁺ free and bound state (supplemental Figure 5). We also prepared and characterized 2 Mac-1 mutants, T209A and F302W, which have previously been reported as "inactive" (T209A) and "pseudo-open" (F302W) variants of the I-domain.^{17,18} The side chain of T209 coordinates directly with the Mg²⁺ ion only in the "open" conformation of the Mac-1 I-domain,^{4,7} and previous crystallographic studies have defined a large allosteric conformational shift between "open" and "closed" states, which links the switching of the coordination of the metal ion from D140 to T209 to a shift in the position of helix α 7.^{7,19} T209 has also been shown to play a direct and critical role in Mac-1 ligand binding to ICAM-1 and iC3b.⁶ The CD spectra of the mutants and wild-type Mac-1 are superimposable, showing no significant changes in protein secondary structure and folding (supplemental Figure 1B). NMR of the ¹⁵N-labeled mutant proteins confirms retention of the core structure; however, extensive perturbations across the MIDAS face are evident in both mutants (supplemental Figure 5).

NMR analysis of the Mac-1 I-domain interaction with GP1b α N

Recombinant mouse GP1b α N was expressed and purified from insect cells (supplemental Figures 1 and 2). NMR chemical shifts (¹H/¹⁵N) are particularly sensitive to changes in chemical environment and the conformation associated with ligand binding and were used to characterize the GP1b α N interaction with the Mac-1 I-domain. A buffered solution of GP1b α N was titrated into a 120 μ M solution of ¹⁵N-Mac-1 I-domain in the presence of 1.5 molar equivalents of Mg²⁺, and ¹H/¹⁵N heteronuclear single quantum coherence spectra were

Table 1. Crystallographic data collection and refinement statistics

Sample	Mouse GP1b α N	Mouse Mac-1 I
Data collection		
Space group	P2 ₁ 2 ₁ 2 ₁	P4 ₁ 2 ₁ 2
Cell dimensions		
a, b, c (Å)	61.5, 72.8, 164.0	62.9, 62.9, 336.2
α , β , γ (°)	90.0, 90.0, 90.0	90.0, 90.0, 90.0
Resolution, Å	29.9-2.0	45.9-2.5
R _{merge} *	11.1	12.5
I/ σ ; CC(1/2)†	7.0 (1.9); 0.997 (0.78)	8.2 (1.5); 0.903 (0.68)
Completeness (%)†	99.0 (90.0)	86.7 (67.2)
Redundancy†	4.3 (1.3)	6.2 (2.5)
Refinement		
Number of reflections	48095	21236
R _{work} /R _{free} ‡	0.192/0.237	0.222/0.271
B factors, Å ²		
Protein	29.8	48.2
RMS deviations		
Bond lengths, Å	0.018	0.021
Bond angles, °	1.96	1.85

RMS, root mean square.

*R_{merge} = Sum(h) [Sum(j) |I(hj) - <Ih>| / Sum(hj) <Ih>], where I is the observed intensity and <Ih> is the average intensity of multiple observations from symmetry-related reflections calculated with SCALA.

†Values in parentheses are for the highest resolution shell.

‡R_{work} = Sum(h) ||F_oh - |F_ch| | / Sum(h)|F_oh|, where F_o and F_c are the observed and calculated structure factors, respectively. R_{free} computed as in R_{work}, but only for 5% randomly selected reflections, which were omitted in refinement, calculated using REFMAC.

collected over a range of ligand concentrations up to a Mac-1 I:GP1b α N molar ratio of 1:2.

A significant number of fast-exchange CSPs are observed in response to ligand binding. However, a number of residues within the Mac-1 I-domain C-terminal region (helix α 7) showed some evidence of signal broadening and peak distortion in the bound state. Statistically significant CSP effects are identified for 16 residues of the Mac-1 I-domain (Figure 1) and enable us to identifying 2 well-defined binding patches. The first group of residues, which are in close proximity or intimately involved in MIDAS Mg²⁺ coordination, includes D140, S142, G143, I145, and D242 together with the adjacent solvent-exposed residues N147, K210, and E244. Residues D242, E244, and D273 cluster on the surface of the I-domain to form a negatively charged patch that is adjacent to the MIDAS. The second group of Mac-1 residues I265, R266, F275, G293, E294, H295, and V296 (Figure 1D-E) are distal to the MIDAS and appear to be linked to it through conformational changes propagated away from the metal center.

Crystal structures of the mouse GP1b α N and Mac-1 I-domain

We next determined the crystal structure of mouse Mac-1 I-domain to 2.5 Å resolution. This revealed the mouse Mac-1 I-domain in an open conformation with the MIDAS metal ion coordinated by T209, S142, S144, and a glutamic acid residue (E314) ligand mimetic supplied by a crystal contact from the C-terminal helix (α 7) of the

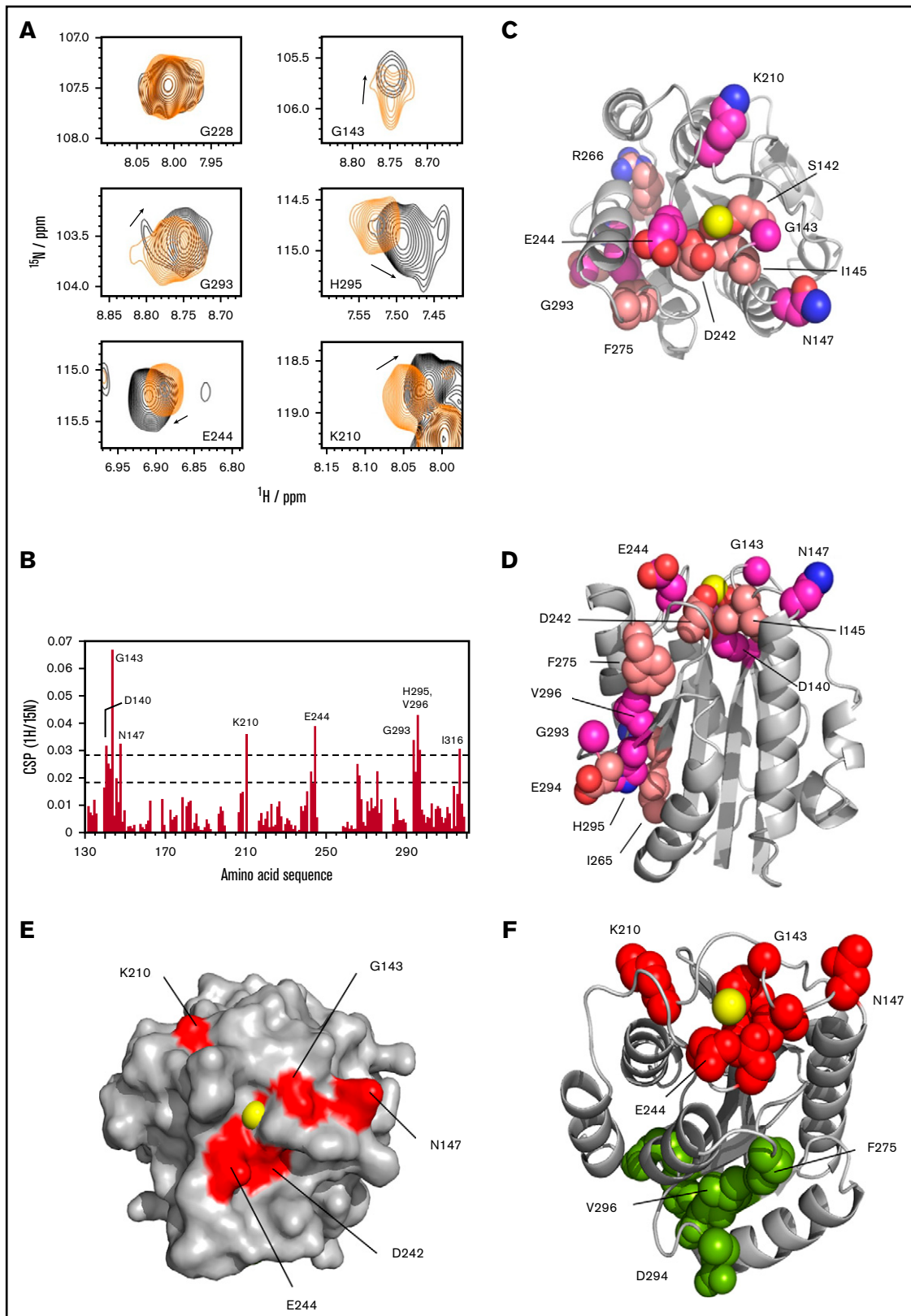


Figure 1. Mapping the Mac-1 I-domain binding site with GP1b α N by NMR. (A) Selected peaks showing CSP effects between bound and unbound (the first panel for G228 is shown as a control, in which no perturbation is observed). (B) Representation of CSP effects, with dashed lines showing statistically significant perturbation (lower line, mean CSP +1 SD; upper line, mean CSP +2 SD) residues above the latter. (C) CSP effects mapped to the Mac-1 surface with residues in darker pink corresponding to

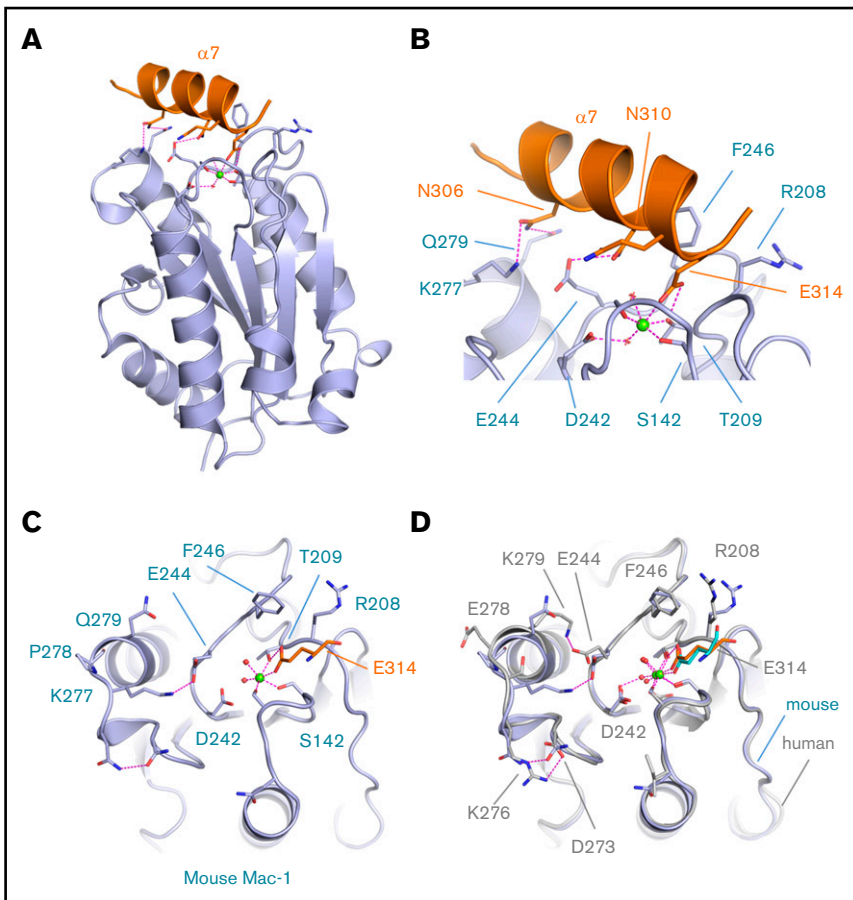


Figure 2. Crystal structure of the mouse Mac-1 I-domain.

(A) A cartoon diagram is shown for the crystal structure of the mouse Mac-1 I-domain (purple) with the crystal contact stabilizing the active conformation derived from the C-terminal helix $\alpha 7$ in orange. The MIDAS bound Mg^{2+} ion is shown as a sphere (green) and key residues are shown as sticks and electrostatic interactions as dashed magenta lines. (B) A close-up view of the MIDAS site with residues from helix $\alpha 7$ labeled. (C) A second view of the Mac-1 I-domain MIDAS site with the crystal contact residue E314 shown coordinating the Mg^{2+} ion and the sidechain hydroxyl of T209. (D) Superposition of the human (gray) and mouse (purple) crystal structures in the region of the MIDAS face. The stabilization glutamic acid is shown as sticks in cyan (human) and orange (mouse).

I-domain (orange in Figure 2A; supplemental Video 1). The C-terminal Mac-1 I-domain α -helix consists of the amino acid sequence 310-NQLQE-314, which resembles the sequence from the α -helix in the GPIIb α N capping region spanning residues 218-HWLQE-222, and is equivalent structurally in terms of the α -helical disposition of the glutamic acid residue. Figure 2B shows that the α -helix sits in a groove on the surface of the Mac-1 I-domain packing against residues F246 and R208, forming multiple hydrogen bonded contacts between the sidechains of N310 and N306 and the sidechains of E244 and K277 on the MIDAS face. For comparison, crystal structures are available for the human Mac-1 I-domain as an open conformation in complex with ligand iC3b,⁴ simvastatin,²⁰ and stabilized by a crystal contact.^{7,8} The mouse and the human Mac-1 I-domain crystal structures are superposed in Figure 2D, revealing the MIDAS metal ion coordinating acidic residue consistently occupies an identical position simultaneously bonding to both the metal ion and the hydroxyl of T209. Subtle differences occur across the MIDAS face with the introduction of K277 in the mouse Mac-1 forming a salt bridge to E244, replacing the K279-E244 salt bridge observed in the human structure (Figure 2C).

We then determined the crystal structure of the mouse GPIIb α N to 2.0 Å resolution to examine the conformation of the LRR C-terminal

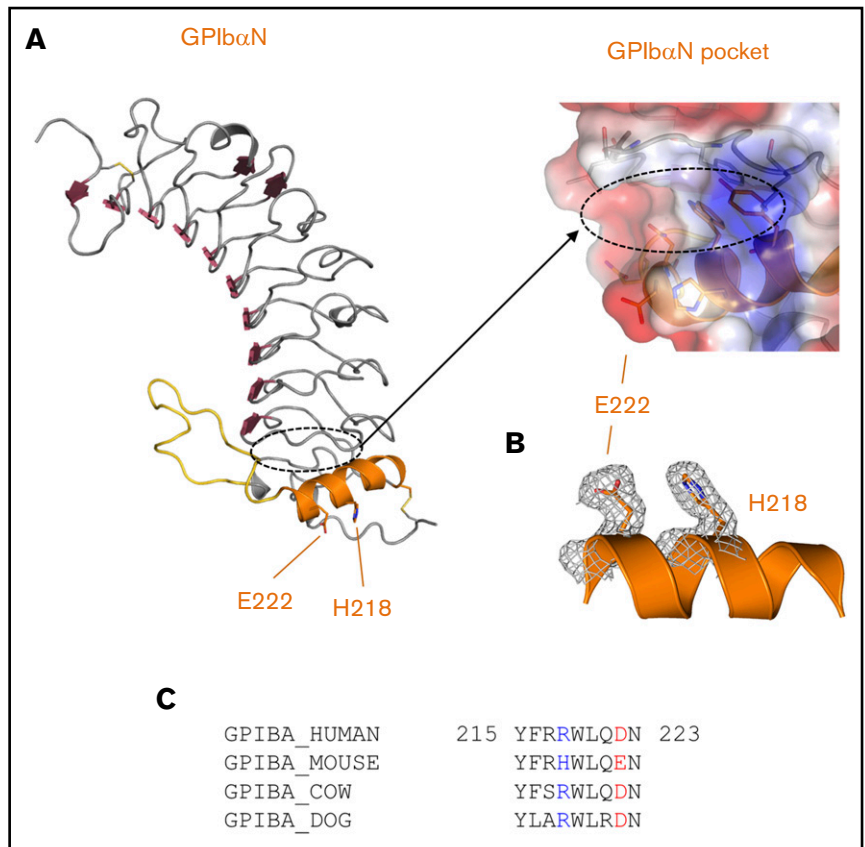
capping α -helix, which is important for the interaction with Mac-1.² This structure reveals overall features of the LRR solenoid flanked by N- and C-terminal disulphide-bonded capping regions and R-loop, as observed previously for the human GPIIb α N structure.²¹ The junction between the C-terminal capping region α -helix and the LRR solenoid forms an elongated pocket (Figure 3A, dotted elliptical line). In a notable departure from the human sequence, Figure 3B-D reveals that the mouse GPIIb α N has the substitutions H218 and E222 that, by analogy with the human GPIIb α N structure, are also surface exposed (Figure 3B). A comparison of structures shows that mouse GPIIb α N H218 and E222 occur with the same disposition as human GPIIb α N R218 and D222, respectively, and a sequence alignment of this region reveals that charged residues at these positions are conserved across other mammalian species (Figure 3C).

Docking of the GPIIb α N and Mac-1 I-domain 3-dimensional structures

Previous cell-based and biochemical studies identified residues 218 through 228 from GPIIb α as the principal determinants of the Mac-1 interaction² encompassing the human GPIIb α N LRR C-terminal

Figure 1. (continued) mean +2 SD, top-down view onto the MIDAS (Mg^{2+} shown in yellow). (D) Side view showing distal CSP effects removed from the MIDAS site (top). (E) Surface representation showing the binding patch around the MIDAS with key residues in red. (F) Differentiation between the MIDAS binding patch, shown in red, and those distal residues (in green) affected through allosteric changes from binding GPIIb α N at the MIDAS site (a few key residues are highlighted).

Figure 3. Crystal structure of the mouse GP1b α N. (A) A cartoon diagram is shown for the crystal structure of the mouse GP1b α N with the C-terminal LRR capping α -helix colored orange and R-loop colored yellow. GP1b α residues H218 and E222 involved in the interaction with Mac-1 are shown as sticks. The elongated pocket formed in the GP1b α LRR capping region is indicated by a dashed line and represented as a transparent charged surface with key residues lining the pocket shown as sticks. (B) Electron density (2Fo-Fc) from the crystal structure of the mouse GP1b α N C-terminal capping α -helix residues H218 and E222 (gray mesh). (C) Amino acid sequence alignment of GP1b α residues 215 through 223 (human sequence numbering without the signal sequence) from the GP1b α LRR capping α -helix for human, mouse, dog, and cow. Key charged residues 218 and 222 are colored blue and red, respectively.



capping α -helix, but did not specify whether a critical acidic residue was involved. Studies using a variety of techniques implicated Mac-1 as residing on the MIDAS face and residues T213 and R216 as being located away from the MIDAS face, but did not propose a role for the Mg²⁺ ion or an overall model for the complex with GP1b α N.^{10,22} We thus set out to investigate this by generating models of the Mac-1 I:GP1b α N complex using the program HADDOCK.¹⁶ Multiple crystal structures are available for the human Mac-1 I-domain in the open conformation^{4,7,8,18,23} and human GP1b α N domain,^{15,21,24-26} and they were used as templates for docking.

Docking was performed using NMR distance restraints between the Mac-1 MIDAS Mg²⁺ ion and residues G143, N147, D242, and E244, which were identified as having significant CSP effects in the NMR titration experiments, together with GP1b α N surface-exposed residues R218, D222, and N223.² The top scoring structure and details of the interface are shown in Figure 4A. The Mac-1 I:GP1b α N complex buries a surface area of 593 Å² (calculated using PISA), and 2 key features that define the unique orientation are that GP1b α N D222 coordinates to the Mac-1 MIDAS Mg²⁺ ion and GP1b α N R218 forms bidentate salt bridges with Mac-1 residues D242 and E244 (supplemental Videos 2 and 3). As viewed down the central projection of the GP1b α N capping α -helix, colored orange in Figure 4B, residues G143 and S144 of Mac-1 (lower left) hydrogen bond to the side chains of N221 and R217 of GP1b α , respectively. Flanking interactions also occur on the right of Figure 4B as R208 from Mac-1 forms hydrogen bonds to the side chains of N223 of GP1b α , and its own H195 main chain carbonyl. The Mac-1 I-domain F246 side chain inserts into an elongated hydrophobic

pocket in the GP1b α N LRR capping region defined by GP1b α residues F192, S194, Y215, and W219 (Figure 4C).

SPR analysis of the Mac-1 I:GP1b α N interaction

To further evaluate the contribution of amino acid side chains at the center of the proposed mouse Mac-1 I:GP1b α N interface, we used site-directed mutagenesis coupled with SPR. To establish the assay, the SPR response was first observed with Mac-1 I-domain analyte and immobilized full-length fibrinogen, which has been characterized in detail from previous studies.³ SPR analysis (Figure 5A) shows a high-affinity interaction with evidence of multiple binding modes; an estimated K_d of 30 μM was derived from the equilibrium values of response units was consistent with previous reports.³

A ligand surface of GP1b α N was generated through amine-coupling to a CM5 chip. To determine the integrity of the GP1b α N surface, a dilution series with the antibody Xia.B2 (emfret ANALYTICS) was conducted and analysis of sensorgrams resulted in a K_d of 34 nM, consistent with the anticipated high affinity for an antibody-antigen interaction, confirming the immobilized GP1b α N is forming an active surface. Sensorgrams were recorded using wild-type Mac-1 I-domain as the analyte and high-quality SPR curves were obtained (Figure 5B, top), such that association and dissociation profiles fitted well to a 1:1 Langmuir simultaneous binding model. A calculated K_d of 64 μM was in good agreement with the steady-state equilibrium K_d value of 121 μM.

Our model of the Mac-1 I:GP1b α N complex shows that the Mac-1 I-domain T209 is a pivotal residue at the binding interface not only

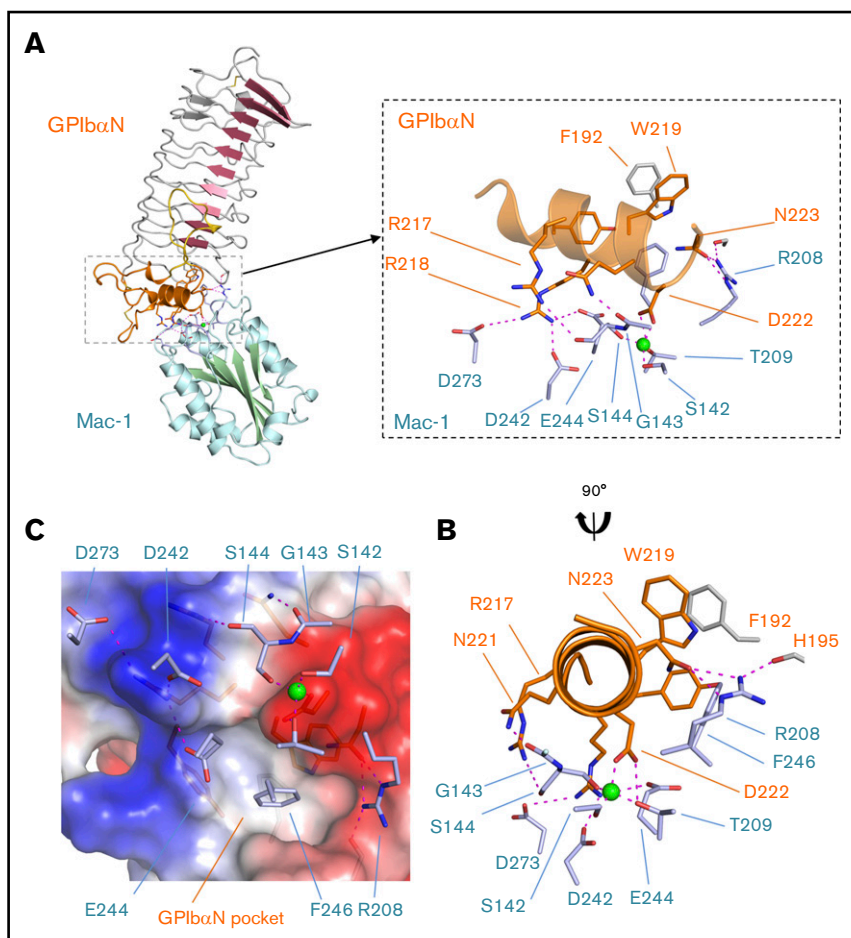


Figure 4. The Mac-1 I:GP1b α N complex. (A) A cartoon diagram of the docked complex of the crystal structures of human GP1b α N and the Mac-1 I-domain with the Mac-1 secondary structures colored blue/green and GP1b α N colored red/orange. Boxed is a close-up view of the interface where the Mg²⁺ ion bound to the Mac-1 MIDAS site is shown as a sphere and electrostatic interactions are shown as dashed purple lines. The GP1b α N C-terminal LRR capping region is colored orange and residue F192 from the LRRs is in gray. (B) A second view related by a 90° rotation. GP1b α N residue D222 coordinates the Mac-1 MIDAS Mg²⁺ ion and surrounding residues on the MIDAS surface (light blue). (C) Charged surface representation of the GP1b α N LRR capping region showing an elongated pocket formed by GP1b α residues Y215, W219, and F192 is flanked by a region of positive charge (GP1b α R218) and negative charge (GP1b α D222) that form complementary interactions with Mac-1 residues shown as sticks (light blue).

by coordinating the MIDAS metal ion, but also in forming a hydrogen bond from the β -OH to the carboxylate of GP1b α N D222. The binding to GP1b α N of the Mac-1 T209A alanine variant (T209A) elicited neither a significant signal response nor a clear association binding curve (Figure 5B, bottom), which was similarly the case with wild-type Mac-1 with EDTA in excess (5 mM) to remove the metal ion from the MIDAS (data not shown). These experiments demonstrate that the bound Mg²⁺ ion in the MIDAS is critical to the interaction and confirm that the observed distal CSP effects in the NMR titration data arise from allosteric effects propagated away from the MIDAS site, rather than to a second ligand-binding site. In contrast with GP1b α N, at least 1 of these binding interactions is still evident from the SPR analysis with full-length fibrinogen and Mac-1 T209A, although the overall binding affinity is greatly reduced (Figure 5A, bottom). This is consistent with a primary binding site around the MIDAS surface, along with a second, low-affinity Mac-1 I-domain site.³

We further tested the model of the Mac-1 I:GP1b α N complex by generating recombinant mouse GP1b α N variants H218A and E222A, which were purified with a similar protocol and produced a similar yield to the wild-type protein (supplemental Figure 1A). SPR studies using the immobilized GP1b α N mutants with the wild-type Mac-1 I-domain as the analyte revealed that GP1b α N H218A and E222A substitutions led to a loss of SPR response, preventing the sensorgrams from being fitted using either kinetic or equilibrium

binding models, indicating significant impairment of the interaction with the Mac-1 I-domain (Figure 5C).

Discussion

Interactions among innate immune cells, platelets, and plasma proteins have been implicated in pathways leading to thrombosis,^{10,27-29} and yet very few of these protein interactions have been fully described. To the best of our knowledge, we provide the first detailed description of the complex formed between the leukocyte integrin Mac-1 I-domain and the platelet receptor GP1b α N using a combination of NMR, crystallography, site-directed mutagenesis, and SPR binding studies. We establish the principle that the Mac-1 I-domain in the open conformation has a groove on the surface formed by the extended sidechains of F246 and R208 and other residues that can accommodate an α -helix appropriately positioned by an acidic side chain coordinating the MIDAS metal ion and the side chain hydroxyl of T209. We show that the GP1b α N receptor LRR C-terminal capping α -helix uses a key acidic residue (D222 in the human; E222 in the mouse ortholog) to coordinate to the vacant site in the metal ion coordination shell of the Mac-1 MIDAS Mg²⁺ ion.

The presence of the integrin-binding acidic residue at the end of an α -helix in GP1b α N is reminiscent of the crystal structure of the Mac-1 I-domain complex with iC3b, whereby the integrin MIDAS metal ion binding residue D1247 of iC3b is also placed at the end of an

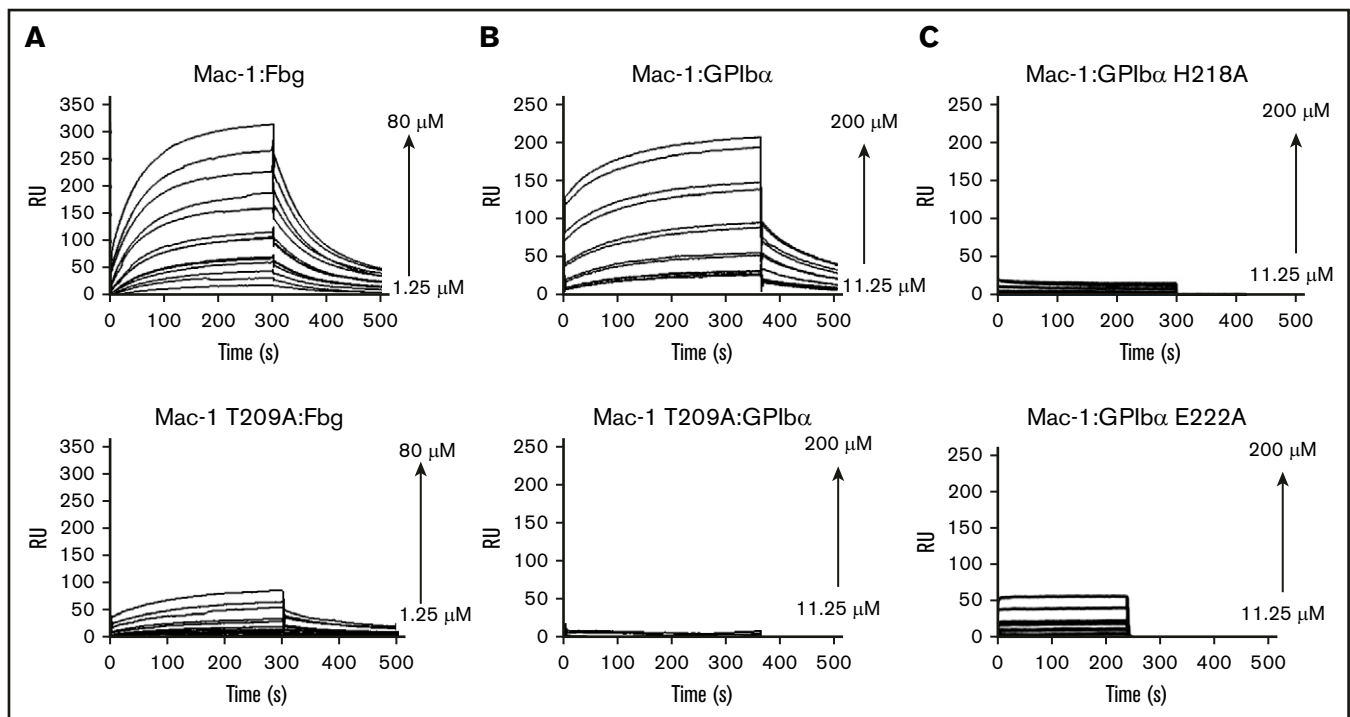


Figure 5. SPR analysis of Mac-1 I-domain interactions. Plots of SPR sensorgrams measured in RU on the y-axis illustrating wild-type Mac-1 (top) or Mac-1 T209A (bottom) binding to fibrinogen (Fbg; A) and GPIb α N (B), and shown in wild-type Mac-1 I-domain binding to the GPIb α N mutants H218A (C, top) and E222A (C, bottom). RU, response unit.

α -helix.⁴ A comparison of the Mac-1 I:iC3b and Mac-1 I:GPIb α N complexes reveals the MIDAS-coordinating α -helix is oriented differently (90° rotation) in each ligand relative to the Mac-1 I-domain surface. In addition, unlike GPIb α N, iC3b does not use the negatively charged patch (E244) on the surface of the Mac-1 I-domain for binding, but in common with GPIb α N it does use Mac-1 I-domain residue R208.⁴ These interactions suggest that the ability of the Mac-1 I-domain to bind a host of specific ligands involves a common binding surface and fits with the previously proposed mosaic model of Mac-1 ligand binding.³⁰ By contrast, GPIb α N uses 3 spatially separated regions of the receptor to bind the structurally distinct ligands of the von Willebrand factor A1 domain^{24,26} (GPIb α LRRs and R-loop), thrombin^{25,31} (GPIb α anionic region), and Mac-1 (GPIb α C-terminal LRR cap).

The Mac-1 MIDAS interaction with an acidic residue from an α -helix of the ligand contrasts to the closely related integrin leukocyte function associated antigen-1 (LFA-1), which exploits a glutamate residue located within a β strand of the ligand ICAM-1³² to bind the LFA-1 MIDAS α -subunit I-domain. A variety of studies have defined a large allosteric conformational shift between open and closed states of the leukocyte integrins Mac-1 and LFA-1 linked to the disposition of α -I-domain helix α 7 and switching of the coordination shell within the MIDAS between D140 and T209.^{7,9} NMR studies have been previously reported for the related LFA-1 I-domain,³³ but to date no NMR characterization has been applied to Mac-1. LFA-1 does not bind GPIb α N, and a series of LFA-1/Mac-1 chimeras have been used previously to localize the binding site for GPIb α N within the Mac-1 I-domain to specific residues. Many of these are consistent with the Mac-1:GPIb α N

interface shown in Figure 4,²² but residues T213 and R216 are not part of the MIDAS face of the Mac-1 I-domain structure.¹⁰ The crystal structures of Mac-1 I-domain reveals the T213 side chain hydroxyl group is buried and hydrogen bonded by contact with the D248 side chain in both the open and closed conformation and thus is not present on the surface to interact with a ligand. Mac-1 residues T213 and R216 are very close to key MIDAS face residues R208 and F246 and thus may affect binding indirectly, or, alternatively, R216 could form part of a second binding site for GPIb α N.

Overall, we believe this is the first detailed description of the interface formed between an integrin receptor and an LRR protein revealing an elongated pocket in the GPIb α N LRR C-terminal capping region, which is key to the interaction. The Mac-1 integrin is not the only integrin to engage multiple ligands and, interestingly, the more distantly related integrin $\alpha_2\beta_1$ binds to collagen¹⁹ as the principal ligand and has a well-characterized additional interaction with the LRR C-terminal capping region of the matrix protein chondroadherin.³⁴ The binding site for the $\alpha_2\beta_1$ integrin is localized to the chondroadherin sequence of residues LRRWLEAK,³⁴ which resembles the FRRWLQDN sequence we report for GPIb α . Chondroadherin crystal structures reveal the LRRWLEAK sequence has an acidic residue that is surface exposed within the α -helix of the LRR capping region in a similar manner to GPIb α N. Previous authors also have speculated that the groove adjacent to the α -helix could be used by the integrin as described here.^{35,36} This structural scaffold describing an integrin-LRR protein interaction will facilitate the development of agents targeting Mac-1 or GPIb α N for the modulation of thrombotic and inflammatory disorders.

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Authorship

Contribution: J.A.L., M.S.S., and J.E. designed the research project; J.M. collected and assigned the nuclear magnetic resonance data with technical assistance from H.E.L.W., with nuclear magnetic resonance analysis of additional mutants by C.A.; cloning, protein expression, and isotopic labeling was carried out by J.M. with additional mutants

produced by A.F.; X-ray analysis of mouse GP1b α N was carried out by M.S. and Mac-1 I-domain by R.N. and S.G.C.; surface plasmon resonance experiments were conducted by J.M., S.S.W., and A.D.M.; all authors discussed the results; and J.E., M.S.S., J.A.L., and J.M. contributed to the writing of the paper.

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References

1. Diamond MS, Garcia-Aguilar J, Bickford JK, Corbi AL, Springer TA. The I domain is a major recognition site on the leukocyte integrin Mac-1 (CD11b/CD18) for four distinct adhesion ligands. *J Cell Biol.* 1993;120(4):1031-1043.
2. Simon DI, Chen Z, Xu H, et al. Platelet glycoprotein Ibalpha is a counterreceptor for the leukocyte integrin Mac-1 (CD11b/CD18). *J Exp Med.* 2000;192(2):193-204.
3. Lishko VK, Podolnikova NP, Yakubenko VP, et al. Multiple binding sites in fibrinogen for integrin alphaMbeta2 (Mac-1). *J Biol Chem.* 2004;279(43):44897-44906.
4. Bajic G, Yatime L, Sim RB, Vorup-Jensen T, Andersen GR. Structural insight on the recognition of surface-bound opsonins by the integrin I domain of complement receptor 3. *Proc Natl Acad Sci USA.* 2013;110(41):16426-16431.
5. Wachtfogel YT, DeLa Cadena RA, Kunapuli SP, et al. High molecular weight kininogen binds to Mac-1 on neutrophils by its heavy chain (domain 3) and its light chain (domain 5). *J Biol Chem.* 1994;269(30):19307-19312.
6. Kamata T, Wright R, Takada Y. Critical threonine and aspartic acid residues within the I domains of beta 2 integrins for interactions with intercellular adhesion molecule 1 (ICAM-1) and C3bi. *J Biol Chem.* 1995;270(21):12531-12535.
7. Lee JO, Rieu P, Arnaout MA, Liddington R. Crystal structure of the A domain from the alpha subunit of integrin CR3 (CD11b/CD18). *Cell.* 1995;80(4):631-638.
8. Xiong JP, Li R, Essafi M, Stehle T, Arnaout MA. An isoleucine-based allosteric switch controls affinity and shape shifting in integrin CD11b A-domain. *J Biol Chem.* 2000;275(49):38762-38767.
9. Sen M, Koksal AC, Yuki K, Wang J, Springer TA. Ligand- and cation-induced structural alterations of the leukocyte integrin LFA-1. *J Biol Chem.* 2018;293(17):6565-6577.
10. Wang Y, Gao H, Shi C, et al. Leukocyte integrin Mac-1 regulates thrombosis via interaction with platelet GPIb α . *Nat Commun.* 2017;8:15559.
11. Carestia A, Kaufman T, Rivadeneyra L, et al. Mediators and molecular pathways involved in the regulation of neutrophil extracellular trap formation mediated by activated platelets. *J Leukoc Biol.* 2016;99(1):153-162.
12. Wolf D, Anto-Michel N, Blankenbach H, et al. A ligand-specific blockade of the integrin Mac-1 selectively targets pathologic inflammation while maintaining protective host-defense. *Nat Commun.* 2018;9(1):525.
13. Maignel D, Faridi MH, Wei C, et al. Small molecule-mediated activation of the integrin CD11b/CD18 reduces inflammatory disease. *Sci Signal.* 2011;4(189):ra57.
14. Dickinson CM, LeBlanc BW, Edhi MM, et al. Leukadherin-1 ameliorates endothelial barrier damage mediated by neutrophils from critically ill patients. *J Intensive Care.* 2018;6(1):19.
15. McEwan PA, Andrews RK, Emsley J. Glycoprotein Ibalpha inhibitor complex structure reveals a combined steric and allosteric mechanism of von Willebrand factor antagonism. *Blood.* 2009;114(23):4883-4885.
16. Dominguez C, Boelens R, Bonvin AM. HADDOCK: a protein-protein docking approach based on biochemical or biophysical information. *J Am Chem Soc.* 2003;125(7):1731-1737.
17. Li R, Rieu P, Griffith DL, Scott D, Arnaout MA. Two functional states of the CD11b A-domain: correlations with key features of two Mn²⁺-complexed crystal structures. *J Cell Biol.* 1998;143(6):1523-1534.

18. McCleverty CJ, Liddington RC. Engineered allosteric mutants of the integrin alphaMbeta2 I domain: structural and functional studies. *Biochem J*. 2003; 372(Pt 1):121-127.
19. Emsley J, Knight CG, Farndale RW, Barnes MJ, Liddington RC. Structural basis of collagen recognition by integrin alpha2beta1. *Cell*. 2000;101(1): 47-56.
20. Jensen MR, Bajic G, Zhang X, et al. Structural basis for simvastatin competitive antagonism of complement receptor 3. *J Biol Chem*. 2016;291(33): 16963-16976.
21. Uff S, Clemetson JM, Harrison T, Clemetson KJ, Emsley J. Crystal structure of the platelet glycoprotein Ib(alpha) N-terminal domain reveals an unmasking mechanism for receptor activation. *J Biol Chem*. 2002;277(38):35657-35663.
22. Ehlers R, Ustinov V, Chen Z, et al. Targeting platelet-leukocyte interactions: identification of the integrin Mac-1 binding site for the platelet counter receptor glycoprotein Ibalpha. *J Exp Med*. 2003;198(7):1077-1088.
23. Mahalingam B, Ajroud K, Alonso JL, et al. Stable coordination of the inhibitory Ca²⁺ ion at the metal ion-dependent adhesion site in integrin CD11b/CD18 by an antibody-derived ligand aspartate: implications for integrin regulation and structure-based drug design. *J Immunol*. 2011; 187(12):6393-6401.
24. Huizinga EG, Tsuji S, Romijn RA, et al. Structures of glycoprotein Ibalpha and its complex with von Willebrand factor A1 domain. *Science*. 2002; 297(5584):1176-1179.
25. Celikel R, McClintock RA, Roberts JR, et al. Modulation of alpha-thrombin function by distinct interactions with platelet glycoprotein Ibalpha. *Science*. 2003;301(5630):218-221.
26. Dumas JJ, Kumar R, McDonagh T, et al. Crystal structure of the wild-type von Willebrand factor A1-glycoprotein Ibalpha complex reveals conformation differences with a complex bearing von Willebrand disease mutations. *J Biol Chem*. 2004;279(22):23327-23334.
27. Pathak M, Kaira BG, Slater A, Emsley J. Cell receptor and cofactor interactions of the contact activation system and factor XI. *Front Med (Lausanne)*. 2018;5:66.
28. Engelmann B, Massberg S. Thrombosis as an intravascular effector of innate immunity. *Nat Rev Immunol*. 2013;13(1):34-45.
29. von Brühl ML, Stark K, Steinhart A, et al. Monocytes, neutrophils, and platelets cooperate to initiate and propagate venous thrombosis in mice in vivo. *J Exp Med*. 2012;209(4):819-835.
30. Ustinov VA, Plow EF. Identity of the amino acid residues involved in C3bi binding to the I-domain supports a mosaic model to explain the broad ligand repertoire of integrin alpha M beta 2. *Biochemistry*. 2005;44(11):4357-4364.
31. Dumas JJ, Kumar R, Seehra J, Somers WS, Mosyak L. Crystal structure of the GpIbalpha-thrombin complex essential for platelet aggregation. *Science*. 2003;301(5630):222-226.
32. Shimaoka M, Xiao T, Liu JH, et al. Structures of the alpha L I domain and its complex with ICAM-1 reveal a shape-shifting pathway for integrin regulation. *Cell*. 2003;112(1):99-111.
33. Huth JR, Olejniczak ET, Mendoza R, et al. NMR and mutagenesis evidence for an I domain allosteric site that regulates lymphocyte function-associated antigen 1 ligand binding. *Proc Natl Acad Sci USA*. 2000;97(10):5231-5236.
34. Haglund L, Tillgren V, Addis L, Wenglén C, Recklies A, Heinegård D. Identification and characterization of the integrin alpha2beta1 binding motif in chondroadherin mediating cell attachment. *J Biol Chem*. 2011;286(5):3925-3934.
35. Paracuellos P, Kalamajski S, Bonna A, Bihan D, Farndale RW, Hohenester E. Structural and functional analysis of two small leucine-rich repeat proteoglycans, fibromodulin and chondroadherin. *Matrix Biol*. 2017;63:106-116.
36. Rämisch S, Pramhed A, Tillgren V, Aspberg A, Logan DT. Crystal structure of human chondroadherin: solving a difficult molecular-replacement problem using de novo models. *Acta Crystallogr D Struct Biol*. 2017;73(Pt 1):53-63.