Integrin cell-adhesion receptors transduce signals bidirectionally across the plasma membrane via the single-pass transmembrane segments of each α and β subunit. While the β3 transmembrane segment consists of a linear 29-residue α-helix, the structure of the αIIb transmembrane segment reveals a linear 24-residue α-helix (Ile-966–Lys-989) followed by a backbone reversal that packs Phe-992–Phe-993 against the transmembrane helix. The length of the αIIb transmembrane helix implies the absence of a significant transmembrane helix tilt in contrast to its partnering β3 subunit. Sequence alignment shows Gly–991–Phe–993 to be fully conserved among all 18 human integrin α subunits, suggesting that unusual structural motifs are prototypical for integrin α subunits. The αIIb transmembrane structure demonstrates a level of complexity within the membrane that is beyond simple transmembrane helices and forms the structural basis for assessing the extent of structural and topological rearrangements upon αIIb–β3 association, i.e. integrin transmembrane signaling.

Integrins are a major class of adhesion receptors that mediate cell migration and extracellular matrix assembly, as well as inflammation, thrombosis, and tumor metastasis. In addition to responding to signals that originate inside the cell and regulate receptor affinity, integrins can respond to the binding of extracellular ligands by activating intracellular signaling pathways. Integrins are noncovalent heterodimers with large extracellular and small intracellular domains that are connected by single-pass transmembrane (TM) segments. An extensive set of experiments provides evidence that signals are transduced bidirectionally across the plasma membrane via the TM segments of the α and β subunits. Specifically, the inactive receptor state is stabilized by the hydrophobic packing of the TM helices and adjacent electrostatic αIIb–β3 interactions, whereas integrin activation ensues from the separation of the TM segments (2, 4–8). However, the precise structural and topological processes underlying integrin inside-out and outside-in signaling events are currently unknown.

Single-pass TM segments likely traverse the membrane as α-helices. The recently determined structure of the integrin β3 TM segment reconstituted in small bicelles fulfills this expectation, but the TM helix is not restricted to residues between the first charged residues, Asp-692 and Lys-716, on the extracellular side and, respectively, Lys-716 on the intracellular side, encoding a helix tilt relative to the membrane. Apparently, this allows Lys-716 to snorkel its side chain out of the hydrophobic lipid core. Integrin α subunits exhibit a similar pattern of hydrophobicity; a lysine residue constitutes the first charged residue on the intracellular side and is followed by four mostly apolar residues (see Fig. 1). Nevertheless, the structure of the αIIb TM segment reported herein shows that the TM helix ends at Lys-989 and that the following four residues loop back to immerse Phe-992–Phe-993 in the hydrophobic lipid core. Moreover, the 24-residue TM helix can immerse in a typical bilayer without a significant helix tilt. The αIIb TM structure reveals an unexpected level of complexity within the integrin TM region that appears crucial for understanding the association with its partnering β3 TM segment. Furthermore, the structure highlights that single-pass TM segments are not restricted merely to helical conformations.

**EXPERIMENTAL PROCEDURES**

*Integrin αIIb Expression and Purification.*—Synthetic oligonucleotides coding for human integrin αIIb residues Ala–958–Pro–998 were assembled by PCR and subcloned into the pET-44 expression vector (Novagen, Inc.) with the third IgG-site. Expression and purification proceeded as described for the β3 TM peptide except that the αIIb peptide was cleaved from the fusion protein using tobacco etch virus protease cleavage site. Expression and purification proceeded as described for the β3 TM peptide except that the αIIb peptide was cleaved from the fusion protein using tobacco etch virus protease cleavage site. Introduction of a high-affinity Cu^{2+}-binding domain, Gly-Gly-His ATRCUN motif, 10−11, was introduced by PCR for residues corresponding to αIIb(957–959). GGH-αIIb peptide expression and purification proceeded in the same manner as with αIIb. Isotope labeling
was achieved by supplying 15N-NH4Cl and 13C-D-glucose and culturing in 99 and 50% D2O for highly and moderately deuterated peptide, respectively.

**NMR Sample Preparation**—αIIb peptide concentration was measured in acetonitrile-water solution by UV spectroscopy (ε280 nm = 16,500 M⁻¹ cm⁻¹), and defined amounts of peptide were freeze-dried. Peptide was taken up in 320 μl of bicelle solution to give an αIIb peptide concentration of ~0.6 mM. The bicelle solution was comprised of 105 mM long-chain lipids and 350 mM 1,2-dihexanoyl-sn-glycero-3-phosphocholine (DHPC), 25 mM HEPES-NaOH, pH 7.4, 6% D2O, 0.02% w/v NaN₃. As long-chain lipids, 1-palmitoyl-2-oleoyl-sn-glycero-3-phosphocholine (POPC) and 1,2-dimyristoyl-sn-glycero-3-phosphocholine (PODG) were used, respectively. No significant H-N αIIb chemical shift changes were detected between the two lipid environments (supplemental Fig. 1), showing that equivalent structures were adopted. Supplemental Fig. 2 illustrates the spectral quality obtained.

To align the αIIb-bicelle complex relative to the magnetic field stretched, negatively charged polyacrylamide gels were used (12, 13). A 320-μl gel was polymerized in 150 mM Tris-HCl, pH 8.0, from a 4.2% w/v solution of acrylamide, 2-acrylamido-2-methyl-1-propanesulfonate (AMPS), and bisacrylamide with a monomer-to-cross-linker ratio of 49:1 (w/w) and a molar ratio of 94:6 of acrylamide to AMPS. For easier comparisons between gels, AMPS was counted as acrylamide in the given weight ratios (14). The gel was dialyzed overnight in 50 ml of 50 mM 1,1,2,2-tetrafluoroethane (HFTE) solution for 24 h and transferred into an open-ended NMR tube, as described (15). A 3H splitting of 1.5 Hz was observed.

**NMR Spectroscopy**—NMR experiments were carried out on a cryoprobe-equipped Bruker Avance 700 spectrometer at 35 °C except for the gel-aligned sample, which was measured at 40 °C. Data were processed and analyzed with the nmrPipe package (16) and CARA. Throughout all experiments, the solvent was kept at +1H whenever possible (17, 18), and TROSY-type H-N detection was used (19).

Highly deuterated αIIb peptide (99% D₂O cultures) was used for the following procedures: HN, N, Cα, Cβ, and Cγ assignments from HNCA, HN(CA)Cβ, HNCO, and HN(CA)CO experiments. J_CCY and J_NCY couplings for aromatic and aliphatic residues were obtained from quantitative J-correlation spectroscopy (20, 21) with dephasing times of 50 and 100 ms, respectively. J_NH, J_CαCβ, J_CαCγ, and J_NH+1DIH coupling constants were determined from 1HODized HNCO experiments (22) and from quantitative J-correlation HNCO experiments (23, 24) on isotropic and aligned samples, respectively. [1H]15N NOE measurements were carried out using 5 s of presaturation preceded by a recycling delay of 4 s for the NOE experiment and a 9-s recycle delay for the reference experiment. To measure HN•H•N and HN•H•D NOEs, a 13C-edited NOESY spectrum (150-ms mixing time) was recorded.

Moderately deuterated αIIb peptide (50% D₂O cultures) was used for the detection of side-chain NOEs. Moreover, deuterated lipids, 1,2-dihexanoyl(d25)-sn-glycero-3-phosphocholine-1,2,2-δ5-N,N,N-trimethyl-d9, and 1,2-dimyristoyl(d45)-sn-glycero-3-phosphocholine-1,2,2-δ5-N,N,N-trimethyl-d9 were employed. [1H]13C-edited NOESY spectra, optimized for the detection of aromatic and aliphatic nuclei, respectively, and a 15N-edited NOE spectrum were recorded with mixing times of 130 ms. Aliphatic side-chain assignments were achieved using an (H)CCH-COSY experiment (25). Full side-chain assignments of the two Phe and partial assignments of the three Trp side chains resulted directly from CT-HSQC spectra and their NOE patterns (26).

**Structure Calculations**—Structures of the well folded Ile-966–Lys-994 residues (see Fig. 4A) were calculated by simulated annealing, starting at 3000 K using the program XPLOR-NIH (27). The peptide termini were represented by random-coil conformations. Besides standard force field terms for covalent geometry (bonds, angles, and improper torsions) and nonbonded contacts (van der Waals repulsion), the following experimental and empirical potentials were included. The difference between predicted and experimental residual dipolar couplings was described by a quadratic harmonic potential. Dihedral angle restraints were implemented using a quartic square-well potential and merely aided in the conversion of the residual dipolar coupling restraints. φ, ϑ backbone dihedral angle constraints were extracted from N, Cβ, Cγ, and Cδ chemical shifts using the program TALOS (28). χ₁, χ₂ side-chain angle restraints were derived from the J_CCY and J_NCY coupling constants (supplemental Table 2). NOE interproton distance restraints were referenced to the linear α-helical structure of Ile-966–Lys-989 and were incorporated using a quadratic square-well potential. A backbone-
backbone hydrogen-bonding potential (29) was used. A torsion angle potential of mean force (30), which was modified to implement the preferred rotamers in transmembrane helices (31, 32), was employed. Moreover, the higher side-chain rotamers (χ2-χ4) of Lys-989 were selected to extend toward the C terminus, i.e. snorkeling its side chain out of the membrane core. A total of 20 structures were calculated. The structural statistics are summarized in supplemental Table 1. The structures have been deposited in the Protein Data Bank (accession number 2k1a) together with the energy-minimized average structure.

RESULTS AND DISCUSSION

Structure Determination and Description—The structure of the αIIb TM segment reconstituted in small bicelles was determined by multidimensional heteronuclear NMR spectroscopy, making use of 2H/13C/15N-labeled peptide encompassing residues Ala-958–Pro-998 (Fig. 1). Bicelles provide a small, symmetric lipid bilayer whose rim is stabilized by short-chain lipids (33, 34), and they appear well suited for the study of single-pass TM segments (9). Under the experimental conditions employed, the presence of a paramagnetic tag on αIIb was not sensed by untagged αIIb peptides, and no significant concentration-dependent chemical shift changes were observed (supplemental Fig. 3), clearly showing that the αIIb structure is not affected by any homo-oligomerization tendency. To validate the choice of bicelle lipids, the effects of lipid headgroup and tail length on αIIb spectral parameters were examined. Between glycerophosphatidylcholine-based lipids with 16/18 and 14/14 hydrocarbon tails, no significant 1H/N and 15N chemical shift changes were detected (supplemental Fig. 1). Moreover, the introduction of lipids carrying phosphatidylserine headgroups had no effects on the αIIb backbone structure, as evidenced by the invariance of 13Cα secondary shifts (Fig. 2A), which correlate with backbone torsion angles (35). The newly introduced lipids were sensed predominantly by αIIb residues on the extracellular face (Fig. 2, B and C). Thus, the detected αIIb structure is stable in lipid environments that resemble the composition of its native platelet membrane (36).

The αIIb TM structure was calculated from an extensive set of bond vector orientation, internuclear distance, and torsion angle constraints (supplemental Tables 1 and 2). For the ensemble of 20 calculated simulated annealing structures (Fig. 3A), coordinate precisions of 0.22 Å were obtained for the backbone-heavy atoms of the structured αIIb residues (Ile-966–Lys-994). The structure is characterized by a linear α-helix for Ile-966–Lys-989 followed by a reversal of backbone direction that conversely immerses Phe-992–Phe-993 toward the lipid hydrocarbon core of the intracellular membrane face (Fig. 3, D and E). By exhibiting increased protection from the paramagnetic Mn2+/EDDA2− agent present in the aqueous phase as well as from H2O exchange, the increased Phe-992–Phe-993 immersion was directly detectable within the bicelle membrane model (Fig. 4, B and C). Phe-993 packs against the TM helix, as evidenced by close interproton distances between its aromatic ring and nuclei from Trp-988, Leu-985, and Val-984 and its well defined χ1, rotameric state (supplemental Table 2). The packing of Phe-992 is characterized by hydrophobic contacts to Val-990

FIGURE 2. Effect of bicelle lipid composition on αIIb chemical shifts. The substitution of one-third of bicellar POPC lipids with 1-palmitoyl-2-oleoyl-sn-glycero-3-[phospho-L-serine] (POPS) lipids has no effects on the 13Cα secondary shifts (A) and, thus, αIIb backbone structure. Secondary 13Cα chemical shifts, defined as the difference between the observed and tabulated random-coil 13Cα shift of a residue, correlate with the underlying backbone conformation (35). Pro-965 and Pro-998 limit the backbone conformation of their preceding residues to extended conformations (42), as evidenced by their large negative 13Cα secondary shifts. 13Cα secondary shifts are reported relative to the random-coil values of Ref. (35) but have not been corrected for the 0.5-ppm isotope shift resulting from perdeuteration of the carbon-attached hydrogens. B and C, the introduction of POPS is sensed predominantly by the αIIb backbone 1H and 15N nuclei on the extracellular αIIb TM helix side. Changes in chemical shifts for these nuclei types are sensitive to changes in the chemical environment as well as to structural changes (43). However, overall 1H and 15N chemical shift changes are relatively small. The transmembrane helix (Ile-966–Lys-989) borders and the Lys-994–Arg-995 transition are marked by blue dashed lines in all panels.
and Met-987; however, overall packing appears less defined, leading to an averaged $\chi_1$ rotameric state (supplemental Table 2). C-terminal to Lys-994, fast time scale backbone dynamics increase significantly (Fig. 4A), leading to dynamically unstructured conformations, with Arg-997 forced into extended backbone conformations by Pro-998 (Fig. 2A). Incidentally, the extracellular side of the TM segment is also strongly affected by a proline residue (Fig. 3, B and C); Pro-965 restricts Ile-964 to extended conformations, as evidenced by its large negative $^{13}$C\alpha secondary shift (Fig. 2A), which may benefit dynamic interactions with the hydrophobic lipid tails (Fig. 3, B and C). Backbone dynamics increase steadily from the start of the TM helix at Ile-966 to the N terminus (Fig. 4A).

Structural similarities to the integrin $\alpha$IIb TM segment were found in the Protein Data Bank for a peptide (1 nkz, chain F) of the light-harvesting complex II from *Rhodopseudomonas acidophila* (37) (supplemental Fig. 5A). Somewhat more remotely related is a TM helix of cytochrome c oxidase from *Thermus thermophilus* (1xme, chain A) (38) (supplemental Fig. 5B).

**Membrane Embedding and Relationship to Other Integrin $\alpha$ Subunits**—Sequence alignment of all 18 human integrin $\alpha$ subunits (Fig. 1 and supplemental Fig. 4) suggests that the $\alpha$IIb extra- and intracellular transitions to the TM helix are characteristic of $\alpha$ subunits. At the extracellular helix border, half of all $\alpha$ subunits exhibit a proline residue preceded by one that is hydrophobic. The Gly-Phe-Phe motif on the intracellular side is the only fully conserved tri-residue sequence element among integrin $\alpha$ TM segments, suggesting that the observed backbone reversal and Phe-Phe membrane immersion are general structural features of integrin $\alpha$ subunits. These transitions are fundamentally different from the $\beta$3 TM segment, where irregular structure and helix fraying characterized the transitions to the extra- and intracellular domains, respectively (9). Moreover, in contrast to the 29-residue $\beta$3 TM helix, the 5-residue-shorter $\alpha$IIb TM helix does not require any tilt within a typical lipid bilayer (Fig. 5). The structural complexity of the Gly-991–Phe-993 motif and ensuing complicated $\text{Mn}^{2+}$EDDA$^{2-}$ protection make it difficult to determine the TM helix center.
Integrin αIIb Transmembrane Structure

A

\[ \frac{I}{I_0} \]

Residue No.

B

\[ \frac{I}{I_0} \]

Residue No.

C

\[ \frac{I}{I_0} \]

Residue No.

FIGURE 4. αIIb backbone dynamics, protection from paramagnetic Mn\(^{2+}\)EDDA\(^{-2}\) and H\(^{2+}\)-H\(_2\)O chemical exchange. A, pico- to nanosecond dynamics, correlative to backbone order (44), is evaluated in the form of heteronuclear \([1\text{H}]-^{15}\text{N}\) NOE values. Low values, as seen for the peptide termini, correspond to dynamically unstructured residues. B, signal broadening due to paramagnetic relaxation enhancement arising from the presence of net neutral Mn\(^{2+}\)EDDA\(^{-2}\) in the aqueous phase. The normalized ratio of H-N TROSY signal intensities in the presence and absence of 1 mM Mn\(^{2+}\)EDDA\(^{-2}\), \(I/I_0\), is used to quantify signal broadening for each residue. C, exchange of the backbone amide protons (\([1\text{H}]-^{15}\text{N}\)) with water, depicted as the ratio of H-N TROSY signal intensities obtained with and without selective solvent inversion, \(I/I_0\), at pH 8.4 and 25 °C. The solvent was inverted 100 ms prior to the start of the pulse sequence, and radiation damping was suppressed by the application of a weak gradient pulse during this period. An \(I/I_0\) value of −1 corresponds to the exchange of the \([1\text{H}]-^{15}\text{N}\) nucleus of a residue with water in all αIIb molecules present, whereas a value of +1 denotes the absence of any exchange. Resonances that were absent at pH 8.4, when compared with pH 7.4, were assigned \(I/I_0\) values of −1. The transmembrane helix (Ile-966–Lys-989) borders and the Lys-994–Arg-995 transition are marked by dashed lines in all panels.

within the bicelle (Fig. 4B); in the absence of this information, the helix is assumed to be centered in the membrane (Fig. 5).

The lipid tail-to-headgroup transition on the extracellular side is estimated to take place before the TM helix commences, i.e. between Pro-965–Ile-966 (Fig. 1). In this region, protection from paramagnetic Mn\(^{2+}\)EDDA\(^{-2}\) is significantly reduced (Fig. 4B). This transition is also in agreement with the Pro-998–Leu-999 transition detected for the α5 subunit (39), which is similar to αIIb (Fig. 1). It is also noted that, at the extracellular helix border, no sharp hydrophobic-hydrophilic transition takes place (Fig. 1), which may result in a border variation between α subunits (39). The membrane borders depicted in Fig. 1 should therefore be considered as minimal for monomeric α subunits. On the intracellular side, the αIIb TM segment is expected to leave the lipid core after Phe-993 at the earliest. Lys-994 dynamics and protection are still somewhat high (Fig. 4), but, in conjunction with glycosylation mapping studies of integrin α2 and α5 subunits (40), the membrane border is estimated to lie between Phe-993–Lys-994. By virtue of a higher sequence homology on the intracellular TM face (Fig. 1 and supplemental Fig. 4), this transition appears to be more uniform among integrin α subunits. Finally, it should be noted that, on the basis of the Mn\(^{2+}\)EDDA\(^{-2}\) protection data of the αIIb and β3 TM segments (9) (Fig. 4B), immersion of the αIIb TM helix in the membrane is nevertheless slightly deeper than is the case for the β3 TM helix, as depicted in Fig. 5.

A stretch of continuous α-helix for residues Val-990–Phe-993 appears less preferable than the observed conformation for three reasons. First, the two Phe-992–Phe-993 aromatic rings prefer to immerse in the hydrophobic lipid tail environment instead of in the electrostatically complex headgroup interface (41). Second, any TM helix extension would necessitate a helix tilt, which will require Phe-992–Phe-993 to reposition further outward into the headgroup region because they would come to lie on the same helix side as snorkeling Lys-989. In contrast, the β3 subunit has Ile-719 and Thr-720 following Lys-716 as the transmembrane helix (39, 40). It appears likely that the highly conserved Gly-988–Phe-Phe motif has additional functional meaning. Thus, as a paradigm for integrin α and β subunits, the four residues following αIIb(Lys-989) would equally well cause a short helix formation, and the absence of a tilt, whereas the five residues following β3(Lys-716) create a bias for a long TM helix and a substantial tilt (Fig. 5).

The use of the Gly-Phe-Phe sequence motif is remarkable since, for example, the introduction of a second charged residue after αIIb(Lys-989) would equally well cause a short αIIb TM helix (39, 40). It appears likely that the highly conserved Gly-Phe-Phe motif has additional functional meaning.

Implications for Integrin αIIb β3 Association—Integrin bidirectional signaling proceeds with changes in α-β TM helix association (2–4). The individual αIIb and β3 TM structures, which correspond to activated integrin conformations (2–4), therefore lay the foundation for evaluating the extent of structural and topological rearrangements upon αIIb-β3 association, i.e. integrin TM signaling. Moreover, the structures show that, in the activated receptor, the α-β extracellular domains connect to their respective transmembrane segments at different membrane crossing angles (Fig. 5).
The monomeric depiction likely orientations of the \( \alpha \)IIb and \( \beta \)3 TM segments, shown in green and cyan, respectively, relative to the indicated functional groups of a lipid bilayer composed of 1,2-dioleoyl-sn-glycero-3-phosphocholine (45), which is closely related to the lipids employed herein. The side chains of Val-696, Leu-697, Val-700, Gly-708, Trp-715, and Lys-716 are shown for the \( \beta \)3 subunit; for \( \alpha \)IIb, the side chains of Trp-968, Gly-672, Gly-976, Thr-981, Trp-988, Lys-989, Phe-992, and Phe-993 are depicted (2, 8). Initial \( \alpha \)IIb-\( \beta \)3 association is compatible with the individual \( \alpha \)IIb and \( \beta \)3 structures; the final dimer structure will involve additional structural rearrangements.

Alanine substitution of Phe-992 or Phe-993 leads to receptor activation (5), showing that interactions of these residues with the \( \beta \)3 subunit are required for maintaining the inactive state. By either interacting with the \( \beta \)3 subunit in the conformation of the monomeric \( \alpha \)IIb TM segment or shifting Val-990–Phe-993 to another, for example helical, conformation, the Gly-991–Phe-993 structural motif is central to determining the relative orientation and topology not only in the activated but critically reading the manuscript.
Integrin αIIb Transmembrane Structure

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