

Role of the β -Subunit Arginine/Lysine Finger in Integrin Heterodimer Formation and Function¹

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Formation of the integrin $\alpha\beta$ heterodimer is essential for cell surface expression and function. At the core of the $\alpha\beta$ interface is a conserved Arg/Lys “finger” from the β -subunit that inserts into a cup-like “cage” formed of two layers of aromatic residues in the α -subunit. We evaluated the role of this residue in heterodimer formation in an α A-lacking and an α A-containing integrin α V β 3 and α M β 2 (CD11b/CD18), respectively. Arg261 of β 3 was mutated to Ala or Glu; the corresponding Lys252 of β 2 was mutated to Ala, Arg, Glu, Asp, or Phe; and the effects on heterodimer formation in each integrin examined by ELISA and immunoprecipitation in HEK 293 cells cotransfected with plasmids encoding the α - and β -subunits. The Arg261Glu (but not Arg261Ala) substitution significantly impaired cell surface expression and heterodimer formation of α V β 3. Although Lys252Arg, and to a lesser extent Lys252Ala, were well tolerated, each of the remaining substitutions markedly reduced cell surface expression and heterodimer formation of CD11b/CD18. Lys252Arg and Lys252Ala integrin heterodimers displayed a significant increase in binding to the physiologic ligand iC3b. These data demonstrate an important role of the Arg/Lys finger in formation of a stable integrin heterodimer, and suggest that subtle changes at this residue affect the activation state of the integrin. *The Journal of Immunology*, 2008, 180: 1713–1718.

Integrins are $\alpha\beta$ heterodimeric cell surface receptors, which mediate cell-matrix and cell-cell adhesion functions that underlie vital cellular processes such as differentiation, migration, proliferation, and survival (1). Integrins also contribute to many pathologies, including cardiovascular disease, cancer invasion, immune dysfunction, and osteoporosis, thus constituting important therapeutic targets. Normally expressed in an inactive state, integrins switch rapidly to the physiologic ligand-binding (active) state in response to chemical signals generated from within cells (inside-out signaling) (reviewed in Ref. 2). Physiologic ligand binding then transmits conformational signals back to the cytoplasmic tails, thus activating intracellular signal transduction pathways (outside-in signaling). This bidirectional signaling involves conformational changes the nature of which is incompletely understood.

Crystal structure of α V β 3 ectodomain revealed a globular “head” (Fig. 1A) sitting on top of two nearly parallel “legs” (3). The integrin head comprises a seven-bladed β -propeller from the α -subunit and a von Willebrand factor type A (VWFA)⁴ domain

(β A or I-like) from the β -subunit (Fig. 1A). β A is inserted into an Ig-like “hybrid” domain that is inserted in turn into an N-terminal plexin/semaphorin/integrin domain (3, 4). Four epidermal growth factor-like domains followed by a novel β -tail domain form the rest of the β -subunit leg. The α -subunit leg is formed of an Ig-like “thigh” domain followed by two large β -sandwich domains “calf 1” and “calf 2”. The α - and β -subunit legs fold back at a $\sim 135^\circ$ angle, forming a V-shaped structure, bent at its knees “genu” (between the thigh domain and calf 1 domains of the α -subunit) and EGF-1 and 2 of the β -subunit. The leg from each subunit terminates into a rod-shaped single membrane-spanning segment and a short cytoplasmic tail. The ligand-binding site in integrins is localized on the upper side of the integrin head, formed by a metal-ion-dependent adhesion site (MIDAS) in β A and an adjacent region in the α -subunit’s propeller domain. In some integrins (α A-integrins), the α -subunit contains an extra VWFA domain (α A or I) that acts as a ligand relay, capturing exogenous ligands through its own MIDAS and engaging the β A MIDAS as an endogenous ligand (5).

The largely hydrophobic propeller/ β A interface buries a surface area of $\sim 1,600 \text{ \AA}^2$ (2), and is responsible for formation of the integrin heterodimer. It bears a striking structural resemblance to the interface between G β - (a seven-bladed β -propeller) and G α (a VWFA domain) subunits of G-proteins (6). At the core of the propeller/ β A interface in integrins lies a basic residue (Arg261 in β 3, Lys in all other β -subunits except β 4) within a 3_{10} -helix; its side chain “finger” inserts into a cup-like structure lined by aromatic side chains in the central cavity of the propeller, and stabilized by hydrophobic and cation- π contacts (7) (Fig. 1, A and B). Most naturally occurring mutations that disrupt heterodimer formation in patients with β 2- or β 3-integrin deficiency (8–11) lie within or in the immediate vicinity of the 3_{10} helix, but do not involve the Arg/Lys finger. A forward genetic screen for mutations in the integrin β subunit (β PS) of *Drosophila melanogaster* also identified mutations in the vicinity of the 3_{10} helix, all resulting in a strong lethal phenotype (12). These observations suggest an important structural role for the Arg/Lys finger in formation of the

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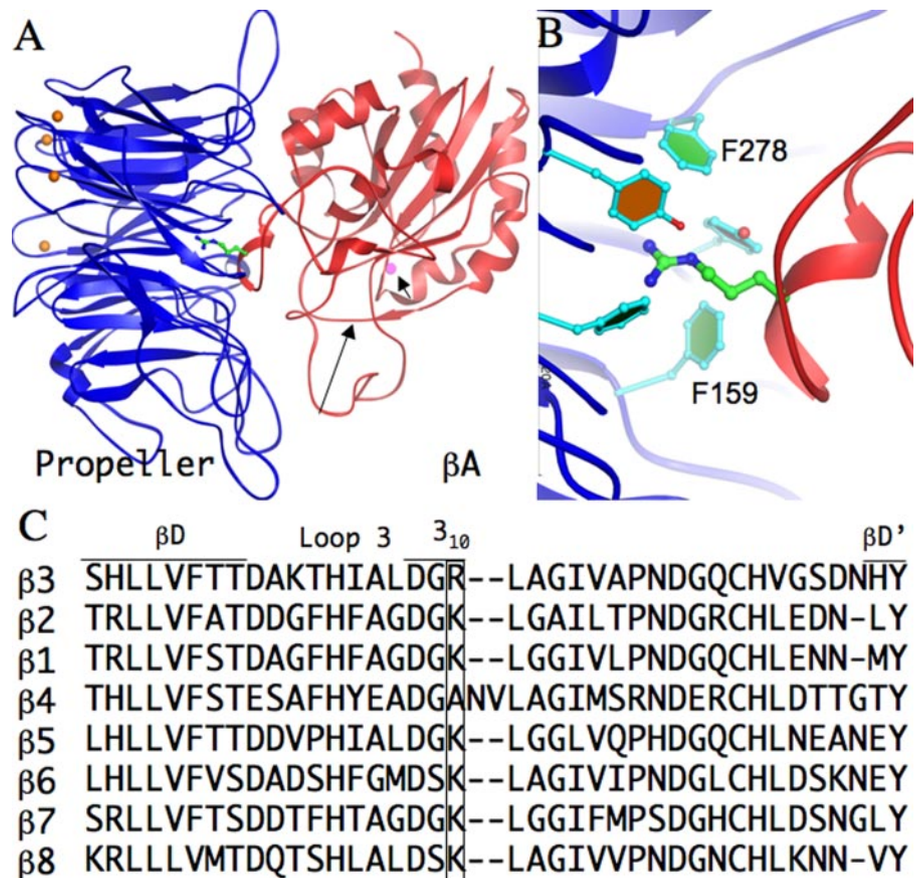
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⁴ Abbreviations used in this paper: VWFA, von Willebrand factor type A; MIDAS, metal-ion-dependent adhesion site; WT, wild type.

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FIGURE 1. Structure and conservation of the Arg/Lys finger in integrin β -subunits. *A*, Three-dimensional crystal structure of unliganded integrin $\alpha V\beta 3$ head segment (as a ribbon diagram with the αV propeller domain in blue and the βA domain in red) showing the position of the arginine finger (Arg261, carbon atoms in green, amides in blue) projecting from the 3_{10} helix. Loop 3 (long arrow) and the ADMIDAS cation (magenta sphere, short arrow) are indicated. The four metal ions at the base of the propeller are indicated as orange spheres. *B*, Higher magnification of the Arg/Lys coordination pocket, with a portion of the aromatic side chains forming the upper ring of the cup-like “cage” structure of αV propeller (adapted from Refs. 7 and 33). Orientation is the same as in *A*. *C*, Structure alignment of all eight mammalian β -subunits around the Arg/Lys finger (boxed). The amino acid sequence of strands βD and $\beta D'$, loop 3 and the 3_{10} helix are indicated in single letter code. The $\beta 4$ -subunit structure model was generated using Modeler (34) with the $\alpha V\beta 3$ structure (PDB code: 1jv2) as template.



integrin heterodimer. In this report, we investigated the role of the Arg/Lys finger in heterodimer formation in the αA -lacking and αA -containing integrins $\alpha V\beta 3$ and CD11b/CD18, respectively.

Materials and Methods

Reagents and Abs

Restriction and modification enzymes were obtained from New England Biolabs, Life Technologies, or Fisher Scientific. All cell culture reagents were from Invitrogen Life Technologies. The anti- $\beta 3$ mAb (mAb) AP3 (IgG1) (13) was from American Type Culture Collection (ATCC); the anti- αV mAb LM142 (IgG1) and the $\alpha V\beta 3$ heterodimer-specific mAb LM609 (IgG1) (14), the anti- $\beta 3$ mAb CBL479 and the anti- αV polyclonal AB1930 were from Millipore. The anti-CD11b mAbs 44a (IgG2a) (15), 903 (IgG1) (16), 107 (IgG1) (17), and a rabbit polyclonal anti-human CD18 Ab have been described previously (18).

DNA constructs

Full-length human wild-type (WT) αV and $\beta 3$ were provided by Dr. Simon Goodman (Merck-Serono KGaA Darmstadt, Germany) (19). Both were recloned into the pcDNA3 expression vector using standard recombinant DNA protocols. In brief, for generating WT αV in pcDNA3, the unique *HindIII* site in the multiple cloning region of an empty pcDNA3 vector was removed and WT αV cDNA from the original vector was transferred into this region using *EcoRI*. For generating WT $\beta 3$ in pcDNA3, WT $\beta 3$ cDNA from the original vector was released using *XhoI* and *XbaI* restriction enzymes and transferred into the multiple cloning site in an empty pcDNA3 vector. Site-directed mutagenesis in $\beta 3$ - and $\beta 2$ -subunits (*CD18*) was conducted using standard recombinant DNA protocols (20) in the newly created $\beta 3$ pcDNA3 and *CD18* pcDNA3 expression vectors (21). PCR-amplified DNA was transferred back into the original $\beta 3$ - or *CD18*-containing expression vector and each mutation was confirmed by DNA sequencing.

Cell culture and transfection

Transient transfection assays were conducted in HEK 293 cells (ATCC) as described (5). In brief, HEK 293 cells were plated on tissue culture treated p-10 plates (BD Clontech) and maintained in complete medium consisting

of DMEM (Invitrogen Life Technologies) supplemented with 10% heat-inactivated FBS (Invitrogen Life Technologies) and 50 IU/ml penicillin and streptomycin (Invitrogen Life Technologies) at 37°C. Cells were detached with 0.05% Trypsin-EDTA solution (Invitrogen Life Technologies) and plated in wells of six-well plates at a concentration of 1 million cells per well. Cells at ~70% confluence were transfected with super coiled cDNAs encoding WT or mutant β -subunits together with corresponding α -subunit, using Lipofectamine2000 reagent (Invitrogen Life Technologies), following the manufacturer's protocol. Transfected cells were grown for 48 h at 37°C in 5% CO₂. Cells were then carefully washed and detached with 10 mM EDTA in PBS, counted, and seeded in replicates onto poly-L-lysine (Sigma-Aldrich) coated 48-well plates (BD Clontech) for 2–4 h at 37°C in 5% CO₂. Plates were blocked with 1% gelatin in TBS for 2 h at 37°C in 5% CO₂. Confluent monolayers in the 48 wells were then used for integrin cell surface expression and ligand-binding assays.

mAb binding studies

Triplicate or quadruplet wells containing confluent monolayers of transfected cells, ~48 h posttransfection, were incubated with mAbs LM142 (10 μ g/ml), AP3 (10 μ g/ml), LM609 (10 μ g/ml), 903 (10 μ g/ml), 44a (10 μ g/ml), TS1/18 (10 μ g/ml), or 107 (10 μ g/ml) in TBS buffer containing 0.1% gelatin and 0.02% sodium azide for 1 h at 4°C. Cells were washed and incubated with biotin-labeled rabbit anti-mouse Ab (Vectastain ABC-kit; Vector Laboratories) for 1 h at 4°C, washed three times, and subsequently fixed with 4% paraformaldehyde for 30 min at room temperature. After blocking, wells were developed using streptavidin-alkaline phosphatase (AP, Vectastain ABC-kit; Vector Laboratories) and p-nitrophenyl phosphate (Sigma-Aldrich) to generate a colored product. The enzymatic reaction was quenched with an equal volume of 2N NaOH, and OD (405) was measured. Binding to sham transfected cells conducted in parallel (which did not exceed 5% in all cases) was subtracted.

Immunoprecipitation

Transfected HEK 293 cells in six-well plates were extracted using radioimmunoprecipitation assay buffer (Boston Bioproducts) supplemented with 1 mg/ml DNase and a mixture of protease inhibitors (Complete; Roche Diagnostics). The detergent-soluble fraction was harvested upon centrifugation and immunoprecipitated using the anti- $\beta 3$ mAb AP3 or the

anti-CD18 mAb TS1/18 and Protein G Sepharose-4 (Amersham Pharmacia Biotech). Washed immunoprecipitates or whole cell lysates were separated by SDS-PAGE on a 4–15% gradient Tris-HCl gel (Bio-Rad) under non-reducing (for $\alpha V\beta 3$) or reducing (for CD11b/CD18) conditions and electroblotted onto polyvinylidene difluoride membranes (Bio-Rad). After blocking with 5% nonfat milk in 25 mM Tris-HCl (pH 7.4), 137 mM NaCl, 2.7 mM KCl (TBS, Boston Bioproducts), the respective membranes were probed with polyclonal anti- αV , anti- $\beta 3$ CBL479 mAb, anti-CD11b, or anti-CD18 polyclonal Abs (5). The anti- $\beta 3$ mAb CBL479 was probed with a polyclonal rabbit anti-mouse Ig Ab. Detection of proteins was performed using HRP linked anti-mouse or anti-rabbit Abs (eBioscience), and SuperSignal chemiluminescent kit (Pierce). The luminescent signal was detected using BioMax x-ray films (Eastman Kodak Company). Band intensity from Western blots was quantitated using the National Institutes of Health ImageJ software (22).

Ligand binding studies

Binding of the activation-dependent physiologic ligand iC3b, and the activation-independent ligand-mimetic mAb 107 to transiently transfected HEK 293 cells expressing CD11b/CD18 was performed in 48-well plates as previously described (5). Sheep erythrocytes coated with iC3b (EiC3b) were prepared following published protocols (5, 21). EiC3b binding was assessed by adding 150 μ l of EiC3b (60×10^6 /ml) to triplicate confluent wells in a total volume of 300 μ l followed by a 15 s spin at 500 rpm. After 35 min of incubation at 37°C, wells were carefully washed and subsequently fixed with 1% glutaraldehyde. Wells were developed using alkaline phosphatase and p-nitrophenyl phosphate as above and binding was quantitated. Specific binding was obtained by subtracting background binding of mock-transfected cells (typically <5% of total binding). Integrin cell surface expression was analyzed simultaneously using anti-CD11b 44a and anti-CD18 TS1/18 mAbs as described above. mAb binding results are reported as histograms representing mean \pm SD of triplicate or quadruplet wells. To correct for the effect of differences in surface expression on ligand or ligand-mimic binding, binding of TS1/18 to the WT receptor was considered 100. In the case of EiC3b and 107, relative binding to mutant vs WT integrin was normalized to that obtained in the presence of TS1/18 as previously described (16), according to the following formula: Percent binding to ligand = (percent mutant ligand binding/percent mutant TS1/18 binding) / (percent WT ligand binding/percent WT TS1/18 binding) \times 100.

Results

Design of Arg/Lys finger mutations in $\beta 3$ and CD18 subunits

Sequence and structure alignments show that the Arg/Lys finger is conserved in all mammalian β -chains except $\beta 4$, where it is replaced with an Ala (Fig. 1C). Conservation of the Arg/Lys finger suggests that this interfacial residue plays an important role in integrin heterodimerization and/or function. Using site-directed mutagenesis, we substituted WT $\beta 3$ residue Arg261 with Glu (to disrupt the cation- π interaction observed in the $\alpha V\beta 3$ crystal structure), and with Ala (replacing the charged residue with a hydrophobic residue). Because the structure of $\beta 2$ integrin heterodimers is currently not known, we generated a larger number of mutations in the integrin CD11b/CD18 model system by substituting the WT CD18 residue Lys252 (which corresponds to Arg261 in $\beta 3$) with Arg (a conservative substitution), Asp and Glu (to disrupt the cation- π interaction), Ala (replacing the charged residue with a hydrophobic residue), and Phe (introducing a large hydrophobic side chain).

Effect of $\beta 3$ Arg261 mutations on cell surface expression and heterodimer formation in $\alpha V\beta 3$

We assessed expression of the WT and mutant $\alpha V\beta 3$ on the surface of HEK 293 cells, which lack endogenous $\alpha V\beta 3$, but endogenously express the related $\alpha V\beta 1$ (23). A pcDNA3 expression vector containing a cDNA encoding full-length WT αV was co-transfected with plasmids encoding WT- or mutant $\beta 3$ into HEK 293 cells using Lipofectamine 2000. Mock transfectants were prepared using the empty pcDNA3 vector. Cell surface expression was measured in an ELISA-based assay, using the anti- αV mAb LM142, the anti- $\beta 3$ mAb AP3 and the $\alpha V\beta 3$ heterodimer-specific mAb LM609. The results, presented in Fig. 2A, show that the

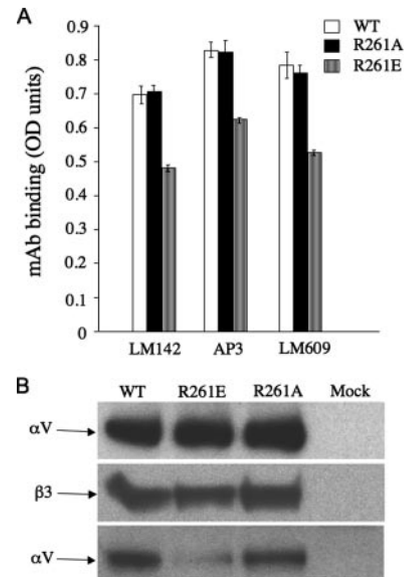


FIGURE 2. Effect of R261A and R261E on surface expression and $\alpha V\beta 3$ heterodimer formation. In this and subsequent figures, amino acids are indicated in the single letter code. **A**, Histograms showing expression of recombinant WT and mutant $\alpha V\beta 3$ on the cell surface of transfected HEK293 cells, probed with the anti- αV mAb LM142, the anti- $\beta 3$ mAb AP3 or the heterodimer specific mAb LM609. Each histogram represents mean \pm SD of triplicate determinations from three independent experiments. **B**, Western blots, following 4–15% gradient SDS-PAGE, developed with polyclonal anti- αV Ab (upper and lower), and anti- $\beta 3$ CBL479 mAb (middle). Equivalent amounts of αV - (upper) and WT or mutant $\beta 3$ -subunits (middle) are detected. Immunoprecipitation with anti- $\beta 3$ mAb AP3 from HEK 293 cells expressing WT or mutant receptors revealed much reduced levels of the associated αV -subunit in the R261E mutant, compared with WT or R216A receptors (one representative experiment of four conducted).

expression level of Arg261Ala mutant was comparable to WT, but the surface expression levels of the Arg261Glu mutant integrins was slightly but significantly reduced (to $\sim 75\%$ of WT).

To determine whether defects in heterodimer formation account for the reduction in cell surface expression, we conducted immunoprecipitation/Western blot studies on detergent-solubilized cell lysates from the transiently transfected HEK 293 cells. Probing Western blots of whole lysate from HEK 293 expressing WT or mutant integrins with polyclonal anti- αV or with anti- $\beta 3$ mAb CBL479 (following immunoprecipitation with anti- $\beta 3$ AP3 mAbs, due to weak reactivity of this mAb in Western blots) showed that equivalent amounts of normal-sized αV and $\beta 3$ -subunits are present in cells transfected with WT or mutant $\beta 3$ (Fig. 2B, upper and middle). Thus, the introduced substitutions did not affect the overall structure or expression levels of the respective subunit. However, immunoprecipitation of the $\beta 3$ subunits from WT- or mutant-expressing cells with mAb AP3 followed by Western blotting with anti- αV polyclonal Ab showed a significant reduction in association of the αV -subunit with Arg261Glu when compared with that associated with WT- or Arg261Ala $\beta 3$ -subunit (Fig. 2B, lower). These findings indicate that the Arg261Glu mutation significantly impairs formation of a stable $\alpha V\beta 3$ heterodimer.

Effect of CD18 K252 mutations on cell surface expression and heterodimer formation

To determine whether the above observations are also applicable to the αA -containing integrin subgroup, we examined whether

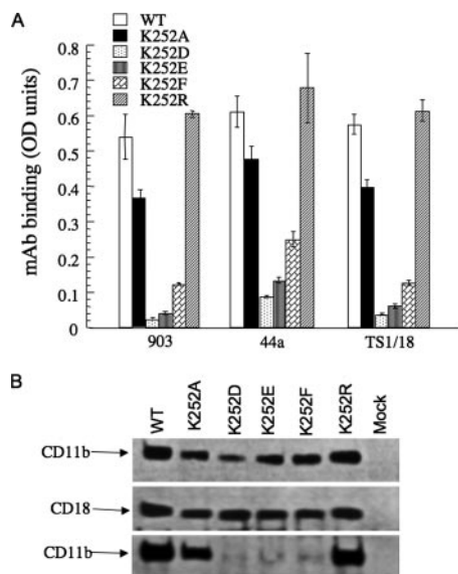


FIGURE 3. Effect of Arg/Lys mutations on surface expression and formation of integrin CD11b/CD18. *A*, Surface expression of WT and mutant CD11b/CD18. Histograms (mean \pm SD of triplicate determinations from a representative experiment, one of three performed) showing expression levels of WT and mutant CD11b/CD18 receptors in transfected HEK 293 cells. Surface expression was evaluated with the anti-CD11b mAbs 903 and 44a and the anti-CD18 mAbs TS1/18 mAbs. *B*, Western blots following 4–15% gradient SDS-PAGE developed with polyclonal anti-CD11b (*upper* and *lower*) or anti-CD18 Abs (*middle*) of whole cell lysate from HEK 293 cells expressing WT or mutant receptors (*upper* and *middle*) or anti-CD18 mAb immunoprecipitates probed with polyclonal anti-CD11b Ab (*lower*) (one representative experiment is shown of three conducted).

Lys252, the equivalent residue to Arg261 in $\beta 2$ integrins, is also important in formation of integrin CD11b/CD18. A pcDNA3 expression vector containing a cDNA encoding full-length human WT CD11b was cotransfected with plasmids encoding full-length human WT or mutant CD18 into HEK 293 cells (which do not endogenously express $\beta 2$ integrins), using Lipofectamine 2000. Mock transfectants were prepared using the empty pcDNA3 vector. Cell surface expression was also measured by ELISA, using the anti-CD11b mAbs 903 and 44a and the anti-CD18 mAb TS1/18. The results, presented in Fig. 3A, show that the expression level of the conserved Lys252Arg substitution in CD11b/CD18 was similar to that of WT, but expression of the other mutants receptors was reduced to various degrees, ranging from $\sim 75\%$ of WT in Lys252Ala mutant to $<10\%$ in each of the remaining mutants.

Immunoprecipitation/Western blot analyses were then used to evaluate the effect of each mutation on formation of the heterodimer more directly, using detergent-extracted cell lysates from transfected HEK 293 as described (5). Detergent extracts from HEK 293 cells transfected with mutant CD18 (together with WT CD11b) expressed a normal-sized CD11b-subunit, as expected (Fig. 3B, *upper*), as well as equivalent amounts of normal-sized WT and mutant CD18-subunit (Fig. 3B, *middle*), indicating that the introduced substitutions did not affect the overall structure or level of mutant CD18. Immunoprecipitation of the CD18 subunit with TS1/18 mAb followed by Western blotting with polyclonal anti-CD11b showed that both Lys252Ala and Lys252Arg mutant CD18 associated with WT CD11b to equivalent degrees as that of WT CD18 (Fig. 3B, *lower*). However, association of Lys252Asp, Lys252Glu, and

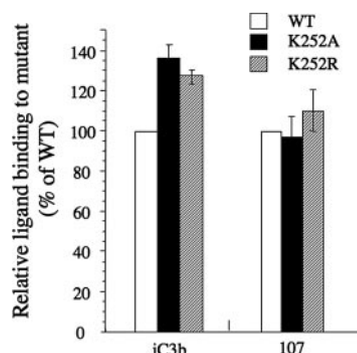


FIGURE 4. Effect of K252A and K252R on ligand binding. *Left*, Histograms showing the binding of the physiologic ligand iC3b or the ligand mimic 107 to mutant CD11b/CD18 relative to that of WT (placed at 100), after correcting for receptor expression using mAb TS1/18. Each histogram represents mean \pm SD of triplicate (for iC3b) or quadruplet (for 107) determinations from a representative experiment (of three performed). Details of the methods used are described in *Materials and Methods*.

Lys252Phe mutant CD18 with WT CD11b was minimal, consistent with the cell surface expression analysis. Thus, charge reversal at the conserved Arg/Lys finger (as in Lys252Asp and Lys252Glu mutants) or introduction of a bulky hydrophobic residue (as in the Lys252Phe mutant) largely abrogates association of the CD11b and the CD18 subunits, whereas the conservative substitution Lys252Arg and the least sterically disruptive substitution Lys252Ala are well tolerated.

Impact of Lys252Ala and Lys252Arg substitutions in CD18 on integrin function

The Arg/Lys finger is just two residues downstream of loop 3 (Fig. 1C), one of the three loops that form the ligand binding MIDAS of the βA domain (1). This structural arrangement is analogous to the activating “switch II” region in G proteins, which is similarly linked to the metal ion binding catalytic site in $G\alpha$ (1). This structural analogy suggests that the packing of the βA domain against the propeller may also affect integrin activation. We examined the impact of the Lys to Arg or to Ala substitutions on the ability of recombinant CD11b/CD18 heterodimer to bind the physiologic ligand iC3b (16, 24). WT CD11b/CD18 expressed on HEK 293 is partially activated, enabling it to bind the activation-sensitive ligand iC3b in the presence of physiologic divalent cations (Ca^{2+} plus Mg^{2+} , each at 1 mM) (25). This explains the intrinsic binding of WT CD11b/CD18-expressing HEK 293 to iC3b (Fig. 4). Lys252Ala and Lys252Arg substitutions in integrin CD11b/CD18 induced a further increase in binding to iC3b by 1.4 and 1.3-fold, respectively, above that of the WT receptor. In contrast, binding of the activation-insensitive ligand-mimetic mAb 107 (17) was similar in both mutant receptors to that of WT (Fig. 4).

Discussion

The results presented in this communication validate the model that the Arg/Lys finger plays a important role in formation of a stable integrin heterodimer, and suggest that as in G-proteins, the βA /propeller interface may also affect integrin activation.

Charge reversal of the Arg/Lys finger led to a reduction in surface expression of both the αA -lacking and containing integrins $\alpha V\beta 3$ and CD11b/CD18, respectively. Cell surface expression of $\alpha V\beta 3$ was less affected by the Arg261Glu substitution in $\beta 3$ than CD11b/CD18 was by the Lys252Glu in $\beta 2$ (compare Figs. 2A and

3A), although the detrimental effect of either substitution on stability of the respective heterodimer was comparable in the immunoprecipitation analysis (compare *lower* in Figs. 2B and 3B). These findings suggest that cell surface expression is a less sensitive measure of integrin heterodimer stability. A similar observation was made in analysis of integrin $\beta 3$ L262P mutant (11), which was also expressed on the surface of HEK 293 cells, but its effect on heterodimer stability was readily detected under the more stringent immunoprecipitation analysis. These findings may also reflect a generally lower affinity of the αA -containing CD11b subunit to CD18 in the WT integrin than that of αV -subunit for $\beta 3$. It remains possible that HEK 293 cells, which constitutively express the related $\alpha V\beta 1$, may possess the cellular machinery to efficiently package and export the mutant $\alpha V\beta 3$ to the cell surface, but not mutant CD11b/CD18 heterodimers that are normally expressed exclusively in a leukocyte context.

The $\beta 4$ -subunit, which pairs only with $\alpha 6$ to form a functional heterodimer, is the only integrin β -subunit lacking the Arg/Lys finger (Fig. 1C). This finger is replaced with an alanine, which is followed by an Asn-Val insertion unique to this subunit. Although there are caveats associated with homology-based modeling, we find that the size and properties of the interface between Ala197AsnVal in $\beta 4$ and the aromatic cup of $\alpha 6$ is similar to that at the $\alpha V/\beta 3$ interface, though the main interaction is now mediated by van der Waals' forces. However, the dipeptide insertion may create a slightly larger interfacial contact with $\alpha 6$, thereby mitigating the reduction in ΔG from the missing cation- π interaction.

Substitution of K252 with Arg or with Ala in the CD11b/CD18 integrin resulted in a significant increase in binding to physiologic ligand by the recombinant integrin (Fig. 4). This increase could result from changes in integrin affinity and/or avidity. Recent evidence suggests that the increase in ligand-binding induced by inside-out signaling in the $\beta 2$ -integrin CD11a is primarily due to affinity modulation rather than avidity changes (26). The proximity of the Arg/Lys finger to the ligand-binding loop 3 of MIDAS, favors an intramolecular (affinity) rather than an intermolecular (clustering/avidity) effect. In contrast to G-proteins, where conformational activation results in complete dissociation of the G α - and G β -subunits, enhanced ligand binding by these two substitutions was elicited with little or no detectable disruption of the integrin heterodimer (Figs. 2B and 3B). These findings suggest that integrin-ligand interactions may be regulated by more subtle allosteric changes at the propeller/ βA interface the nature of which remain to be structurally-defined.

The present results also provide new insights into how some naturally-occurring mutations, which occur most frequently in the βA domains of CD18- (27) and $\beta 3$ - (28) integrin subunits, lead to life-threatening bacterial infections and thrombasthenia, respectively. For example, the naturally occurring mutations G251R in CD18 (equivalent to G260 in $\beta 3$) (29) and L262P in $\beta 3$ (equivalent to L253 in CD18) (11), known to prevent formation of integrin heterodimers, lie immediately after the critical Arg/Lys finger residue in the 3_{10} -helix of the respective integrin. Data presented in this study suggest that these mutations act in part by disrupting the contact of the Arg/Lys finger with the cup-like structure of the propeller. If this interpretation is correct, then mutations in the aromatic cup residues of the propeller domain should likewise impair heterodimer formation. This has recently been borne out: a naturally-occurring Phe171Cys mutation in αIIb (equivalent to Phe159 in αV) (Fig. 1B) found in patients with inherited thrombasthenia, impairs formation of the $\alpha IIb\beta 3$ heterodimer and its surface expression in platelets (30). The $\alpha V\beta 3$ crystal structure shows that Arg261

is closest to Phe159 of the α -subunit, and contributes the primary cation- π bond; its replacement with cysteine is expected to destabilize the Arg/Lys finger contact with the cup structure. Similarly, a mutation of another aromatic cup residue Phe289 in αIIb (corresponding to the conserved Phe278 in αV) also leads to loss of surface expression and heterodimer formation (31). Other naturally occurring mutations in the cup structure (32) may likewise compromise stable contact with the Arg/Lys finger and hence integrin heterodimer formation.

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Disclosures

The authors have no financial conflict of interest.

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