Single-molecule measurements of the effect of force on Thy-1/αvβ3-integrin interaction using nonpurified proteins

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ABSTRACT Thy-1 and αvβ3 integrin mediate bidirectional cell-to-cell communication between neurons and astrocytes. Thy-1/αvβ3 interactions stimulate astrocyte migration and the retraction of neuronal prolongations, both processes in which internal forces are generated affecting the bimolecular interactions that maintain cell–cell adhesion. Nonetheless, how the Thy-1/αvβ3 interactions respond to mechanical cues is an unresolved issue. In this study, optical tweezers were used as a single-molecule force transducer, and the Dudko-Hummer-Szabo model was applied to calculate the kinetic parameters of Thy-1/αvβ3 dissociation. A novel experimental strategy was implemented to analyze the interaction of Thy-1-Fc with nonpurified αvβ3-Fc integrin, whereby nonspecific rupture events were corrected by using a new mathematical approach. This methodology permitted accurately estimating specific rupture forces for Thy-1-Fc/αvβ3-Fc dissociation and calculating the kinetic and transition state parameters. Force exponentially accelerated Thy-1/αvβ3 dissociation, indicating slip bond behavior. Importantly, nonspecific interactions were detected even for purified proteins, highlighting the importance of correcting for such interactions. In conclusion, we describe a new strategy to characterize the response of bimolecular interactions to forces even in the presence of nonspecific binding events. By defining how force regulates Thy-1/αvβ3 integrin binding, we provide an initial step towards understanding how the neuron–astrocyte pair senses and responds to mechanical cues.

INTRODUCTION Cell adhesion, the ability of a cell to bind either to another cell or to the extracellular matrix (Gumbiner, 1996), plays an important role in cell communication and the regulation of many biological processes, including cell proliferation, migration, and survival (Khalili and Ahmad, 2015). In the CNS, cellular adhesion between neurons and astrocytes is critical to regulating neuronal functions, both under physiological conditions as well as in response to CNS damage (Benarroch, 2005). Astrocytes are the most abundant type of glial cells in the CNS and exceed more than fivefold the quantity of neurons (Sofroniew and Vinters, 2010). A single astrocyte can contact up to 105 synapses (Bushong et al., 2002) and hundreds of dendrites (Halassa, Fellin, et al., 2007); hence, a wide variety of molecular interactions mediate the communication between neurons and glial cells.

Our group has previously reported that the glycosyl phosphatidylinositol (GPI)-anchored Thy-1, an abundant neuronal cell adhesion protein from the immunoglobulin (Ig) superfamily...
(Leyton and Hagood, 2014), acts as a receptor for the astrocytic αβ3 integrin (Leyton et al., 2001; Hermosilla et al., 2008). This heterodimeric integrin is upregulated in astrocytes after CNS injury due to increased inflammation (Ellison et al., 1999). We have also provided evidence for a direct Thy-1/αvβ3 integrin interaction mediated by the integrin-binding site (RLD) in Thy-1 using surface plasmon resonance analysis with the functional Fc-tagged fusion proteins Thy-1-Fc and αβ3-Fc (Hermosilla et al., 2008). Furthermore, we have described how the Thy-1/αvβ3 integrin interaction mediates bidirectional neuron-to-astrocyte communication, inducing cell migration in astrocytes (Avalos, Valdivia, et al., 2009; Kong, Munoz, et al., 2013) and promoting an inhibition of neurite outgrowth and retraction of already established neuronal processes in neurons (Herrera-Molina et al., 2012; Maldonado et al., 2016). Therefore, this interaction might play a key role in the responses of both neurons and astrocytes after CNS injury.

Adhesion proteins that maintain cell–cell communication, such as Ig superfamily members and integrins, are constantly exposed to mechanical forces generated as a consequence of different cellular functions (Evans and Calderwood, 2007; Strunz et al., 2000; Liu et al., 2015). In the wound healing process, for instance, cells tug on each other during migration and exert traction forces on the interactions that maintain cell–cell adhesion (Ananthakrishnan and Ehrlicher, 2007; Rakshit and Sivasankar, 2014). As mentioned, astrocyte migration is a process triggered by the Thy-1/αvβ3 integrin interaction. In cell migration, the polymerization of actin filaments and the sliding of the motor protein myosin II over these filaments generate contractile forces. In turn, these forces pull on the cell body to promote forward motion and retraction of the cell rear (Pollard and Borisy, 2003; Fournier et al., 2010). Similarly, neurite retraction also occurs due to increased contraction of the cortical actin cytoskeleton (Kranenburg et al., 1997; Govek et al., 2005; Maldonado et al., 2016). Therefore, both cell migration and neurite retraction generate internal forces exerted to protein–protein interactions that maintain cell–cell adhesion, as is the case of neuronal Thy-1 and astrocytic αβ3 integrin binding.

Force can tilt the energy landscape of the dissociation pathway, alter the height of the energy barrier, and modify unbinding kinetics, that is, forces that regulate molecular interactions by modulating the off-rate (koff) (Bell, 1978; Dembo et al., 1988). Based on how bimolecular interactions respond to mechanical forces, bonds can be classified as either slip bonds, where force shortens bond lifetime (accelerate dissociation); catch bonds, where force prolongs bond lifetime (slow down dissociation); or ideal bonds, where bond lifetime is independent of force (Dembo et al., 1988; Zhu, 2014). Mechanical cues from the surrounding environment clearly influence binding properties and, consequently, modify the effects of cellular signaling, as triggered by adhesion proteins such as Thy-1 and αβ3 integrin. Bidirectional neuron-to-astrocyte communication in the CNS could therefore also be affected. Nonetheless, how mechanical forces may modulate kinetic parameters of the interaction between Thy-1 and αβ3 integrin has never been explored. Therefore, molecular force spectroscopy was used in the present study to characterize the mechanical properties of the Thy-1/αβ3 integrin interaction at the single-molecule level. This methodology has been employed to characterize protein–protein interactions in response to forces, resulting in assessments of bond strength, the dissociation process energy landscape, and the lifetime of bimolecular interactions (Yuan et al., 2000; Stangner et al., 2013).

In the current study, optical tweezers and force-ramp assays were used to obtain rupture force data for the Thy-1/αβ3 integrin interaction. The kinetic parameters of the force-induced Thy-1/αβ3 dissociation were determined using the Dudko-Hummer-Szabo (DHS) model (Dudko et al., 2008). Previous studies using molecular force spectroscopy to characterize the effect of mechanical forces on bimolecular interactions have utilized purified proteins to reduce the possibility of nonspecific interactions (Zhang et al., 2004; Litvinov et al., 2005; Stangner et al., 2013). However, protein purification may represent a limiting factor. To overcome this limitation, we designed experimental and mathematical strategies for performing assessments with the nonpurified αβ3-Fc fusion protein (i.e., directly from crude supernatant of transfected cells). Through this, more accurate kinetic and thermodynamic unbinding parameters at the single-molecule level of the Thy-1/αβ3 dissociation were obtained even when nonspecific rupture events were detected. Additionally, a slip bond behavior for the bimolecular interaction between Thy-1 and αβ3 integrin was identified. The parameters obtained were confirmed using purified αβ3 integrin. Of note, nonspecific interactions were detected even for purified proteins, meaning that our mathematical approach might also be of use when working with purified proteins.

**RESULTS**

Rupture force measurements and the specificity of the Thy-1 and αβ3 integrin interaction

The optical tweezers technique (miniTweezers) (Smith et al., 2003) was used to determine the binding parameters between Thy-1 and αβ3 integrin at the single-molecule level in response to pulling forces. Fc-tagged recombinant molecules were bound to protein G-coated polystyrene beads for the force spectroscopy experiments (see Figure 1A for schematic representation). To promote the Thy-1-Fc/αβ3-Fc interaction, the optically trapped bead coated with αβ3-Fc was approached to the micropipette-attached bead coated with Thy-1-Fc. If a protein-protein interaction occurs, then the optically trapped bead is displaced from the center, and force is generated. When the generated force is greater than the force that disrupts the bond, rupturing of the interaction ensues. The rupture force is obtained from a force-trap position trace (Figure 1A).

To characterize the binding parameters of the interaction between purified Thy-1-Fc and αβ3-Fc integrin in supernatants using molecular force spectroscopy, single-molecule interactions are needed. Therefore, it was first necessary to estimate the concentration of αβ3-Fc in the supernatants, with specific focus on determining the lowest concentration required for single-molecule binding events with Thy-1. Considering that supernatants obtained from HEK293T cells transfected with αβ3-Fc contained other proteins (Supplemental Figure S1A), a semiquantitative Western blot analysis was performed using antibodies against the Fc fragment (Supplemental Figure S1B). Applying the linear regression equation (Supplemental Figure S1C) and a molecular weight of ~300 kDa for the αβ3-Fc molecule, the concentration obtained for the αβ3-Fc integrin in the supernatant was 4 nM. Subsequently, different dilutions of the supernatant containing the αβ3-Fc fusion protein (without dilution, 10-2, 10-3, and 10-fold dilutions) were incubated with protein G-beads to define the lowest concentration needed for single-molecule binding events between Thy-1-Fc and αβ3-Fc. Since many different proteins were present in the HEK293T cell–based supernatant, we recognized some might compete with αβ3-Fc to interact with Thy-1. Such interactions would generate nonspecific binding events, which might partially obscure the signals of the specific interaction between Thy-1-Fc and αβ3-Fc. To try and limit measurements to only specific-binding events, protein G was used to specifically bind αβ3-Fc to the beads. To test whether nonspecific binding events persisted despite this measure, and to estimate
The magnitude of such events, the supernatant of HEK293T cells transfected with an empty plasmid vector was used as a control (mock plasmid). The absence of αvβ3-Fc integrin in this control supernatant was confirmed by Western blotting (Supplemental Figure S1, A and B). The mock supernatant was diluted just as the αvβ3-Fc-containing supernatants.

**FIGURE 1:** Rupture force measurements for specific and nonspecific Thy-1-Fc interactions with different supernatant dilutions with or without αvβ3-Fc integrin using an optical tweezers system. (A) Scheme of the optical tweezers technique used to study the Thy-1-Fc/αvβ3-Fc interaction. Left, MiniTweezers experimental design. Two different sizes of protein G-coated beads were used; the smaller bead (2.1 μm) contained purified Thy-1-Fc and was attached to a micropipette by suction, whereas the larger bead (3.1 μm) contained the αvβ3-Fc protein and was trapped by a laser beam and held in the focus of the microscope. Right, force-trap position trace for one approach-retraction cycle. To promote protein interaction, the laser beam-trapped bead contacted the other bead (approach). After <1 s of contact, the beads were separated by pulling the laser beam-trapped bead at a constant force-loading rate (10 pN/s) (retraction) until interaction rupturing. Dashed lines in the red laser beam represent the center of the optical trap. (B–I) Rupture force histograms obtained for interactions between (B, D, F, and H) Thy-1-Fc/αvβ3-Fc integrin supernatant or (C, E, G, and I) Thy-1-Fc/mock plasmid supernatant, at different dilutions. (B, C) Undiluted supernatants or supernatants diluted (D, E) 10⁶-fold, (F, G) 10⁹-fold, or (H, I) 10¹²-fold. All histograms summarize at least 300 binding events per five pairs of new beads. Solid line corresponds to a Gaussian distribution curve used to illustrate the distribution of rupture forces and characterize the peak rupture force. Representative force-trap position traces obtained from MiniTweezers measurements between Thy-1-Fc and the supernatant (J) containing the αvβ3-Fc integrin or (K) from cells transfected with mock plasmid, both diluted 10⁶-fold.
When using the undiluted supernatant containing αβ3-Fc, the resulting rupture forces showed bimodal distribution, the peaks of which (19.9 ± 3.2 and 50.0 ± 6.4 pN) were indicative of multiple binding events (Figure 1B). However, only the lower peak rupture force, characteristic of single-molecule interactions, was detected following dilution of the αβ3-Fc-containing supernatant (Figure 1, D, F, and H). On the other hand, similar rupture force distributions were found in all spectroscopy experiments using different mock-supernatant dilutions (Figure 1, C, E, G, and I), suggesting that Thy-1-Fc promotes nonspecific interactions characterized by low rupture forces (~10 pN). Differences between the rupture force histograms obtained for Thy-1-Fc/αβ3-Fc and Thy-1-Fc/mock plasmid were only detected up to a 10^3-fold dilution (Figure 1, F and G). The representative force-trap position traces obtained for the 10^6 dilutions of both Thy-1-Fc/αβ3-Fc and Thy-1-Fc/mock plasmid are shown in Figure 1, J and K, respectively. Indeed, when the αβ3-Fc was diluted 10^3-fold, the histogram profile and peak rupture force were similar to that obtained for the Thy-1-Fc/mock plasmid (Figure 1, H and I). These results indicate that a 10^4-fold dilution of the αβ3-Fc-containing supernatant would be the most diluted condition at which rupture forces differ from the control (i.e., Thy-1-Fc/mock plasmid). Consequently, this concentration would also represent the highest dilution at which specific Thy-1-Fc/αβ3-Fc interactions could be detected. Therefore, the 10^4-fold dilution was defined as the most diluted condition needed for specific single-molecule binding events between Thy-1-Fc and αβ3-Fc integrin. These results also suggest the presence of nonspecific binding events, despite that the designed experimental strategy to evaluate purified Thy-1-Fc and αβ3-Fc integrin interactions in supernatants used protein-G to specifically bind the αβ3-Fc integrin to the beads.

Adhesion frequency measurements and the specificity of the Thy-1 and αβ3 integrin interaction

Adhesion frequency was determined to improve the characterization of binding specificity between Thy-1-Fc and the αβ3-Fc integrin obtained from the supernatant of transfected cells. Adhesion frequency, the probability for adhesion based on the total number of contacts, constitutes a measure of the relative on-rate value (Robert et al., 2007). Evaluating this parameter helps discriminate between specific and nonspecific interaction events. The adhesion frequency was evaluated using either undiluted or diluted supernatants (10^5-, 10^4-, and 10^3-fold) for either the αβ3-Fc-containing (Thy-1-Fc/αβ3-Fc) or control supernatant (Thy-1-Fc/mock plasmid) (Figure 2A). The highest adhesion frequency was observed for the interaction between Thy-1-Fc and the undiluted supernatant containing αβ3-Fc (Figure 2A). Values for adhesion frequencies of supernatants at the 10^5- and 10^4-fold dilutions tended to decrease with respect to those obtained for the nondiluted supernatant; however, the differences were not statistically significant. A significant decrease in the adhesion frequency was observed for the interaction between Thy-1-Fc and the αβ3-Fc integrin at the 10^3 dilution. Considering the similarity of this value with those of the adhesion frequency determined for Thy-1-Fc and the mock supernatants (Figure 2A), these results indicate that the binding events occurring between Thy-1-Fc and the αβ3-Fc integrin-containing supernatant at the 10^3 dilution are due exclusively to nonspecific interactions (compare green bar at 10^3 dilution with orange bars, Figure 2A).

To corroborate the specific binding events between Thy-1-Fc/αβ3-Fc integrin at the single-molecule level (i.e., αβ3-Fc at a 10^3-fold dilution), the interaction was characterized using a supernatant predepleted of the αβ3-Fc integrin (see Materials and Methods and Supplemental Figure S2A). Predepletion of the αβ3-Fc integrin...
decreased the adhesion frequency of Thy-1-Fc/depleted αββ3-Fc by ∼30% compared with the Thy-1-Fc/αββ3-Fc condition (Figure 2B). As expected, the adhesion frequency between Thy-1-Fc and the αββ3-Fc-depleted supernatant was equivalent to that obtained for the Thy-1-Fc/mock plasmid (Figure 2B). These results suggest that depletion of the fusion protein from the supernatant eliminates the specific binding events between Thy-1-Fc and the αββ3-Fc integrin and that nonspecific interactions occurring in Thy-1-Fc/depleted αββ3-Fc are comparable to those in Thy-1-Fc/mock supernatant.

As a second strategy to corroborate specific binding between Thy-1-Fc and αββ3-Fc, we used the Thy-1(RLD)-Fc protein, which is mutated in the integrin binding site (RLD) (Hermosilla et al., 2008). The interaction of this protein was tested with the αββ3-Fc supernatant. As anticipated, a lower adhesion frequency than that calculated for wild-type Thy-1-Fc was detected, similar to what was observed when nonspecific binding was evaluated (Figure 2B). These results confirm the specific interaction of Thy-1 with αββ3-Fc and indicate that nonspecific binding events are independent of the Thy-1 integrin-binding site.

Additionally, to study whether nonspecific interactions depended on Thy-1-Fc and the protein G-beads, the adhesion frequency was evaluated between Thy-1-Fc and protein G-beads preincubated only with the HEPES buffer (Thy-1-Fc/buffer). A different control included measuring the interaction using beads incubated only with buffer (buffer/buffer). The adhesion frequency between Thy-1-Fc and the uncoated protein G-beads was ∼53% lower than the condition with supernatant from cells transfected with mock plasmid (i.e., Thy-1-Fc/buffer vs. Thy-1-Fc/mock plasmid, Figure 2B). Notably, a significant decrease in adhesion frequency also occurred with uncoated protein G-beads (i.e., Thy-1-Fc/buffer vs. buffer/buffer, Figure 2B), suggesting that Thy-1-Fc interacts not only with soluble proteins present in the supernatant from transfected cells, but also with the protein G-beads alone. Importantly, these results suggest that the total binding events found in the single-molecule approach, performed with Thy-1-Fc and nonpurified αββ3-Fc integrin, only ∼30% are attributable to specific interactions between Thy-1-Fc and the integrin.

Correction for rupture forces from nonspecific binding events and estimation of Thy-1-Fc/αββ3-Fc binding parameters
The binding parameters of the Thy-1/αββ3-Fc integrin interaction can be obtained from the rupture force distribution graph (Figure 1F) by applying the DHS model (Dudko et al., 2008). The experimental setup provided rupture forces representative of both specific and nonspecific binding. Therefore, to characterize the correct binding parameters between Thy-1-Fc and the αββ3-Fc integrin, the rupture forces generated by nonspecific binding were filtered from the total rupture forces obtained in the supernatant for the interaction between pure Thy-1-Fc and the αββ3-Fc.

To estimate the effects of nonspecific binding events, the rupture forces were measured for supernatant interactions of Thy-1-Fc/mock-transfected cells. The presence of only nonspecific binding events in this condition was corroborated by characterizing rupture forces between Thy-1-Fc and supernatants depleted of αββ3-Fc (Supplemental Figure S3A). Rupture forces were also characterized between the mutant Thy-1(1RL)-Fc and supernatant containing αββ3-Fc (Supplemental Figure S3B). As expected, both conditions showed rupture force distributions and peak rupture forces equivalent to those obtained for Thy-1-Fc and the supernatant from cells transfected with mock plasmid (Figure 1G). These results are consistent with those depicted in Figure 2B, where similar adhesion frequency values were obtained for each condition in which nonspecific interactions were observed.

To correct nonspecific rupture forces from the total rupture forces obtained for Thy-1-Fc/αββ3-Fc in the supernatant interaction, a mathematical strategy was developed (see Materials and Methods and the Supplemental Material). Briefly, the assumption was that the total rupture forces measured between Thy-1 and the mock supernatant must have resulted from only nonspecific binding events, which occur with the probability \( P(AB) \) (Figure 3A). On the other hand, the total rupture forces measured in the supernatant containing αββ3-Fc must have resulted from both nonspecific and specific binding events. The probability of nonspecific events remained \( P(AB) \), while that of specific binding events was defined by \( P(AB) \) (Figure 3B). Since both events are exclusive and complementary, \( P(AB) + P(AB) = 1 \), thus allowing calculation for the value of \( P(AB) \).

To determine the contribution of specific Thy-1-Fc/αββ3-Fc interactions to the rupture forces measured in the Thy-1-Fc/αββ3-Fc-containing supernatant, the measured rupture forces were weighted by the probability \( P(AB) \). After applying this mathematical correction, a rupture force histogram representative of only specific interactions between Thy-1-Fc and the αββ3-Fc integrin was obtained (Figure 3C). Fitting with a Gaussian distribution curve resulted in a peak rupture force of 25.4 ± 1.6 pN.

Thy-1-Fc/αββ3-Fc interaction lifetimes were calculated for each bin of the rupture force histogram by applying the DHS model (Eq. 1; see Materials and Methods; Figure 3D). The lifetime data were then fitted by applying the nonlinear DHS model (Eq. 2; see Materials and Methods), thus characterizing the energy landscape of the unbinding process between Thy-1-Fc and the αββ3-Fc integrin (Figure 3E). Bond lifetime values (\( t^* = 19.30 ± 1.34 s \)), the off-rate constant (\( k_{off} = 0.051 ± 0.003 s^{-1} \)), the distance to the transition state (\( \Delta x^* = 0.50 ± 0.05 nm \)), and the free energy of activation (\( \Delta G^\ddagger = 5.54 ± 0.47 \text{ kJ} \)) were calculated in the absence of force. Therefore, the applied experimental and mathematical strategies allowed us to characterize the force-dependent kinetic parameters of the dissociation process between Thy-1 and αββ3 integrin, despite using a nonpurified integrin.

Experimental validation of the mathematical strategy used to obtain binding parameters for interactions between purified and nonpurified proteins
To validate the implemented mathematical approach, the αββ3-Fc integrin was purified to study the interaction parameters between Thy-1-Fc and the pure αββ3-Fc fusion protein under force. Previous attempts to purify the αββ3-Fc fusion protein have generated a nonfunctional and unstable integrin (Hermosilla et al., 2008). Therefore, mutations were introduced into the C\( \beta \)3 domains of the Fc fragment, as described to purify αββ1-Fc (Coe et al., 2001). These mutations reportedly improve the possibility of maintaining functional heterodimers after purification. SDS-PAGE with silver staining revealed the presence of αβ-Fc and β3-Fc subunits, as well as other proteins with greater electrophoretic mobility in the supernatant obtained from transfected cells. However, only αβ-Fc and β3-Fc subunits were detected after purification (Supplemental Figure S4A). Additionally, the purified product was also detected by Western blotting using antibodies against the Fc fragment. Western blotting analyses of the eluates were performed using antibodies against the Fc fragment under reducing and nonreducing conditions. The αβ-Fc and β3-Fc subunits were found under reducing conditions, while the αββ3-Fc heterodimer was found in nonreducing conditions (Supplemental Figure S4, B and C).
To test whether functionality of the αvβ3-Fc protein was maintained after purification, an enzyme-linked immunosorbent assay was used. In this assay, the nonpurified and pure αvβ3-Fc fusion proteins bound to known RGD-containing ligands, including fibronectin, vitronectin, and the Kis peptide (Leyton et al., 2001). Nevertheless, binding did not occur with RGE-containing peptides.
indicating interaction specificity with integrin-binding molecules (Figure S5). As for the nonpurified protein, the concentration of the pure αvβ3-Fc was estimated by a semiquantitative Western blotting assay (Supplemental Figure S4D) and validated using the bicinchoninic acid assay. The determined concentration was ~130 nM. To ensure correct comparisons with the parameters estimated for the nonpurified fusion protein, pure αvβ3-Fc was diluted 10^6-fold. The peak rupture force obtained between Thy-1-Fc and the diluted pure αvβ3-Fc was 24.9 ± 2.1 pN (Figure 4A).

To test the binding specificity between Thy-1-Fc and the pure αvβ3-Fc integrin, thereby confirming whether the use of a purified protein eliminated nonspecific binding events, a range of adhesion frequency measurements were carried out. A higher adhesion frequency was observed for the pure αvβ3-Fc protein with Thy-1-Fc than with the mutant Thy-1(RLE)-Fc (Figure 4B), indicating that this purified protein can still directly interact with the RLD motif in Thy-1-Fc. Considering that Thy-1-Fc nonspecifically interacted with the buffer-treated beads, the adhesion frequency was evaluated between Thy-1-Fc and beads preincubated with the HEPES buffer. As expected, a similar adhesion frequency was observed as compared with the control Thy-1(RLE)-Fc/pure αvβ3-Fc condition (Figure 4B). Moreover, similar rupture force distribution profiles were observed for both the Thy-1(RLE)-Fc/pure αvβ3-Fc and Thy-1-Fc/buffer conditions (Supplemental Figure S3, C and D). This similarity suggests the presence of comparable nonspecific binding events even when both proteins are pure. Considering that nonspecific events represent 20% of all events, and to correctly determine the binding parameters between Thy-1-Fc/pure αvβ3-Fc, the developed mathematical method was applied to correct rupture forces from nonspecific interactions, resulting in a peak force of 24.5 ± 1.8 pN (Figure 4C).

The force-dependent lifetime data τ(F) obtained using the DHS model are shown in Figure 4D. The lifetime data were fitted by applying the nonlinear DHS model (Eq. 2; see Materials and Methods), thus characterizing the energy landscape of the dissociation process between Thy-1-Fc and the pure αvβ3-Fc integrin. The values obtained at zero force were as follows: binding lifetime (t^0 = 27.25 ± 7.70 s), off-rate constant (koff^0 = 0.036 ± 0.010 s^-1), distance to the transition state (Δx^‡ = 0.54 ± 0.16 nm), and free energy of activation (ΔG^‡ = 6.16 ± 0.55 k_B T). Importantly, these results are very similar to
those obtained with nonpurified αβ3-Fc in the crude supernatant after applying the mathematical model designed to eliminate nonspecific interactions (see Table 1).

Taken together, our findings indicate that the developed mathematical approach allows for precisely characterizing the effects of mechanical forces on the Thy-1-Fc/αβ3-Fc interaction, even when one of the proteins is unavailable in the purified state.

**DISCUSSION**

In this study, we characterized for the first time how force regulates interaction dissociations between the glycoprotein Thy-1 and the αβ3 integrin, proteins that mediate cell–cell adhesion (Saalbach et al., 2005; Herrera-Molina et al., 2013), including bidirectional neuron-to-astrocyte communication (Hermosilla et al., 2008; Herrera-Molina et al., 2012; Alvarez et al., 2016; Maldonado et al., 2016). Direct force measurement results of the interaction between the αβ3 integrin and Thy-1 were obtained using optical tweezers. This force-transducer technique (Ashkin, 1997; Smith et al., 2003) has been employed to research different bimolecular interactions, such as those between αββ2 integrin/fibrinogen (Litvinov et al., 2005, 2011), P- and L-selectins/PSGL-1 (Rinko et al., 2004), and αβ3 integrin/osteonectin (Litvinov et al., 2003).

Molecular force spectroscopy is the experimental approach of choice for studying how mechanical forces regulate bimolecular interactions (Fiore, Ju, et al., 2014; Li et al., 2003), which has been also used to study intact cells and purified proteins (Zhang et al., 2004; Litvinov et al., 2005; Sun et al., 2005). This technique is highly sensitive, and purified proteins are desirable to reduce the possibility of nonspecific interactions. However, the purification of a protein could represent a limiting factor. In our own previous attempt to purify the αβ3-Fc fusion protein, the final purified product was a nonfunctional integrin (Hermosilla et al., 2008). To overcome this limitation, we designed an experimental and mathematical strategy that allowed us to perform the present study with a nonpurified αβ3-Fc fusion protein. For this, supernatants containing the αβ3-Fc protein were incubated with protein G-coated beads to separate the integrin from other molecules, as is commonly achieved in pull-down assays. Surprisingly, even when protein G specifically interacts with the Fc fragment (Kato et al., 1995), rupture forces due to nonspecific interactions were detected between Thy-1 and the other soluble molecules. Moreover, Thy-1-Fc interacted nonspecifically with the beads themselves. Interestingly, the strength determined for these nonspecific interactions reached values of ~10 pN (i.e., peak of rupture forces), which are in line with previous reports for nonspecific protein–protein interactions (Weisel et al., 2003; Litvinov et al., 2005). This rupture force peak differed from that obtained with the supernatants containing the αβ3-Fc protein (~25 pN), however, in the lower force range, the nonspecific interactions partially overlapped the specific Thy-1/αβ3 interaction. To overcome this issue, a mathematical algorithm was developed and used to correct the nonspecific rupture forces and, consequently, obtain an accurate estimation of the dissociation rupture forces of specific Thy-1-Fc/αβ3-Fc interactions.

To confirm the results obtained with the nonpurified αβ3-Fc integrin, and to validate the employed mathematical approximation, pure and stable αβ3-Fc protein was obtained as reported for αβ51-Fc (Coe et al., 2001). The purified αβ3-Fc bound to ligands containing an integrin-binding site (e.g., fibronectin and vitronectin) as expected (Charo et al., 1990), corroborating the notion that mutations in the Fc fragments did not affect the ability of the αβ3-Fc integrin to interact with corresponding ligands, such as Thy-1. Importantly, the unbinding parameters of the Thy-1-Fc/pure αβ3-Fc interaction were highly comparable to those obtained with nonpurified αβ3-Fc in the crude supernatants after filtering nonspecific interactions. This observation demonstrates that our approach permits characterizing how protein–protein interactions respond to mechanical forces, even when one of the molecules involved is not available in the purified state.

In assessing the nonpurified αβ3-Fc protein, rupture forces resulting from single-molecule interactions were corroborated using several criteria. For instance, the obtained histograms showed forces resulting only from single-step bond ruptures (force-trap position graphs, Figure 1, J and K). While multiple binding events are represented by a series of peaks in rupture force histograms, single binding events are characterized by a unique peak (Weisel et al., 2003; Litvinov et al., 2005). The two-peak rupture forces detected when nondiluted supernatants containing αβ3-Fc protein were evaluated (Figure 1B) suggest the presence of multiple binding events. However, a single peak at ~25 pN appeared when the integrin was diluted (Figure 1, D, F, and H), which would be representative of single-binding events. Furthermore, the peak of rupture forces for single-binding events should not be modified by changes in the surface density of the interacting proteins (Zhu et al., 2002; Litvinov et al., 2005). Accordingly, a peak value of ~25 pN was detected for each dilution of the supernatant containing the αβ3-Fc protein. In contrast, variations in surface density can change the adhesion frequency between proteins (Litvinov et al., 2005). In the current assessments, a tendency towards decreased frequency values was detected on reducing the αβ3-Fc concentration. However, a significant decrease in the adhesion frequency value was observed only when the integrin concentration was diluted to the attomolar level (i.e., 10⁷ dilution).

Bond strength is the most probable bond rupture force and can be estimated from rupture–force distribution histograms (Evans and Ritchie, 1997). In this study, a bond strength of ~25 pN was determined for Thy-1/αβ3 binding. The peak rupture force for adhesion proteins are characteristic of each ligand–receptor pair and range from 20 to 150 pN (Weisel et al., 2003). However, it is important to note that for noncovalent interactions, bond strength also depends on the rate at which force is applied (force loading rate) (Evans and Ritchie, 1997; Rico et al., 2007). Common pulling rates used in single-molecule experiments are in the range of 10⁵–10⁶ nm/s (Roca-Cusachs, Iskratsch, and Sheetz, 2012). Here, force-ramp assays were performed under similar conditions (i.e., constant rate of 100 nm/s), and considering the stiffness of the system (spring constant of 0.1 pN/nm), the loading rate value would be 10 pN/s (pulling rate × spring constant).

|                | Lifespan (τ) | Off-rate (kₐff) | Distance to transition state (Δx) | Free energy of activation (ΔG) |
|----------------|-------------|----------------|----------------------------------|-------------------------------|
| Nonpurified αβ3-Fc | 19.30 ± 1.34 s | 0.051 ± 0.003 s⁻¹ | 0.50 ± 0.05 nm | 5.54 ± 0.47 kBT |
| Pure αβ3-Fc    | 27.25 ± 7.70 s | 0.036 ± 0.010 s⁻¹ | 0.54 ± 0.16 nm | 6.16 ± 0.55 kBT |

**TABLE 1: Comparison of kinetic and thermodynamic parameters in the absence of force.**
(Dudko et al., 2008). By comparison, the bond strength for the interaction between fibronectin and αIIbβ3 is ~50 pN using a pulling rate of 160 nm/s (Agnihotri et al., 2009), which suggests a stronger interaction than for Thy-1/αvβ3 integrin binding. In other studies, a lower bond strength (9.2 ± 4.4 pN), as compared with the present Thy-1/αvβ3 interaction, has been reported for actin/myosin binding using a loading rate of 12 pN/s (Nishizaka et al., 2000). Interestingly, results obtained by atomic force microscopy demonstrated that the interaction between the αvβ3 integrin and GRGDS peptide has a bond strength of 42 ± 4 pN at a force loading rate of 30,000 pN/s (Lehenkari and Horton, 1999). Considering that the Thy-1/αvβ3 interaction also depends on the RGD-like integrin-binding domain (RLD), the result obtained with a fast loading rate (30,000 pN/s) suggests that force loading rate increases greater than three orders of magnitude only modify the bond strength found for the Thy-1/αvβ3 interaction by approximately twofold. Bonds that are almost insensitive to stress rates are known as “persistent bonds” and have been reported for the force responses of the interaction between the intracellular cell adhesion molecule-1 (ICAM-1) and the αLβ2 integrin (Evans and Calderwood, 2007; Evans et al., 2010). In this context, we found that in the range of 5 and 100 pN/s, the bond strength for Thy-1/αvβ3 interaction is almost not affected by loading rates (Supplemental Figure S6A). Importantly, current measurements were taken using a force loading rate within the physiological range (Yamazaki et al., 2002; Li et al., 2003), suggesting that the bond strength observed for Thy-1/αvβ3 integrin interactions is likely representative of that existing in the CNS. Despite that bond strength depends on the rate at which force is applied, it is important to note that according to the DHS model, lifetime data obtained at different loading rates (i.e., at different pulling rate) must collapse onto a single-master curve that describes the force-dependent lifetimes (Dudko et al., 2008), prediction that was observed with loading rates up to 100 pN/s (Supplemental Figure S6B). In fact, the low loading rate (10 pN/s) curve obtained was maintained at a higher loading rate (50 and 100 pN/s). Consequently, the unbinding parameters calculated for Thy-1/αvβ3 integrin interaction were not modified by an increased loading rate (Supplemental Figure S6B).

Applying the DHS model (Dudko et al., 2008) to the rupture force data obtained from crude supernatants and adjusted for nonspecific binding effects resulted in an estimated off-rate of ~0.051 s⁻¹ for Thy-1/αvβ3. Interestingly, an off-rate of 0.012 s⁻¹ has been reported for the interaction between fibronectin and high-affinity form of the α5β1 integrin (Li et al., 2003), which is also involved in cellular adhesion and migration (Danen et al., 2005; Morgan et al., 2013). Although the off-rate values are in the same order of magnitude, the absolute values indicate that the interaction between Thy-1 and αvβ3 integrin is less stable in the absence of force than binding between fibronectin and the α5β1 integrin. Additionally, fibronectin/α5β1 integrin binding values of ΔΓ = 0.42 nm have been reported (Li et al., 2003), which are similar to those obtained for the Thy-1/αvβ3 interaction (ΔΓ = 0.5 nm) and indicative of ruptures for similar bond types. In both cases, the reported values are probably attributable to the RGD-like tripeptide. Likewise, the DHS model has been applied to characterize unbinding parameters for the interaction between αIIbβ3 integrin and a cyclic peptide integrin antagonist, a drug used to prevent interaction of the platelet integrin αIIbβ3 with fibrinogen and blood clot formation (Dutta, Horita, et al., 2013). The αIIbβ3/antagonist interaction had an off-rate of ~0.02 s⁻¹, indicating that the rate at which the dissociation between Thy-1 and αvβ3 integrin occurs is faster than that for αIIbβ3/antagonist binding. This result is to be expected given that the free energy of activation calculated for the αIIbβ3/antagonist interaction (~11.7 k_BT) is greater than the values described herein for Thy-1 and the αvβ3 integrin (~5.54 k_BT).

In the present study, force exponentially accelerated Thy-1/αvβ3 integrin dissociation, suggesting a slip bond behavior. This behavior has been described for the interaction between αIIbβ3 and fibrinogen in a force range up to 50 pN (Litvinov et al., 2011). However, the catch bond phenomenon has been described most widely for integrin molecules. For instance, force prolongs bond lifetime up to 15 pN in the interaction between αLβ2 and ICAM-1 (Chen et al., 2010). Interestingly, the interaction between αvβ3 integrin and fibrinectin also shows a catch regime in the range of 0–20 pN (Eliseou-Artola et al., 2016; Chen et al., 2017), indicating differences between both Thy-1/αvβ3 and fibrinectin/αvβ3 integrin interactions. Such differences have also been found for α5β1 integrin. The fibrinectin/α5β1 integrin interaction shows catch bond behavior in which maximum lifetimes occur at forces of 10–30 pN (Kong et al., 2009). Nevertheless, as well as for αvβ3 integrin, an ordinary slip bond behavior has been described for the interaction between Thy-1 and αvβ3 integrin (Fiore, Ju, et al., 2014). For α5β1 integrin, the catch bond mechanism depends on the synergy-site engagement (Friedland, Lee, and Boettiger, 2009; Sun et al., 2016). Particularly in the absence of mechanical tension, the α5β1 integrin interacts with the tripeptide RGD motif present in the 10th type III module of fibrinectin (FNIII10) (Aota et al., 1994); however, under tensile forces, this integrin additionally engage the synergy-site in FNIII10 to form a catch bond (Friedland, Lee, and Boettiger, 2009; Kong et al., 2009). In contrast to fibrinectin, Thy-1 interacts either with α5β1 or αvβ3 integrin only through its RGD-like integrin-binding domain (RLD) (Hermosilla et al., 2008; Herrera-Molina et al., 2013; Fiore, Ju, et al., 2014), which could account for the slip bond behavior. Thus, even when the catch bond mechanism for αvβ3 integrin has not been characterized yet, the differences found with fibrinectin and Thy-1 could depend on the presence of synergy-site contacts for the integrins. As mentioned, Fiore and colleagues identified a slip bond behavior between Thy-1 and α5β1 integrin. However, a triphasic force dependence (slip–catch–slip) was observed for interactions with the α5β1 bound to a cell specifically expressing this integrin in the membrane instead of the purified Fc-protein. Our previous research describing the interactions between Thy-1 and the proteoglycan syndecan-4 (Avalos, Valdivia, et al., 2009; Kong, Munoz, et al., 2013; Alvarez et al., 2016) support that Thy-1 forms a trimolecular bond with integrin and syndecan-4. This trimolecular bond displays a triphasic phenomenon termed dynamic catch, where increasing force abruptly stiffens the complex and is followed by the formation of a catch bond (Fiore, Ju, et al., 2014). Our prior results further indicate that the neuronal Thy-1 and astrocytic syndecan-4 interaction is involved in neuron-to-astrocyte bidirectional communication mediated by the Thy-1/αvβ3 integrin interaction (Avalos, Valdivia, et al., 2009; Kong, Munoz, et al., 2013). Considering that Thy-1 interacts with syndecan-4 and αvβ3, both astrocytic molecules, and that syndecan-4 levels are up-regulated in astrocytes following CNS injury (Iseki et al., 2002), as is also the case for the αvβ3 integrin (Ellison et al., 1999; Lagos-Cabre, Alvarez, et al., 2017), the results reported by Fiore and colleagues (Fiore, Ju, et al., 2014) suggest that the Thy-1/αvβ3 integrin/syndecan-4 complex may also exhibit dynamic catch bond behavior within our model system; however, whether this is indeed the case remains to be determined.

In summary, to our knowledge, our study is the first to evaluate the mechanical response of bimolecular Thy-1/αvβ3 integrin binding, providing a detailed characterization of the energy landscape for the dissociation pathway of this interaction. Importantly, we
developed a simple mathematical methodology to investigate how tensile forces tune the kinetics of protein–protein interactions at the single-molecule level. Clearly, further studies are required to fully understand the mechanical regulation of the Thy-1 and αvβ3 integrin interaction in a physiological context. Nonetheless, our research provides an initial step towards facilitating better comprehensions of how neuron and astrocyte communication processes sense and respond to mechanical forces and, consequently, how these events affect the complex behavior of these cell–cell interactions.

MATERIALS AND METHODS
Preparation of Fc-tagged fusion proteins
The recombinant proteins Thy-1-Fc wild-type and the Thy-1-(RLE)-Fc mutant for the integrin-binding site were obtained and purified as previously reported (Leyton et al., 2001; Hermosilla et al., 2008). The αvβ3-Fc fusion protein was expressed in HEK293T cells by transient cotransfection with equal amounts (10 μg) of both αv-Fc (PS711) and β3-Fc (PS675) expression plasmids encoding for the extracellular domains of human αv and β3 integrin subunits fused to the Fc portion of the IgG1 (kindly donated by Pascal Schneider, Department of Biochemistry, University of Lausanne, Switzerland) (Schneider, 2000), using the X-treme GENE HP DNA Transfection Reagent (Roche). After 16 h of transfection, cells were grown in serum-free medium (DMEM, high glucose) for 2 d. Next, the culture serum-free supernatant containing secreted αvβ3-Fc heterodimer was centrifuged (800 rpm × 5 min), filtered (0.2 μm; Whatman), and stored at −20°C. Control supernatant was obtained by transfection of HEK293T cells with mock plasmids (pCR3.1) in serum-free medium for 2 d and recovered as indicated for αvβ3-Fc-containing supernatant.

Purification of αvβ3-Fc fusion protein
Human αvβ3-Fc integrin was purified from culture supernatants of transfected HEK293T cells via the Fc domain after mixing supernatant (3 ml) with protein-A-sepharose (500 µl) during 16 h. Resin with bound proteins was collected in a column and washed with Tris-buffered saline (Tris-HCl 50 mM, pH 7.5; NaCl 150 mM). Proteins were eluted with citric acid (50 mM, pH 2.7) and neutralized with Tris-HCl (1 M, pH 8.0). Eluates obtained were mixed and concentrated in ultracentrifuge tubes (Amicon 30 K). To enhance the probability of heterodimerization between αv-Fc and β3-Fc after low-pH treatment, specific mutations were introduced into the Cα3 domains of Fc fragments by oligonucleotide-directed PCR mutagenesis as described (Coe et al., 2001). Specifically, in the Fc DNA of the αv-Fc subunit, the small amino acid threonine (residue 366) was changed to the large amino acid tyrosine (αv-Fc), while in the Fc DNA of the β3-Fc subunit the large amino acid tyrosine (residue 407) was changed to a threonine (to create a “hole”) (Ridgway et al., 1996). Both nonpurified and pure αvβ3-Fc proteins used in this study contain the mutations in the Fc fragments.

Characterization of the αvβ3-Fc fusion protein
Nonpurified αvβ3-Fc fusion protein was detected in the supernatant of transfected HEK293T cells by Coomassie blue–stained SDS–PAGE. Eluted αvβ3-Fc integrin was characterized by Silver staining and on immunoblots under reducing and nonreducing conditions using specific anti-human IgG (Fc-specific)-horseradish peroxidase (HRP) antibody (1:2000; Sigma). Semiquantitative Western blotting was performed to estimate the amount of purified αvβ3-Fc fusion protein, as well as the nonpurified αvβ3-Fc proteins in supernatants. The standard curve based on signal intensity was generated using different amounts of the fusion protein of the receptor for the soluble apoptosis-inducing ligand TRAIL (TRAIL-R2-Fc) (Schneider, 2000). The functionality of αvβ3-Fc was tested using an enzyme-linked immunosorbent assay as previously described (Hermosilla et al., 2008). This assay allowed evaluating the ability of both nonpurified and pure αvβ3-Fc to interact with RGD-containing ligands, including fibronectin, vitronectin, and snake venom peptides (Kis RGD-peptide) (Hermosilla et al., 2008; Leyton et al., 2001). As a negative control, RGE peptides were used. Integrin-ligands were immobilized in a 96-well plate (Maxisorp, Nunc). After blocking nonspecific binding sites with 1% bovine serum albumin (BSA) in phosphate-buffered saline (PBS), supernatants containing the αvβ3-Fc protein (25 µl) as well as the purified and concentrated αvβ3-Fc fusion protein (1 µl) were added to the wells, incubated, and washed, and bound Fc-tagged proteins were detected with HRP-coupled goat anti-human IgG. The chromogenic substrate 3,3’-,5,5’-tetramethylbenzidine (TMB; Thermo Scientific) was used to reveal the HRP activity. TMB solution was added to each well (100 µl) and incubated at room temperature for 30 min. Reaction was stopped with sulfuric acid (2 M, 100 µl), and absorbance was read at 450 nm in an Infinite 200 pro NanoQuant Microplate Readers (Tecan).

Depletion of αvβ3-Fc fusion protein from supernatant
Supernatant containing the αvβ3-Fc fusion protein (500 µl) was incubated with an excess of protein A-Sepharose beads (50 µl) during 1 h at 4°C. The solution was then centrifuged at 2000 rpm × 5 min. This process was repeated four times, to ensure complete depletion of the fusion protein. Each supernatant and precipitate obtained was evaluated by Western blotting using anti-human IgG (Fc specific)-HRP antibody (1:2000; Sigma). After the fourth incubation/centrifugation cycle, the level of the αvβ3-Fc protein was undetectable in the precipitate (precipitate 4); thus the fourth supernatant was used in the miniTweezers experiments.

Optical tweezers setup and force measurement protocols
Molecular force spectroscopy experiments were performed with the miniTweezers instrument, which has a dual-laser beam single trap design (Smith et al., 2003). The rupture forces required to dissociate protein–protein interactions are measured optically by the change in momentum of the light beams leaving the trap. This instrument is capable of force, spatial, and time resolution of 0.1 pN, 0.1–2 nm, and 10^{-4} s at 100-Hz bandwidth (Moffitt et al., 2008; Neuman and Nagy, 2008). Details of the miniTweezers instrument and its operation are available on http://tweezerslab.unipr.it. All experiments were carried out in a temperature-controlled room at 25°C employing HEPES buffer (10 mM HEPES, pH 7.4, 150 mM NaCl) containing 1 mM MgCl₂ and using a microchannel with a glass micropipette (inner diameter <1 μm). Considering the properties of our recombinant molecules (Fc-tagged proteins: Thy-1-Fc and αvβ3-Fc), we used protein G-coated polystyrene particles (Spherotech) to attach the molecules via the Fc fragment. To study the mechanical rupture of the Thy-1-Fc/αvβ3-Fc interaction, the bead (3.1 μm) coated with αvβ3-Fc was held in the optical trap, while the bead (2.1 μm) coated with Thy-1-Fc was held on the tip of the micropipette by suction, and cycles of approaching and retraction of the beads were performed. Binding between proteins was observed after bringing the beads into close proximity by moving the optically trapped bead toward the other bead attached to the micropipette (approaching). To dissociate Thy-1-Fc/αvβ3-Fc interaction, a force-ramp assay was performed, where the optically trapped bead is moved in the opposite direction at a constant pulling rate of 100 nm/s (retraction). Considering the stiffness of the system (0.1 pN/nm), the value of the loading rate at which the dissociation was induced corresponds to
10 pN/s (pulling rate × spring constant) (Dudko et al., 2008). If the protein-protein interaction occurs, then the trapped bead is displaced from the center of the optical trap and force is generated. When the force generated is greater than the force required to dissociate the binding, rupture of the interaction is observed; the force is measured directly via the deflection of the trapping laser beam by the trapped bead. The rupture force histograms were obtained with at least 300 binding events per five pairs of new beads and were normalized for the total number of approaching cycles. The bin size (b) was calculated using Scott’s rule: \( b = 3.5 \sigma / (n)^{1/3} \) in which \( \sigma \) is the SD and \( n \) the number of rupture events (Scott, 1979). Adhesion frequency was calculated by measuring the total number of binding events observed in at least 50 approach–retraction cycles per five pairs of new beads.

**Optical tweezers data analysis**

Rupture force histograms were analyzed using the DHS model (Dudko et al., 2008), which permits transforming them into force-dependent lifetimes \( \tau(F) \) as follows:

\[
\tau(F) = \tau_0 \left( 1 - \frac{\Delta G^f}{\Delta G} \right)^{1-\nu_v} e^{-\beta \Delta G \left[ 1 - (1 - \nu_v \Delta G^f / \Delta G)^{1-\nu} \right]}
\]

Here, a rupture force histogram contains \( N \) bins of width \( \Delta F \) that starts at \( F_0 \) and ends at \( F_N = F_0 + N \Delta F \). The number of counts in the \( i \)-th bin is \( C_i \), resulting in height \( h_i = C_i/(N_{tot} \Delta F) \), where \( N_{tot} \) is the total number of counts; \( k = 1, 2, \ldots \).

After applying Eq. 1, data points of the \( (\tau(f))-versus-F \) plot were obtained. The DHS model interpreted the force-dependence lifetime assuming that dissociation of protein–protein bonding can be described as an escape from a deep one-dimensional free energy well (Dudko et al., 2008). Therefore, with the following nonlinear DHS model, the energy landscape and kinetic parameters of the dissociation process at a constant loading rate were characterized as follows:

\[
\tau(F) = \tau_0 \left( 1 - \frac{\nu_v F x^t}{\Delta G} \right)^{1-\nu_v} e^{-\beta \Delta G \left[ 1 - (1 - \nu_v F x^t / \Delta G)^{1-\nu} \right]}
\]

In this expression, \( F \) is the externally applied force and \( \beta = (k_B T)^{-1} \) is the inverse thermal energy, \( \tau_0 \) is the lifetime of the interaction at zero force [inversely related to off-rate constant, \( (g_0)^{-1} \)], \( \Delta x^t \) is the distance to the transition state, and \( \Delta G^f \) is the apparent free energy of activation, describing the energy barrier that must be overcome to dissociate the interaction between Thy-1-Fc and \( \alpha \beta 3-Fc \) in the absence of force. The scaling factor \( v \) describes the shape of the underlying free energy landscape. The variable \( v = 1/2 \) corresponds to a harmonic well with a cusplike barrier, while \( v = 2/3 \) corresponds to a potential that contains linear and cubic terms. For \( v = 1 \), Bell’s formula is recovered (Bell, 1978). Both \( v \) values 1/2 and 2/3 fitted well to our data.

**Mathematical model to correct rupture force due to nonspecific binding events.**

Considering only the cases in which interactions actually occur (these cases are those giving relevant information), two possible outcomes exist: Thy-1-Fc interacts in a specific manner with the \( \alpha \beta 3-Fc \) integrin (event \( AB \)) or Thy-1-Fc interacts in a nonspecific manner with either other proteins present in the integrin-containing supernatant or the surface of the bead (event \( \overline{AB} \)). These events are assumed to be mutually exclusive due to the single-molecule conditions of the experiments, which prohibit both events from occurring simultaneously. The presence of single binding events is supported by the fast, single cooperative rips at which rupture events occurred (see Figure 1J), as opposed to multiple-ripping patterns observed when the rupture of multiple bonds occurs. Applying the law of total probability, the probability of obtaining a rupture force between \( F \) and \( F + dF \), \( P(F) \) will be given by the following:

\[
P(F) = P(F|AB)P(AB) + P(F|\overline{AB})P(\overline{AB})
\]

Here \( P(F|AB) \) and \( P(F|\overline{AB}) \) are the normalized rupture force histograms for the conditions studied between Thy-1-Fc/\( \alpha \beta 3-Fc \)-containing supernatant and Thy-1-Fc/supernatant from mock-transfected cells, respectively. \( P(AB) \) defines the probability of detecting a specific interaction (Thy-1-Fc/\( \alpha \beta 3-Fc \); Figure 3B), while \( P(\overline{AB}) = (1 - P(AB)) \) indicates the probability of being a nonspecific interaction (Thy-1-Fc/other proteins or Thy-1-Fc/bead; Figure 3, A and B).

Using Bayes’s theorem (Bois, 2013), it is possible to obtain useful expression for \( P(AB|F) \) and \( P(\overline{AB}|F) \) as follows:

\[
P(AB|F) = \frac{P(F|AB)P(AB)}{P(F)}
\]

\[
P(\overline{AB}|F) = \frac{P(F|\overline{AB})P(\overline{AB})}{P(F)}
\]

In these expressions, \( P(AB|F) \) and \( P(\overline{AB}|F) \) are the probabilities to find interactions between Thy-1-Fc and the \( \alpha \beta 3-Fc \) integrin or other proteins given the rupture force was between \( F + DF \), respectively. These three relationships (Eqs. 3, 4, and 5) allowed us to obtain the specific interaction events between Thy-1 and the \( \alpha \beta 3 \) integrin by measuring the nonspecific interactions and the mixed (specific + nonspecific) interactions.

In practical terms, if we define \( h(F) = P(F) \), the rupture force histogram for the mixed interactions (i.e., condition studied between Thy-1-Fc/supernatant-containing \( \alpha \beta 3-Fc \) where both specific and nonspecific interactions are detected), \( f_{AB}(F) = P(F|AB) \), the rupture force histogram for the specific Thy-1/\( \alpha \beta 3-Fc \) interactions; \( f_{\overline{AB}}(F) = P(F|\overline{AB}) \), the rupture force histogram for the conditions between Thy-1-Fc/supernatant from mock-transfected cells (without \( \alpha \beta 3-Fc \)); and \( \alpha = P(AB) \), the mixing parameter that defines the probability to detect a specific interaction (Thy-1-Fc/\( \alpha \beta 3-Fc \)), the equations are as follows:

\[
h(F) = \alpha f_{AB}(F) + (1 - \alpha) f_{\overline{AB}}(F)
\]

\[
P(AB|F) = \frac{\alpha f_{AB}(F)}{h(F)}
\]

\[
P(\overline{AB}|F) = \frac{(1 - \alpha) f_{\overline{AB}}(F)}{h(F)}
\]

These expressions represent a direct method of obtaining the specific interactions between Thy-1-Fc and the \( \alpha \beta 3-Fc \) integrin in the nonpurified state, using rupture force data from the Thy-1-Fc/\( \alpha \beta 3-Fc \) supernatant (specific + nonspecific binding events) and the Thy-1-Fc/mock plasmid supernatant (nonspecific binding events) conditions. The equations are derived from basic rules of statistics and correspond to a valid representation, as long as the nonspecific interactions are independent of \( \alpha \beta 3-Fc \) integrin concentration (i.e., the probability of nonspecific interactions do not change in the presence of the \( \alpha \beta 3-Fc \) fusion protein). The validity of the equations considers the following assumptions: 1) the \( \alpha \beta 3-Fc \)
protein does not compete with the nonspecific binding events mediated by Thy-1-Fc, 2) the αvβ3-Fc protein does not interact with other proteins present in the supernatant, and 3) the interactions between Thy1-Fc/αvβ3-Fc are of lower affinity than those nonspecific interactions mediated by Thy-1-Fc (Thy-1-Fc/other proteins or Thy-1-Fc/bead). Under these assumptions, the addition of the αvβ3-Fc integrin could be treated as a small perturbation to the system. Nevertheless, most likely the Thy-1/nonspecific interactions will change with the addition of αvβ3 integrin to the system, since it is clear that the αvβ3 integrin interacts with a high-affinity with its ligands (Xiao and Truskey, 1996). In real conditions, then, the αvβ3-Fc will compete with the nonspecific interactions; however, this competition will mainly affect the nonspecific binding events of lower affinity represented by the low force values in the histograms (see Figure 1, C, E, G, and I). Thus, here, with the assumptions made for our mathematical model, minor changes on the rupture force distribution caused by the competition of specific and nonspecific binding are expected to affect lower rupture force values only.

A step-by-step procedure to obtain the filtered histograms using these expressions is given in the Supplemental Material.

Statistical analysis

Data are expressed as mean ± SEM of three independent experiments. Results were analyzed by non-parametric Mann–Whitney analysis to compare two groups. Statistical significance was set at p < 0.05.

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