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Cryo-EM reveals integrin-mediated TGF- β activation without release from latent TGF- β

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SUMMARY:

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Author Contributions:

M.G.C., and A.C., performed cryo-EM and structural biology. M.G.C. performed cryo-EM data acquisition and processing. A.C. and M.G.C. performed modeling. A.C., S.I. and R.I.S. performed biochemical and cell biologic experiments. S.I., A.C., M.G.C., R.I.S., and S.L.N. designed, generated and characterized mutant integrins and TGF- β constructs. A.C., A.J.B., R.I.S., S.I., J.D.M., J.L., J.L.B. and S.L.N. produced, characterized, cloned and engineered monoclonal antibodies. M.G.C., A.C., S.I., R.I.S., Y.C. and S.L.N. conceived experiments and wrote the manuscript.

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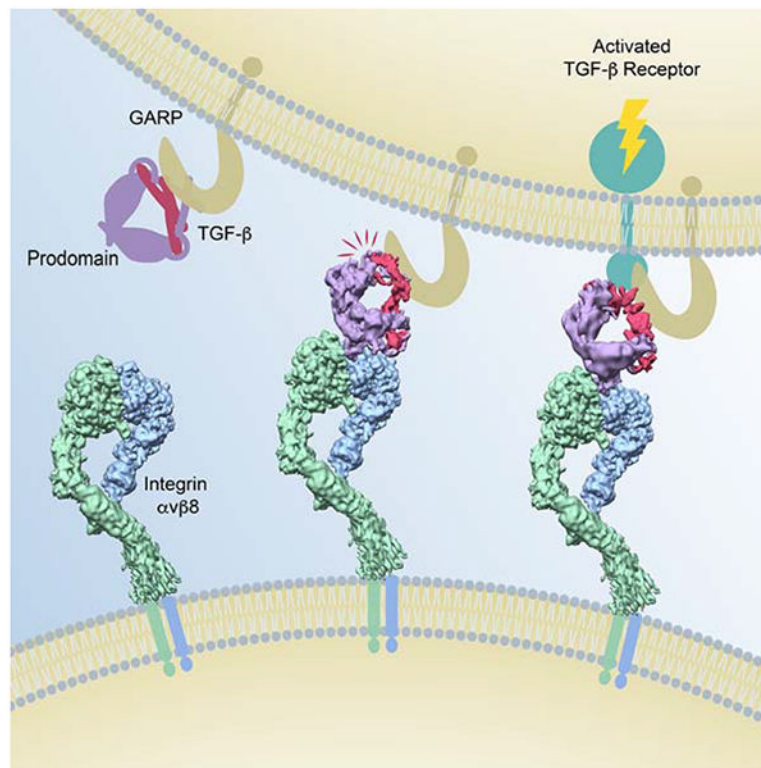
DATA AND CODE AVAILABILITY

All structures and models have been deposited into the EMDB and the PDB. The accession numbers for primary conformation of α v β 8/L-TGF- β (conformation iv) are EMD-20794 and PDB ID 6UJA. For the additional α v β 8/L-TGF- β conformations, i-iii and v-vii the accession numbers are EMD-20797 to EMD-20802. A consensus map of all seven conformations focused at the headpiece region with a nominal resolution of 2.9Å was used for modeling and is deposited with the accession number EMD-21125. For α v β 8/C6D4, the accession numbers are EMD-20795 and PDB ID 6UJB and for α v β 8/C6-RGD3 they are EMD-20796 and PDB ID 6UJC. Three cryo-EM dataset collections containing movies, aligned micrographs, and stacks of α v β 8 bound to L-TGF- β or C6D4 have been deposited into the EMPIAR database: EMPIAR-10343 (α v β 8 bound to L-TGF- β collected on holey carbon grids), EMPIAR-10344 (α v β 8 bound to L-TGF- β collected on graphene oxide grids) and EMPIAR-10345 (α v β 8 bound to C6D4 collected on holey carbon grids).

ADDITIONAL RESOURCES N/A

Integrin $\alpha\text{v}\beta\text{8}$ binds with exquisite specificity to latent transforming growth factor- β (L-TGF- β). This binding is essential for activating L-TGF- β presented by a variety of cell types. Inhibiting $\alpha\text{v}\beta\text{8}$ -mediated TGF- β activation blocks immunosuppressive regulatory T-cell differentiation, which is a potential therapeutic strategy in cancer. Using cryo-electron microscopy, structure-guided mutagenesis, and cell-based assays, we reveal the binding interactions between the entire $\alpha\text{v}\beta\text{8}$ ectodomain and its intact natural ligand, L-TGF- β , as well as two different inhibitory antibody fragments to understand the structural underpinnings of $\alpha\text{v}\beta\text{8}$ binding specificity and TGF- β activation. Our studies reveal a mechanism of TGF- β activation where mature TGF- β signals within the confines of L-TGF- β and the release and diffusion of TGF- β is not required. The structural details of this mechanism provide a rational basis for therapeutic strategies to inhibit $\alpha\text{v}\beta\text{8}$ -mediated L-TGF- β activation.

Graphical Abstract



Abstract

Analysis of intermediate conformations of the interaction between $\alpha\text{v}\beta\text{8}$ integrin and latent TGF- β suggest an activation mechanism that does not require release and diffusion of mature TGF- β , which has implications for current approaches to targeting TGF- β signaling therapeutically.

INTRODUCTION

Studies in the past three decades have elucidated the diverse and essential functions of TGF- β in immunity and vascular biology (Sanjabi et al., 2017; van Meeteren et al., 2011). While

TGF- β and its receptors are ubiquitously expressed (Gordon and Blobel, 2008), TGF- β is always expressed in a latent form (L-TGF- β). During biosynthesis, TGF- β is cleaved by furin from its prodomain, latency associated peptide (LAP), but remains associated with it forming a latent complex, in which LAP forms a ring like structure shielding the active domains of mature TGF- β from its receptors (Fig. S1A)(Shi et al., 2011). Therefore, activation is a major point of regulation of TGF- β function (Miyazono et al., 1990). It is widely presumed that TGF- β activation requires mature TGF- β to be released from LAP to enable receptor binding (Annes et al., 2003; Massagué, 2012; Moses et al., 2016). The majority of therapeutics in development are conceptually tied to inhibiting freely diffusible TGF- β , but have been associated with toxicities (Akhurst, 2017). Targeting TGF- β activation thus has the potential to increase specificity while limiting toxicities associated with such global TGF- β inhibition strategies (Nishimura, 2009).

Integrin $\alpha v \beta 8$ is particularly important for TGF- β activation since it is essential for T-cell, myeloid and endothelial cell differentiation during development (Aluwihare et al., 2009; Arnold et al., 2014; Qin et al., 2018; Travis et al., 2007). In post-natal life, $\alpha v \beta 8$ plays important roles in fibroinflammatory processes (Kudo et al., 2012; Melton et al., 2010; Minagawa et al., 2014), and anti-tumor immunity (Takasaka et al., 2018). In particular, the immunosuppressive function of regulatory T cells (Tregs), which is a major mechanism of tumor immune evasion, is determined by the activation of L-TGF- β presented by GARP (glycoprotein A repetitions predominant) on their cell surface to $\alpha v \beta 8$ (Stockis et al., 2009). How $\alpha v \beta 8$ binds to and activates L-TGF- β has not been elucidated.

A structural model of TGF- β activation by a different integrin, $\alpha v \beta 6$, suggests that L-TGF- β binding to $\alpha v \beta 6$ triggers a global conformational change from extended-closed to extended-open, which allows actin cytoskeletal force to be transmitted through the β -subunit to release mature TGF- β from its latent complex (Fig. S1B)(Dong et al., 2017). However, the $\alpha v \beta 8$ integrin has a divergent cytoplasmic domain that does not interact with the actin cytoskeleton and exists only in an extended-closed conformation whether alone or in complex with L-TGF- β (Cormier et al., 2018; Minagawa et al., 2014; Mu et al., 2002; Wang et al., 2017). Therefore, $\alpha v \beta 8$ -mediated L-TGF- β activation can occur without drastic conformational rearrangements and without cytoskeletal force. Thus, its mechanism of activation must be different from the model proposed for $\alpha v \beta 6$ (Fig. S1C).

$\alpha v \beta 8$ only binds efficiently to L-TGF- β , which is distinct from all other αv -integrins, which bind to many other ligands in addition to L-TGF- β (Humphries et al., 2006; Mu et al., 2002; Ozawa et al., 2016). Therefore, determining the basis of $\alpha v \beta 8$ specificity for L-TGF- β is of general importance for understanding integrin-mediated TGF- β activation, and to guide development of therapeutic antibodies and integrin-specific small molecules, the latter of which has proven difficult (Hatley et al., 2018).

Using cryo-electron microscopy we determined structures of three complexes between the $\alpha v \beta 8$ ectodomain and L-TGF- β , or two inhibitory Fabs, one of which is ligand-mimetic. Together with structure-guided mutagenesis, we reveal a binding intermediate and the two key regions that define the specificity of $\alpha v \beta 8$ for L-TGF- β , suggesting an alternative strategy to inhibit $\alpha v \beta 8$ /L-TGF- β binding. The dynamic nature of $\alpha v \beta 8$ /L-TGF- β

interactions suggest an alternative activation mechanism that does not require release and diffusion of mature TGF- β , which we confirmed through cell-based assays. Our findings present a model and conceptual framework to understand the cell-specific regulation of TGF- β activation.

RESULTS

Structure of the $\alpha v\beta 8$ /L-TGF- β complex

To gain mechanistic insights into TGF- β activation and $\alpha v\beta 8$ ligand specificity, we used single particle cryo-EM to determine structures of a stable complex of the human $\alpha v\beta 8$ ectodomain with porcine L-TGF- $\beta 1$, which has 94% sequence identity with human L-TGF- β . This L-TGF- β has two point substitutions: C4S that increases expression and R249A that disrupts the furin cleavage to stabilize the L-TGF- β complex for structural studies. The biochemical relevance of this complex was established in previous studies (Shi et al., 2011). From the entire cryo-EM dataset, we isolated a subset of particles that give a structure with an overall nominal resolution of 3.3 Å, however, the local resolution varies (Figs. 1A–C and S2A). In our map, most of the integrin headpiece has well resolved side chains for reliable atomic model building. In the αv subunit, the resolution of the thigh domain near the headpiece is sufficient to separate β -strands and to trace the backbone, but it becomes less well-resolved towards the lower leg. In the $\beta 8$ subunit, the resolution of the hybrid domain is also sufficient to trace the backbone of all β -strands with the bulky side chains resolved. However, the resolution of the PSI/EGF1 domain is significantly lower, suggesting that the linkage between the PSI/EGF1 and hybrid domains is flexible. Overall, our reconstruction suggests that there is considerable motion between the headpiece and the legs, which is consistent with our previous studies of the $\alpha v\beta 8$ ectodomain (Cormier et al., 2018).

The integrin-binding loop of L-TGF- $\beta 1$ is strikingly well resolved with clear density for the side chains suggesting that this binding interface between integrin $\alpha v\beta 8$ and L-TGF- $\beta 1$ is very stable. In comparison, the rest of L-TGF- $\beta 1$ is at a lower resolution (5–10 Å) suited for rigid body fitting suggesting that bound L-TGF- $\beta 1$ is flexible (Figs. 1B and 1C). Using the arm domain of L-TGF- β as a mainstay, the crystal structure of L-TGF- $\beta 1$ (PDB: 3RJR (Shi et al., 2011)) is well accommodated by our map (Fig. 1B).

The L-TGF- β RGD loop is comprised of two main components: the proximal RGD loop, which shows significant variance in length and sequence between TGF- $\beta 1$ and TGF- $\beta 3$, and the RGD LXXI/L integrin-binding motif, which is comprised of the tripeptide RGD recognition motif followed by a consensus accessory binding site (²¹⁸LATI²²¹ in L-TGF- $\beta 1$ and ²⁴⁴LGRL²⁴⁷ in L-TGF- $\beta 3$) (Mu et al., 2002; Ozawa et al., 2016). From here on we refer to the RGD LXXI/L consensus motif as the integrin-binding motif. A crystal structure of L-TGF- β alone shows proximal loop is disordered, and LXXI/L motif forms a loop (Shi et al., 2011). When crystallized in complex with $\alpha v\beta 6$, the LXXI/L motif organized into a short one and half turn α -helix (Dong et al., 2014; Dong et al., 2017), which we will refer to as the integrin-binding helix.

In our map, the density of the entire integrin-binding loop is well resolved. The RGD sequence is located in the ligand-binding cleft immediately followed by the integrin binding

helix (Fig. 1D). Overall, the integrin-binding motif and helix and the $\alpha\beta$ 8 ligand-binding cleft are similar to the crystal structures of the $\alpha\beta$ 6 headpiece in complex with L-TGF- β 1 or with a L-TGF- β 3 RGD-peptide (Fig. S3) (Dong et al., 2014; Dong et al., 2017).

SDL2 is a loop between the α 1 and α 2 helices of β 8 and is located at the back of the ligand binding cleft. A disulfide bond is formed between two conserved residues (C169 and C176) forming the SDL2 C-C loop. The SDL2 C-C loop bends inwards towards the integrin binding helix of L-TGF- β 1 and positions its Y172 to form a hydrophobic patch with multiple residues of SDL1–3 (Figs. 1E, S3E). The Arg^{RGD} interacts with the sidechain of α D218. The Asp^{RGD}, together with S114 and S116 of the SDL1 α 1-helix, coordinate the MIDAS cation consistent with other α v-integrins in complex with RGD (Fig. S3J–M) (Takagi, 2007).

The $\alpha\beta$ 8/C6D4 structure reveals the unliganded integrin binding pocket

In our previous ligand-free structure of $\alpha\beta$ 8, we found that SDL2 was flexible, and therefore could not be modeled (Cormier et al., 2018). To define the structure of $\alpha\beta$ 8 in the ligand-free state, we used the Fab domain of C6D4 to capture SDL2 in a specific conformation. C6D4 is a potent neutralizing antibody that we previously developed to bind to the $\alpha\beta$ 8 headpiece with high affinity and to effectively inhibit binding of L-TGF- β (Takasaka et al., 2018). To further facilitate high-resolution structure determination, we added an additional Fab, 11D12V2, that binds the α v-thigh. Since 11D12V2 is used solely for facilitating alignment and does not impact L-TGF- β binding nor headpiece conformation, for simplicity we refer to this complex as $\alpha\beta$ 8/C6D4. We determined a cryo-EM structure of $\alpha\beta$ 8/C6D4 in an extended conformation at an overall nominal resolution of 3.9 Å. With further focused alignment on the region of interest, which includes the headpiece and the variable domain of C6D4 Fab, the resolution was improved to 3.5 Å (Figs. 2A, 2B, S3C). The variable domains of C6D4 contact all SDL loops and D218 of the α v-head, thus directly competing with L-TGF- β (Fig. 2B, 3, S3C, Table S1, S3 and S4).

Although C6D4 interacts with the ligand-binding pocket, it does not participate in coordination of the MIDAS cation (Fig. 2B). Comparison between the $\alpha\beta$ 8/L-TGF- β 1 and $\alpha\beta$ 8/C6D4 structures shows that upon ligand binding the S116 at the tip of the SDL1 α 1-helix position shifts slightly towards the ligand-binding pocket (~1 Å) but the majority of the SDL1 α 1-helix itself remains at a similar position (Fig. 2C, S3G, S3R and S3S). This result is surprising because other integrin structures show both larger shifts and straightening of the SDL1 α 1-helix upon ligand binding (Fig. S3N–S) (Dong et al., 2014; Dong et al., 2017; Zhu et al., 2013).

The differences in SDL2 positioning are the largest overall conformational differences between the $\alpha\beta$ 8/L-TGF- β and $\alpha\beta$ 8/C6D4 complexes. C6D4 captures the SDL2 C-C loop in an outward position through interactions with complementary determining regions (CDR) CDR_{L1,L3} and CDR_{H1,H3} (Fig. 2D, 3 and Table S1). Notably, the inhibitory Fab Act-1 captures the β 7 integrin SDL2 C-C loop in the same position, indicating that SDL2 can assume similar ligand free conformations in other integrins (Fig. S3T and S3U) (Yu et al., 2012). Overall, our structure demonstrates that C6D4 captures $\alpha\beta$ 8 in a conformation corresponding to an unliganded state and completely occludes the ligand binding cleft. Thus,

it provides a clear mechanism of inhibition and explains the potency of C6D4 *in vitro* and *in vivo* (Takasaka et al., 2018).

An intermediate $\alpha v\beta 8$ binding state

We hypothesized that the inward position of the SDL2 C-C loop in the $\alpha v\beta 8$ /L-TGF- β structure is a critical feature that facilitates formation of the integrin-binding helix, which is necessary for positioning of the RGD sequence into the $\alpha v\beta 8$ ligand binding pocket. To address this hypothesis, we took advantage of the binding footprint of C6D4, which covers SDL1, 2 and 3 in addition to having its CDR_{L1} extended into the ligand-binding cleft. It is therefore possible to exchange the CDR_{L1} with the RGD loop of L-TGF- β while still maintaining binding of the Fab to $\alpha v\beta 8$. This would allow visualization of the RGD loop in the ligand binding cleft with SDL2 in an inactive conformation. We designed a chimeric Fab with the CDR_{L1} loop replaced by the RGD loop of L-TGF- $\beta 3$. This antibody (C6-RGD3) binds to and blocks function of $\alpha v\beta 8$ (Fig. S3V–W). We determined the structure of $\alpha v\beta 8$ in complex with C6-RGD3 and 11D12V2 Fabs to a resolution of 3.9Å (Figs. 2E, S2C and S3D). The angle and position at which the C6-RGD3 binds integrin is very similar to C6D4 with the major differences in the engineered CDR_{L1} loop (Fig. 2E–G, 3, Table S2, S3, and S4). The conformation of this loop is distinct from the canonical position of the RGD in the $\alpha v\beta 8$ /L-TGF- $\beta 1$ structure with Arg^{RGD} in a pocket formed by SDL2/3, which we will refer to as the proximal SDL2/3 pocket, where it interacts with D¹⁴⁸ of the αv -head (Fig. 2F, S3D and 3SF). This residue is conserved in $\alpha 8$, the only other integrin α -subunit that binds to L-TGF- β , suggesting it may play a key role in L-TGF- β binding (Table S3). In the $\alpha v\beta 8$ /C6-RGD3 structure, the integrin-binding helix does not form and SDL2 remains in an outward position as in the $\alpha v\beta 8$ /C6D4 complex (Fig. 2G, 2H, and S3D). In contrast, the position of the SDL1 $\alpha 1$ -helix is similar to its position in $\alpha v\beta 8$ /L-TGF- β and the Asp^{RGD} coordinates with the MIDAS cation (Fig. 2F, 4 and S3H, S3R, S3S). These features suggest that this is an intermediate state of L-TGF- β binding and its biological relevance is tested below.

C6-RGD3 binds efficiently to $\alpha v\beta 8$ and $\alpha v\beta 6$, but not $\alpha v\beta 1$, $\alpha v\beta 3$, or $\alpha v\beta 5$, mirroring the binding preference of these integrins to L-TGF- β (Fig. S3W). Since C6D4 does not bind to $\alpha v\beta 6$, we assume that C6-RGD3 binding to $\alpha v\beta 6$ is entirely through canonical positioning of the integrin-binding loop of CDR_{L1}. We reasoned that the C6-RGD3 CDR_{L1} loop could also reach a canonical conformation with $\alpha v\beta 8$ without a contribution from rest of the Fab binding footprint. Indeed, upon intensive classification, we identified a minor population (~6% of particles) that showed the C6-RGD3 integrin-binding motif in a canonical binding position with helix formation (Fig. S4). These results suggest that the C6-RGD3 Fab is able to bind to $\alpha v\beta 8$ in two distinct modes, one reflecting an alternative L-TGF- β RGD position with no integrin-binding helix formation and no contact with the $\beta 8$ SDL2, and the other, a canonical positioning of L-TGF- β RGD with the integrin-binding helix.

$\alpha v\beta 8$ lacks the ADMIDAS cation

When $\alpha v\beta 8$ is bound with L-TGF- β the position of the SDL1 $\alpha 1$ -helix tip is similar to the ligand-bound state of other integrins. In other integrins, the inward motion of the SDL1 $\alpha 1$ -helix is accompanied by swinging out of the hybrid domain, a process that is inhibited by Ca²⁺ and stimulated by Mn²⁺ (Zhu et al., 2013). We previously found that $\alpha v\beta 8$ binding to

L-TGF- β is not significantly enhanced by Mn^{2+} over Ca^{2+} and that hybrid domain swing out is not seen in Mn^{2+} containing buffers, suggesting that metal-binding properties are different for $\alpha v\beta 8$ than other integrins studied so far (Minagawa et al., 2014). Existing crystal structures of integrin ligand binding pockets show that three adjacent conserved cation-binding sites SyMBS, MIDAS and ADMIDAS, are coordinated by residues in SDL1 and SDL3 (Takagi, 2007; Xiong et al., 2001). Such cation coordination is widely thought to be a key determinant of integrin binding affinity and conformational dynamics (Takagi, 2007). Although coordinating residues are conserved across all β -subunits in the MIDAS and SyMBS sites, in the $\beta 8$ ADMIDAS site, two Asn residues replace otherwise conserved adjacent Asp residues. We find no density for the ADMIDAS cation in any of our structures while densities corresponding to the SyMBS and MIDAS cations are clearly observed (Figs. 4A–H). The absence of an ADMIDAS in $\alpha v\beta 8$ provides a structural explanation for the lack of cation preference and movement of SDL1 $\alpha 1$ -helix upon L-TGF- β binding, which are expected features in other integrins where conformational coupling between the hybrid domain and ligand binding were observed.

Flexibility of L-TGF- β when bound to $\alpha v\beta 8$

To gain further insight into conformational flexibility of L-TGF- $\beta 1$ bound to $\alpha v\beta 8$, we analyzed the complete $\alpha v\beta 8$ /L-TGF- $\beta 1$ dataset through comprehensive classification (Fig. S5). We define seven subclasses that represent conformational snapshots of the movements adopted between L-TGF- $\beta 1$ and the integrin headpiece. Each structure exhibits high resolution in the integrin headpiece and binding interface (3.3–3.6 Å) and a well-defined shape for L-TGF- $\beta 1$ prodomain. This conformational lineup shows different degrees of rotation between L-TGF- $\beta 1$ and the $\alpha v\beta 8$ headpiece (Fig. 5). In all seven classes the key structural elements that mediate major interactions between $\alpha v\beta 8$ and L-TGF- β are similar to those shown in Fig. 1. The proximal RGD loop of L-TGF- β is not as well resolved in most of these structures, and the distal linker that follows the integrin binding helix is resolved but varies considerably from subclass to subclass. This suggests that the proximal RGD loop and the distal sequence act as flexible linkers, which allow the rest of L-TGF- β to tilt around the $\alpha v\beta 8$ headpiece in a wide range. This allows the arm domain of L-TGF- β to make contacts with the $\beta 8$ headpiece in some of the subclasses. The overall density of the prodomain and TGF- β itself varies noticeably indicating a range of plasticity throughout L-TGF- β (Fig. 5).

The position of the SDL1 $\alpha 1$ -helix is the same and no movement of the $\beta 8$ hybrid domain is observed in any subclass. The lower leg domains in these subclasses are similarly and consistently at a lower resolution than the rest of the complex. This is in line with what we observed previously and suggests that movement in the lower legs is not correlated with the motion of bound L-TGF- β (Cormier et al., 2018). Such flexibilities in both $\alpha v\beta 8$ and L-TGF- β are unlikely to generate any force to liberate the TGF- β from the latent complex. Therefore, our structural data reveal a model where TGF- β can be activated without being released from the latent complex. In such a model, the structural flexibility of L-TGF- β upon binding to $\alpha v\beta 8$ would be sufficient to expose the active domain of TGF- β to its receptors.

Dynamic interactions within the ligand binding pocket determine $\alpha v\beta 8$ specificity for L-TGF- β

Differences in the proximal SDL2/3 binding pocket and the SDL2 C-C loop define three distinct conformational states captured in our structures. We hypothesized that a series of sequential restructuring steps of the L-TGF- β integrin-binding motif and $\alpha v\beta 8$ headpiece lead to stable ligand binding (Fig. 6A). To test this model, we carried out a series of mutations. To ensure that all mutants were properly folded, we confirmed that C6D4, which recognizes a non-linear epitope on the $\alpha v\beta 8$ headpiece (Takasaka et al., 2018), binds to these mutants as expected (Fig. S6A).

To rule out the possibility that the alternative binding mode is an artifact of grafting the RGD L-TGF- β loop onto the C6D4 Fab, we mutated the SDL3 residue I208 to its corresponding Arg residue in $\beta 3$ (I208R). Based on our structure and hypothesis, this substitution would occlude the proximal SDL2/3 binding pocket and prevent the alternative binding mode, while not affecting the placement of RGD in the canonical binding position (Fig. S6E–G). The I208R mutant completely inhibited L-TGF- β binding and activation suggesting that the alternative RGD binding mode is a required initial or intermediate step in the L-TGF- β binding and activation process (Fig. 6B). Mutational validation of the importance of the proximal SDL2/3 pocket including I208 in ligand binding, reinforces the potential of this domain as a therapeutic target.

The differences in position of the SDL2 C-C loop in the $\alpha v\beta 8$ /C6D4 and $\alpha v\beta 8$ /RGD3 compared to the $\alpha v\beta 8$ /L-TGF- β structures suggest a critical role for Y172 in the formation of the L-TGF- β integrin-binding helix (Fig. 6C). The conservation of the hydrophobic patch in $\beta 6$ suggests a similar role for $\beta 6$ I183 (Fig. 6C). The positioning of the aromatic ring of Y172 in our structure, along with the knowledge that in $\alpha v\beta 6$ the corresponding residue is an Ile (I183) suggest that the hydrophobic interaction is imperative for L-TGF- β integrin-binding helix formation (Fig. 3). In order to test the importance of this residue we made the following substitution mutants: one conservative, Y172M, and one non-conservative, Y172N. As expected, the conservative substitution, Y172M did not affect L-TGF- β activation. Replacing Y172 with the polar Asn significantly reduced L-TGF- β activation to a level similar to when the entire SDL2 region is removed (Fig. 6B) (Dong et al., 2014). A secreted version of $\alpha v\beta 8$ expressing Y172A dramatically reduced L-TGF- β binding (Fig. S6C).

The highly flexible SDL2 is poorly conserved between integrin b-subunits and provides the main $\beta 8$ residues of the hydrophobic patch interacting with the L-TGF- β helix. Despite large sequence variation in SDL2 between the $\beta 6$ and $\beta 8$ subunits, the footprint of hydrophobic interactions that promotes stabilization of the L-TGF- β helix appears to be absolutely conserved (Fig. 6C) (Dong et al., 2014). As a test of the generalizability of our model to the $\alpha v\beta 6$ integrin, we made the non-conservative mutation of $\beta 6$ I183N, which corresponds to $\beta 8$ Y172N (Fig. 3). As expected I183N dramatically inhibits $\alpha v\beta 6$ -mediated L-TGF- β activation and binding (Fig. 6B, S6D). This suggests that the $\beta 6$ and $\beta 8$ SDL2 have similar roles.

Our data reveal two main features that are required for efficient L-TGF- β binding: a conserved hydrophobic residue in the $\beta 8$ SDL3 I208 position of the proximal SDL2/3 pocket and a bulky hydrophobic residue at the tip of the SDL2 C-C loop. The $\beta 3$ and $\beta 5$ integrins contain a charged residue in the $\beta 8$ SDL3 I208 position, while integrin $\beta 1$, $\beta 2$, $\beta 4$ and $\beta 7$ subunits do not have a bulky hydrophobic residue in the position of $\beta 8$ SDL2 Y172 (Fig. 3). Thus, the absence of one or the other features explain the relatively inefficient L-TGF- β binding to these integrins (Dong et al., 2014).

Previous studies showed that peptides derived from the TGF- $\beta 3$ compared to the TGF- $\beta 1$ RGD β LXXI/L sequence have increased ability to inhibit L-TGF- β binding to $\alpha v\beta 6$ (Dong et al., 2014). Because the formation of integrin binding helix is a critical step in L-TGF- β binding to integrin, the ability of the LXXI/L motif to form a helix may underlie this difference. Therefore, we synthesized a series of peptide mutants that were designed to differ in their predicted abilities to form a helix. As expected, the ability to form a helix correlated with the IC₅₀ of each peptide and the peptides with proline substitutions in the terminal hydrophobic position of LXXI/L that were designed to disrupt the helix had the highest IC₅₀. Similarly, the TGF- $\beta 3$ peptide has a lower IC₅₀ than the TGF- $\beta 1$ peptide because TGF- $\beta 3$ peptide has a higher propensity to form a helix (Fig. 6D). Overall, our findings suggest that conformational dynamics within the L-TGF- β integrin-binding helix are critical for binding ability to $\alpha v\beta 8$.

Integrin $\alpha v\beta 8$ -mediated TGF- β activation does not require TGF- β release

In the latent form, the active domain of TGF- β is shielded by a flexible loop of the prodomain known as the latency lasso. Without extensive conformational changes and cytoskeletal linkage, force cannot be transmitted through the $\beta 8$ subunit to release TGF- β from its latent complex. However, the flexible nature of L-TGF- β we observed in our structure may expose the active domain of mature TGF- β without releasing it.

A recent crystal structure of the L-TGF- β /GARP complex reveals how GARP functions as a stable platform for L-TGF- β presentation, which suggests that a complex of $\alpha v\beta 8$ with L-TGF- β /GARP forms between two opposing cells with one presenting $\alpha v\beta 8$ and the other presenting L-TGF- β (Lienart et al., 2018). Furthermore, it is already known that L-TGF- β /GARP presenting immune cells also present its receptor, TGF- β R2 (Li et al., 2006). We thus envision a new TGF- β activation model in which L-TGF- β binding to $\alpha v\beta 8$ exposes the active domain of mature TGF- β so that it can bind to TGF- β R2, recruit a type I TGF- β R (i.e. ALK5) to form the minimal TGF- β signaling module to initiate the TGF- β signaling cascade (Huang et al., 2011). We generated a structural model using our cryo-EM structure of $\alpha v\beta 8$ /L-TGF- β with crystal structures of L-TGF- β /GARP (Lienart et al., 2018) and TGF- β in complex with TGF- β R2 (Groppe et al., 2008; Hart et al., 2002; Radaev et al., 2010) (Fig. 7A). In this multicomponent model, TGF- β signaling can occur within the confines of L-TGF- β and release and diffusion of TGF- β is not required.

To test this hypothesis, we sought to express a form of TGF- β that could not be released from L-TGF- β in a model system that reproduces a physiologic milieu where $\alpha v\beta 8$ normally activates TGF- β . Stimulated T-cells present L-TGF- β on their cell surfaces via covalent linkage to transmembrane adaptor proteins such as GARP and demonstrate TGF- β -

dependent differentiation into regulatory T-cells (Treg) (Edwards, et al, 2014). These findings indicate that cell surface L-TGF- β can be activated to interact with its own TGF- β receptors in a T cell-intrinsic mechanism of TGF- β activation and signaling (Stockis, et al, 2012). The integrin $\alpha v \beta 8$ either provided from contacting cells or Treg cells themselves has been implicated in stimulating such cell-intrinsic T-cell differentiation (Takasaka, et al, 2018; Edwards, et al, 2014). No reporter system currently exists to investigate the mechanistic basis of cell-intrinsic TGF- β activation, whereby the L-TGF- β presenting cell is also the cell that responds to TGF- β signaling. The most widely used in vitro TGF- β activation system relies on co-culturing a highly sensitive and specific TGF- β reporter cell line (TMLC) with integrin-expressing and/or L-TGF- β /GARP presenting cells (Wang, et al 2012). In such systems, TMLC must be in contact with these other cell-types for detection of TGF- β activation (Munger, et al, Cell, 1999). TMLC cells are a stable subclone of mink lung epithelial cells (MLEC) stably transfected with an expression construct containing a TGF- β specific promoter fragment, consisting of a truncated plasminogen activator inhibitor-1 (PAI-1) promoter, fused to the firefly luciferase reporter gene (Abe, et al, 1994). TMLC cells are widely used to measure active TGF- β because of their low background, specificity for TGF- β and abundant expression of TGF- β receptors and downstream signaling molecules that allow measurement of dose-dependent increases in TGF- β concentration in the physiologic range (Abe, et al, 1994). However, TMLC cells cannot report cell-intrinsic TGF- β activation since they do not present L-TGF- β themselves.

To build a cell-intrinsic TGF- β activation system, TMLC cells were stably transfected with wild-type (WT) TGF- β . Without co-transfecting GARP, TMLC do not present L-TGF- β on their cell surface. When co-transfected with TGF- β and GARP, high levels of cell surface expression of L-TGF- β can be detected (Fig S7A, C). To build a cell-intrinsic TGF- β activation system, which express a non-releasable form of TGF- β , we similarly expressed the L-TGF- β (R249A)/GARP complex on the surface of TGF- β reporter cells (TMLC) in which the L-TGF- β carries the same mutation (R249A) as described above that prevents cleavage of TGF- β from its prodomain during biosynthesis (Fig S7B, D). We verified that the transfected L-TGF- β (R249A)/GARP reporter cells expressed the cell surface complex at equivalent levels as the WT L-TGF- β /GARP reporter cells (Fig. S7C, D), and confirmed the absence of cleaved mature TGF- β from LAP in the R249A mutant (Fig. S7G). The resulting cell lines were used to determine cell-intrinsic TGF- β activation and signaling and as well as the role for release of mature TGF- β .

TMLC non-transfected or transfected with WT L-TGF- β , L-TGF- β (R249A), WT L-TGF- β /GARP, or L-TGF- β (R249A)/GARP were plated on the immobilized $\alpha v \beta 8$ ectodomain, or control substrates ($\alpha v \beta 3$ as a low-affinity TGF- β binding integrin, an antibody to LAP as a high-affinity L-TGF- β binder, or BSA) (Fig. S7E). In this configuration, actin cytoskeletal force cannot be generated or applied from the integrin to cause release of TGF- β . Robust TGF- β activation was observed at similar levels using cells expressing L-TGF- β (R249A)/GARP or WT L-TGF- β /GARP when plated on $\alpha v \beta 8$, while no activation was seen when these same cells were plated on control substrates (Fig. 7B). Activation in this system was significantly enhanced by cell surface presentation of L-TGF- β by GARP since minimal TGF- β activation was seen when TMLC transfected with WT L-TGF- β or L-TGF- β (R249A) without GARP were plated on $\alpha v \beta 8$ (Fig. S7F). The ability of GARP to present L-

TGF- β and to increase efficiency of integrin-mediated TGF- β activation is consistent with previous findings (Wang, et al, 2012). No active TGF- β was detected in the supernatant from reporter cells expressing either WT or L-TGF- β (R249A)/GARP when plated on α v β 8 (Fig. 7B). These results demonstrate that α v β 8-dependent TGF- β activation can occur independently of actin-cytoskeletal force and does not require release of mature TGF- β (Fig. 7C).

DISCUSSION

Our studies reveal an intermediate step of L-TGF- β binding to α v β 8, and the flexible nature of L-TGF- β in complex with α v β 8. Together with mutagenesis and cell-based assays, we reveal an unanticipated mechanism for TGF- β activation, where mature TGF- β signals within the confines of L-TGF- β . Thus, α v β 8-mediated TGF- β activation does not require actin-cytoskeletal force nor release and diffusion of mature TGF- β . This model differs from all other activation models that require release and diffusion of mature TGF- β (Annes et al., 2003; Massagué, 2012; Moses et al., 2016). Instead, α v β 8-mediated TGF- β activation directs TGF- β signaling to the opposing L-TGF- β /GARP expressing cell through formation of a large multicomponent cell-cell protein complex.

Protein inhibitors in clinical development that globally inhibit mature TGF- β or TGF- β receptors have been associated with toxicities (Akhurst, 2017; Tolcher et al., 2017). Furthermore, our model predicts that such global TGF- β inhibition strategies would be ineffective in blocking α v β 8 mediated TGF- β activation due to the inaccessibility of their targeted epitopes within the geometrically constrained cell-cell α v β 8/L-TGF- β /GARP complex. In contrast, targeting α v β 8 prevents the complex from forming, which would increase specificity and efficacy while mitigating the risks of global TGF- β inhibition.

This TGF- β activation mechanism provides the structural basis to explain why conditional deletion of α v β 8 on one cell type primarily affects another (Arnold et al., 2019; Kitamura et al., 2011; Mohammed et al., 2016; Travis et al., 2007). This mechanism also explains why deletion of GARP, or its homologue, on immune cells negatively affects their TGF- β -dependent differentiation state (Qin et al., 2018; Salem et al., 2019; Wallace et al., 2018). In tumors, we have found that α v β 8 expressing tumor cells direct the immunosuppressive differentiation of L-TGF- β /GARP expressing Treg and myeloid cells (Takasaka et al., 2018).

Inhibiting α v β 8 in tumors can specifically target local immunosuppressive mechanisms of tumor immune evasion by blocking α v β 8-mediated contact of tumor cells with immune cells (Takasaka et al., 2018). We have defined the mechanisms of action of two antibody prototypes that efficiently block binding of α v β 8 to L-TGF- β . One of these is ligand-mimetic and defines an alternative RGD binding mode that not only reveals the initial phase of ligand recognition for L-TGF- β , but also reveals the proximal SDL2/3 pocket as a possible therapeutic target for α v β 8. Based on the variation of this pocket between other α v-integrins, it is possible that small molecules designed to bind within this pocket could increase specificity, which has been difficult with small molecules that directly target the conserved RGD binding site (Hatley et al., 2018). After the initial phase of ligand binding, it is clear from our structures that SDL2 plays a prominent role in stable binding of α v β 8 to L-

TGF- β , a mechanism that we have experimentally extended to the α v β 6 integrin. We anticipate that the dynamic interactions occurring during binding between α v β 8 and L-TGF- β could provide a roadmap for understanding how flexible ligands conform to the SDL loops deep within the ligand binding clefts of other integrins as well.

In summary, our data provide unprecedented structural insights into the TGF- β activation process that expand the basic concepts of TGF- β activation for improved biologic understanding of TGF- β function in the immune and vascular systems and lay the foundation for structural investigations of other intact integrin ectodomains with their ligands to improve strategies to target those functions for therapeutic benefit.

STAR Methods

LEAD CONTACT AND MATERIALS AVAILABILITY

Further information and requests for reagents may be addressed to the Lead Contact, Stephen 5 Nishimura (stephen.nishimura@ucsf.edu). Antibodies, cell lines and plasmids used in this manuscript will be available from the Lead Contact upon execution of a materials transfer agreement.

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Human Subjects—Adult airways were collected from first through fourth order bronchi from lobectomy specimens from resections performed for primary lung cancer. Informed consent was obtained from all surgical participants as part of an approved ongoing research protocol by the University of California San Francisco Committee on Human Research, in full accordance with the declaration of Helsinki principles. The specimens were de-identified.

Normal bronchial epithelial cells (HBEC)—Normal human bronchial epithelial cells (HBEC) were isolated from human lung specimens, as previously described (Araya et al., 2007). Specifically, airway epithelium was stripped freshly from bronchi from surgical specimens and after washing in PBS with dithiothreitol (5 mM) were digested overnight at 4°C with a protease XIV solution (0.4 mg/ml, Sigma). After washing with bronchial epithelial growth medium (BEGM, Lonza) with 2.5% FCS and titration, HBEC were plated onto rat-tail Col I-coated (10 μ g/ml, Corning, cat. no. 354236) dishes and incubated overnight; then the medium was changed to fresh BEGM. HBEC were passaged under conditional reprogramming conditions with BEGM with ROCK inhibitor (10 mM) and irradiated NIH3T3 fibroblasts, as described (Liu et al., 2012).

Human embryonic kidney cells (HEK293)—HEK293 are an authenticated female line from a commercial vendor (ATCC). HEK293 cells were grown in DMEM + 10% FBS + penicillin-streptomycin + amphotericin B + 0.11mg/mL Sodium Pyruvate + 2mM L-glutamine + MEM nonessential amino acids, cultured at 37°C with 5% CO₂. HEK293 and all other cell lines were routinely tested for mycoplasma contamination. For all established cell lines cell culture media and antibiotics were prepared by the University of California,

San Francisco Cell Culture Facility using deionized water and analytical grade reagents. Fetal calf serum was obtained from Invitrogen (Carlsbad, CA).

Chinese Hamster Ovary Cells (CHO)—CHO-K1 are an authenticated female line from a commercial vendor (ATCC). CHO Lec 3.2.8.1 have four independent mutations in the N- and O-glycosylation pathways producing proteins all of the high mannose glycosylation pattern (Stanley, 1989). CHO Lec 3.2.8.1 cells were grown in CHO-S-SFM II medium + 10% FBS + penicillin-streptomycin + amphotericin B, cultured at 37°C with 5% CO₂. CHO Lec 3.2.8.1 cells were provided by Pamela Stanley (Stanley, 1989). Authentication of CHO Lec 3.2.8.1 cells was performed by changes in glycosylation patterns as estimated by migration of secreted proteins by SDS-PAGE.

Transformed Mink Lung Epithelial Cells (TMLC)—TMLC is a stable mink lung epithelial reporter cell line derived from mixed sex American Mink fetal lung epithelial cells that stably expresses a portion of the plasminogen activator inhibitor 1 promoter (Abe et al., 1994). TMLC cells were grown in DMEM + 10% FBS + penicillin-streptomycin + amphotericin B, cultured at 37°C with 5% CO₂. TMLC are a gift from John Munger (NYU medical center, New York City, New York). Authentication was performed by increased luciferase activity in response to recombinant TGF- β .

METHOD DETAILS

Antibody isolation, characterization and production—The C6D4, 68, 8B8, 3G9, 1D11 antibodies have been previously described (Minagawa et al., 2014; Mu et al., 2002; Takasaka et al., 2018; Weinreb et al., 2004). Mouse anti-human α v, clone 11D12V2 was isolated as described, with the following modifications. Screening of hybridomas produced a polyclonal hybridoma 11D12 with reactivity against cell lines expressing the human α v-subunit (CHO- α v β 8, or SW480- β 6). Subcloning produced two clones 11D12V1 and 11D12V2 both with reactivity against the α v-subunit. Variable (V) genes were isolated, sequenced, and the V_H/V_K genes cloned into mouse IgG_{2a} expression vectors and stably transfected into CHO-K1 cells. C6-RGD3 was created by splice overlap extension (SOE) polymerase chain reaction (PCR) into the VL CDRL1 region of a C6D4 IgG_{2a} expression vector (Minagawa et al., 2014) with the following oligonucleotide primers using the C6D4 IgG_{2a} expression construct as a template: 5'-GATCTGGGGCGCCTCAAGAAGAACGCCTTGGCTTGGTACCAGCAG-3'; 5'-CTTGAGGCGCCCCAGATCTCCACGGCCGAGCAGACTCTGACTGGATTTG-3'. C6-RGD3 was transfected into CHO-K1 cells. Antibodies were produced and purified as described previously (Minagawa et al., 2014). Briefly, stable transfected CHO-K1 cells were grown in spinner cultures in CHO SFMII media with antibiotics. Antibodies were purified from culture media using protein G sepharose columns (HiTrap, GE Healthcare). C6D4, C6-RGD3, and 11D12V2 fragments were generated by papain digestion (Pierce) of the IgG followed by Protein-A Agarose (Millipore) incubation and to remove Fc fragments and intact antibodies and final separation using Mono S ion-exchange chromatography (GE Healthcare).

The 11D12V1 and 11D12V2 binding epitopes were estimated by negative staining electron microscopy (ns-EM), essentially as described below and in previous studies (Minagawa et al., 2014; Takasaka et al., 2018). Clone 11D12V1 bound to the α v-head and 11D12V2 to the α v-thigh. Anti- β 6 (MAB4155) was purchased (R&D Systems).

A bioassay to measure cell-intrinsic TGF- β signaling—To develop a cell intrinsic TGF- β activation system we used stable transfection of TMLC (Amaxa) with a vector containing either a WT human TGF- β 1 IRES GFP, or a human TGF- β 1 (R249A) IRES GFP cassette with puromycin resistance to obtain TMLC cells expressing WT L-TGF- β either capable of dissociating into LAP and mature TGF- β or not, due to the R249A mutation that normally allows furin cleavage of LAP from mature TGF- β (Shi, et al, 2011). Human TGF- β 1 IRES GFP or human TGF- β 1 (R249A) IRES GFP TMLC cells were sorted for equal expression using GFP fluorescence and did not present any L-TGF- β on their cell surface (Fig. S7). In contrast, stable transfection of these lines with a HA-GARP construct with a blastacidin resistance cassette followed by selection and sorting resulted in high surface expression of TGF- β 1/GARP or TGF- β 1 (R249A)/GARP, as measured by anti-HA (clone 5E11D8, GenScript, Piscataway, NJ) or anti LAP (R&D Systems, AF426). To confirm lack of releasable of TGF- β from the L-TGF- β R249A/GARP cell surface complex, L-TGF- β (R249A)/GARP expressing or WT L-TGF- β /GARP TMLC cells were surface biotinylated using EZ-link sulfo-NHS biotin (Thermo Fisher Scientific) and immunoprecipitated using anti-HA, resolved by 4–12% gradient SDS-PAGE, under non-reducing conditions, immunoblotted, probed with streptavidin-HRP and detected by chemiluminescence, essentially as described (Mu, et al, 2002). Immunoprecipitations confirmed association of GARP with WT L-TGF- β , L-TGF- β (R249A), and absence of cleavage of LAP from mature TGF- β in the L-TGF- β (R249A)/GARP TMLC cells (Fig. S7).

A TMLC assay to measure integrin-mediated cell-intrinsic TGF- β signaling—The α v β 8 ectodomain was coated along with the controls α v β 3 (R&D Systems), BSA (Sigma-Aldrich) or anti-LAP (R&D AF426, 1 μ g/ml) onto ELISA plates in PBS (1mM Ca²⁺ and 1mM Mg²⁺) 1 hour at RT. Wells were subsequently washed in PBS and blocked in PBS with 1% BSA WT L-TGF- β 1, L-TGF- β 1 (R249A), WT L-TGF- β /GARP, L-TGF- β (R249A)/GARP expressing TMLC cells were plated at a density of 1×10^5 cells/ml in basal media. After 16 hrs, media was removed and applied to wells containing TMLC reporter cells to measure diffusible mature TGF- β , which were incubated overnight prior to lysis and determination of luciferase activity (Promega) as reported (Mu, et al, 2002). To measure cell-intrinsic TGF- β 1 activation, the attached L-TGF- β 1, L-TGF- β 1 (R249A), WT L-TGF- β /GARP, L-TGF- β (R249A)/GARP or parental TMLC cells were lysed and assayed for luciferase activity. To facilitate comparison between different TMLC lines expressing WT L-TGF- β 1, L-TGF- β 1 (R249A), WT L-TGF- β /GARP, L-TGF- β (R249A)/GARP (or parental TMLC cells) normalized luciferase activity was expressed as activated TGF- β in pg/ml. Normalization was performed by interpolating luciferase activity against standard curves generated using each TMLC line with varying doses of recombinant human TGF- β 1 (R&D Systems).

Integrin DNA constructs—Wild-type and mutant recombinant human integrin $\alpha\text{v}\beta\text{8}$ and $\alpha\text{v}\beta\text{6}$ truncated at the junction of the ectodomains and transmembrane domains in pcDNA1neo have been described (Nishimura et al., 1994; Weinreb et al., 2004). β8 SDL was prepared as described (Takasaka et al., 2018). To create mutant β8 subunits, SOE was performed using PCR with WT β8 as a template to create mutant constructs all in pcDNA6 V5HisA (Invitrogen) with a stop codon inserted before the V5/His tag. The following mutagenic primers (all in 5' to 3' orientation) were used: β8 I208R 5'-CAGAAGATCTCTGGAAACAGAGATACACC-3'; 5'-GAAGTTTGGTCGACATAATGC-3' β8 Y172N: 5'-CAATGCAGTGACAACAATTTAGACTGC-3', 5'-GCAGTCTAAATTGTTGTCACTGCATTG-3'; β8 Y172M: 5'-GATTCATAATCAATGCAGTGACATGAATTTAGACTGCATGCC-3', 5'-GGCATGCAGTCTAAATTCATGTCAGTGCATTGATTATGAATC-3'; β8 Y172A: 5'-GATTCATAATCAATGCAGTGACGCCAATTTAGACTGCATGCC-3', 5'-GGCATGCAGTCTAAATTGGCGTCACTGCATTGATTATGAATC-3'; β6 I183N: 5'-CCCTTGCAGTAGTAATCCATACTTCTG-3', 5'-CAGAAGTATGGATTACTACTGCAAGGG-3'.

TGF- β DNA constructs—Porcine L-TGF- β1 with a C4S mutation, to improve secretion and prevent association with L-TGF- β binding proteins, and an N-terminal cleavable (HRV 3C) 7x Histidine-streptavidin binding protein tag to facilitate purification (Shi et al., 2011), were joined by SOE PCR from pcDNA-GS-TGF- β1 (gift from Dr. Sun, National Institutes of Health, Bethesda, MD (Zou and Sun, 2004)) and subcloned into pcDNA6 (Invitrogen). Porcine L-TGF- β1 C4S pcDNA6 was modified using the following mutagenic primers, RGE (a mutation that disrupts the integrin binding recognition sequence on L-TGF- β and therefore minimizes the number of L-TGF- βs that bind two integrins simultaneously): 5'-CCGCCGGGGTGAAGTGGCCAC-3'; 5'-GTGGCCAGTTCACCCCGGCGG-3'; R249A: 5'-ACCTGCACAGTCCCGGCACCGCGCAGCCCTGG-3'. Wild type human TGF β1 was derived from human TGF β1 _pLX307 (Rosenbluh et al., 2016) by removing the C-terminal V5 tag using PCR with the following primers; 5'-ATGCCACCCCGCTGG-3', 5'-CTCTACTAGTCTCGAGTTATCAGCTGCACTTGCAGGAGCGCAC-3'. The WT human TGF- β1 IRES GFP cassette was then subcloned into a version of pcDNA6 (Invitrogen) with a puromycin resistance cassette which was amplified from TGF β1 _pLX307 using 5'-ATCGTTTCAGACCCACCTCCC-3' and 5'-CTCTGCTTAGCGAATTCGTTAACTGGCACCGGG-3'. A R249A mutant of this construct was produced using SOE PCR employing the following mutagenic primers; 5'-CACCGCGCAGCCCTGGACACCAAC-3', 5'-CCAGGGCTGCGCGGTGCCGGGAG-3'.

GARP DNA constructs—N-terminal HA tagged human GARP pcDNA3 (Cuende et al., 2015) was provided by Sophie Lucas (de Duve Institute, UCLouvain, Brussels, Belgium) and the entire HA-GARP reading frame transferred into pcDNA6 (Invitrogen). All constructs were verified by sequencing.

Secreted protein expression and purification—Integrin constructs were expressed using stably expressing CHOlec 3.2.8.1 cells grown in spinner cultures in CHO SFMII

(Thermo Fisher) with antibiotics, for structural studies, or transiently transfected in HEK293 cells using 293 SFMII (Thermo Fisher) for biochemical studies. Integrin purification was carried out by affinity chromatography using a Protein G-clone 8B8 column followed by size exclusion chromatography (Superdex 200 Increase 10/300 GL, GE Healthcare) in 20mM Tris-HCL pH 7.5, 150 mM NaCl, 1mM CaCl₂ and 1mM MgCl₂. To produce L-TGF- β for structural studies, 293 cells were transiently transfected with equal amounts of porcine L-TGF- β 1 C4S R249A RGD and C4S R249A RGE plasmids, to favor formation of L-TGF- β 1 with a single intact RGD binding site to favor 1:1 binding stoichiometry to reduce sample heterogeneity, and purified as described (Shi et al., 2011). Briefly, supernatants were collected from spinner cultures, clarified by centrifugation, filtered through a PES (polyethersulfone) membrane, 0.2 mm pore size (Millipore), concentrated, and purified using Ni-NTA agarose (Qiagen), washed with three column volumes of 0.6 M NaCl, 0.01 M Tris (pH 8.0) and eluted with 0.25 M imidazole in TBS. The was adjusted to pH 7.4 then applied to Strep-tactin agarose (IBA) (1 ml per 1 L of culture supernatant) and washed with TBS (pH 7.4). Tag was cleaved with recombinant His-tagged HRV-3C protease (Novagen, 100 U mg⁻¹, 1 mg ml⁻¹), diluted 20-fold in TBS (pH 7.4) with 10% glycerol, applied to the column, and incubated at 4 °C for 16 h. The flow-through was washed with two column volumes of TBS (pH 7.4), containing untagged proTGF- β 1, then concentrated using centrifugal concentrators (Millipore) to about 1 mg ml⁻¹ in 10 mM Tris (pH 7.5), 75 mM NaCl. The homogeneity and purity of all protein preparations were verified by SDS-PAGE stained with Coomassie blue and protein concentrations were measured by bicinchoninic acid assay (Pierce).

Cryo-EM sample preparation—To prepare integrin-Fab or integrin-L-TGF- β complexes, 100 mg of recombinant α v β 8 was incubated in a 2-fold molar excess of each Fab or L-TGF- β , incubated at room temperature for 30 min, subjected to size exclusion chromatography and concentrated to 6 to 9 mg/ml. For cryo-EM grid preparation, 2.5 μ L of purified α v β 8 complex were deposited onto Quantifoil grids. For the α v β 8/L-TGF- β -R249A at 0.25 mg/ml, a 400 mesh 1.2/1/3 holey carbon gold grid was used. For the α v β 8/L-TGF- β /RGD-RGE complex at 0.075 mg/mL, a 400 mesh 2/2 holey carbon copper grid that had been covered with a thin layer of graphene oxide was used. For the α v β 8/C6D4/11d12v2 and the α v β 8/C6-RGD3/11d12v2 complexes, both at 6.8 mg/mL, 300 mesh 1.2/1.3 holey carbon gold grids were used. Except for the graphene-coated grid, grids were glow-discharged for 60s at 15 mA prior to sample application and freezing. The α v β 8/L-TGF- β C4S R249A RGD and α v β 8/L-TGF- β C4S R249A RGD/RGE complexes were frozen using a FEI Vitrobot Mark IV using a 4 second blot time. The α v β 8/C6D4/11d12v2 and α v β 8/C6-RGD3/11d12v2 complexes were frozen using a FEI Vitrobot Mark III using a 3–4 second blotting time. All grids were frozen with 100% humidity at 20°C and plunge-frozen in liquid ethane cooled by liquid nitrogen.

Cryo-EM data acquisition—Four datasets were acquired on a FEI Titan Krios transmission electron microscope operated in nano-probe mode at 300 kV equipped with a Gatan Quantum GIF energy filter, operated in zero-loss mode with a slit width of 20 eV and a Gatan K2 Summit direct detector. Automated data collection was carried out using the SerialEM software (Mastrorade, 2005). Movies were recorded in super-resolution mode

with a super-resolution pixel size of 0.673 Å/px and a nominal magnification of 105kx at a dose rate of ~ 8 e⁻/px/s. For the $\alpha v\beta 8/L$ -TGF- β -R249A complex and the $\alpha v\beta 8/L$ TGF- β /RGD-RGE, each 16 second movie contained 80 frames of 200 ms each, which corresponds to a total dose of ~ 70 e⁻/Å². For the $\alpha v\beta 8/C6D4/11d12v2$ complex and the $\alpha v\beta 8/C6RGD3/11d12v2$ complex, each 12 second movie contained 60 frames of 200 ms, which corresponds to a total dose of ~ 50 e⁻/Å². Each dataset was collected in a single session with a nominal defocus range of 1.0 – 2.5 μ m under focus. Total micrographs collected for each dataset are as follows: $\alpha v\beta 8/L$ -TGF- β -R249A complex: 1684 micrographs; $\alpha v\beta 8/L$ -TGF- β -RGDRGE complex: 2682 micrographs; $\alpha v\beta 8/C6D4/11d12v2$ complex: 1644 micrographs; $\alpha v\beta 8/C6RGD3/11d12v2$ complex: 4033 micrographs.

Imaging Processing—Dose fractionated super-resolution image stacks were motion corrected and binned 2 × 2 by Fourier cropping using MotionCor2 (Zheng et al., 2017). Motion corrected sums without dose-weighting were used for contrast transfer function (CTF) determination using GCTF (Zhang, 2016) or CTFFIND4 (Rohou and Grigorieff, 2015). Particles were picked using the reference-free method using Gautomatch (<http://www.mrc-lmb.cam.ac.uk/kzhang/Gautomatch>) and boxed out using Relion 3.0 (Zivanov et al., 2018) with a box size of 300 pixels and binned to 64 pixels. After 2D alignment and classification was carried out using cryoSPARC (Punjani et al., 2017), selected particles were re-extracted in Relion 3.0 and binned to 128 pixels to generate *ab initio* initial models using cryoSPARC. 3D classification schemes are outlined for the $\alpha v\beta 8/L$ TGFB-R249A complex and the $\alpha v\beta 8/L$ -TGF- β -RGD-RGE complex in Fig. S5 and for the $\alpha v\beta 8/C6RGD3/11d12v2$ complex in Fig. S4. No 3D classification was used for the $\alpha v\beta 8/C6D4/11d12v2$ complex dataset. For all final maps, non-uniform refinement, local resolution refinement, local resolution estimation, sharpening, and local filtering was carried out using cryoSPARC to yield maps with a final pixel size of 1.345 Å/px. The number of particles contributing to the final maps are as follows: $\alpha v\beta 8/L$ -TGF- β (conformation iv): 43,600 particles; $\alpha v\beta 8/C6D4/11d12v$: 84,266 particles; $\alpha v\beta 8/C6$ -RGD3/11d12v2: 221,159 particles. Images were rendered using UCSF Chimera (Pettersen et al., 2004) and PyMol (DeLano, 2002).

Model Building and Refinement—The atomic model of the αv headpiece from the crystal structure of $\alpha v\beta 6$ (PDB: 4UM8) with glycans removed was fitted to the cryo-EM map as a rigid body. An atomic model of the $\beta 8$ headpiece was generated by rigid body fitting of a homology model based on the same crystal structure (4UM8) using Modeller (Webb and Sali, 2014) and adjusted around the SyMBS cation area using the crystal structure of $\alpha v\beta 3$ (PDB: 3IJE), then fitted into the cryo-EM density map as a rigid body. Atomic models of Fabs were generated with RosettaAntibody using multiple-template grafting and H3 loop modelization (Lyskov et al., 2013) based on the primary sequence of their V_H/V_K. An atomic model of the arm domain of L-TGF- β 1 was generated from the crystal structure of L-TGF- β 1 bound to $\alpha v\beta 6$ (PDB: 5FFO). The models for Fab C6D4 and Fab C6-RGD3 or L-TGF- β 1 were then fitted as a rigid body to the map. Prototypical CHOLec3.2.8.1 glycans were added back to the model at the solvent exposed N-glycosylation consensus sites using GLYCAM (Singh et al., 2016). The sugar base of glycans were trimmed to fit into the corresponding densities and further refined in Rosetta

using a dedicated protocol that uses physically realistic geometries based on prior knowledge of saccharide chemical properties (Frenz et al., 2019). After rigid body fitting, all models were manually adjusted to fit the cryo-EM density maps in COOT (Emsley et al., 2010), followed by real space refinement using Phenix (Adams et al., 2010), and Rosetta (Wang et al., 2016). All modeling was aided by using EM maps that were focused on specific regions, as well as sharpened and unsharpened maps. All maps used for modeling have been deposited.

Antibody binding assays—ELISA plates were coated with recombinant α v-integrins (1 mg/ml coating concentration, all from R&D systems) blocked with 5% BSA in PBS for 1 hour, and antibodies allowed to bind for 2 hours at RT and detected with anti-mouse-HRP.

Cell adhesion assays—ELISA plates were coated with integrins (2 mg/ml coating concentration) and blocked with PBS with 5% BSA for 1 hour and then CHO-GARP/L-TGF- β , or CHO mock (5×10^5) transfectants in the presence of various concentrations of C6D4, C6-RGD3 or 3G9 were centrifuged onto integrin coated wells ($10 \times g$) for 5 min allowed to adhere for 30 min at RT after which the plates were inverted and centrifuged ($10 \times g$) for 5 min and then immediately fixed and stained (1% formaldehyde, 20% MeOH, 0.5% crystal violet) and after extensive washing, dye was solubilized in PBS with 1% T-X100 for 1 hr at RT and attached cells estimated by absorbance (A_{595}).

L-TGF- β 1 binding assays—ELISA plates were coated with recombinant porcine L-TGF- β 1 (0.5 μ g/ml coating concentration), blocked with 5% BSA in PBS for 1 hour, and integrins at various concentrations were allowed to bind for 2 hours at RT, and detected with 8B8 antibody and anti-mouse-HRP.

Cell staining, flow cytometry and cell sorting—Cell staining for L-TGF- β 1 was confirmed by flow cytometry using anti-LAP (R&D biotinylated anti-LAP BAF246 and streptavidin APC) and GARP cell surface expression was confirmed by anti-HA staining (clone 5E11D8 (Genscript, Piscataway, NJ)). EGFP expression was also used as a surrogate marker for L-TGF- β 1 expression. High-expressing pools of expressing cells were established by sorting (BD FACSAria, BD Biosciences, US).

Peptide competition assays—For peptide competition assays, ELISA plates were coated with recombinant porcine L-TGF- β 1 (0.5 μ g/ml coating concentration), blocked with 5% BSA in PBS for 1 hour, and integrins at 0.5 μ g/ml were pre-incubated with peptides at various concentrations for 20 minutes, allowed to bind for 2 hours at RT, and detected with 8B8 antibody and anti-mouse-HRP.

Sequence alignments—Multiple protein sequence alignments for integrins were generated using Clustal Omega (Madeira et al., 2019).

QUANTIFICATION AND STATISTICAL ANALYSIS

ELISA and TMLC assays are reported as means \pm s.e.m. All assays were repeated a minimum of 3 times. All statistical analyses were performed using the software package Prism 7 (GraphPad Software, San Diego, CA).

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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HIGHLIGHTS:

- Integrin $\alpha v \beta 8$ in the extended-closed conformation binds to and activates L-TGF- β
- The L-TGF- β RGD loop initiates contact with $\alpha v \beta 8$ through an alternate binding mode
- L-TGF- β is flexible when bound to $\alpha v \beta 8$
- Mature TGF- β activates signaling without being released from the latent TGF- β complex

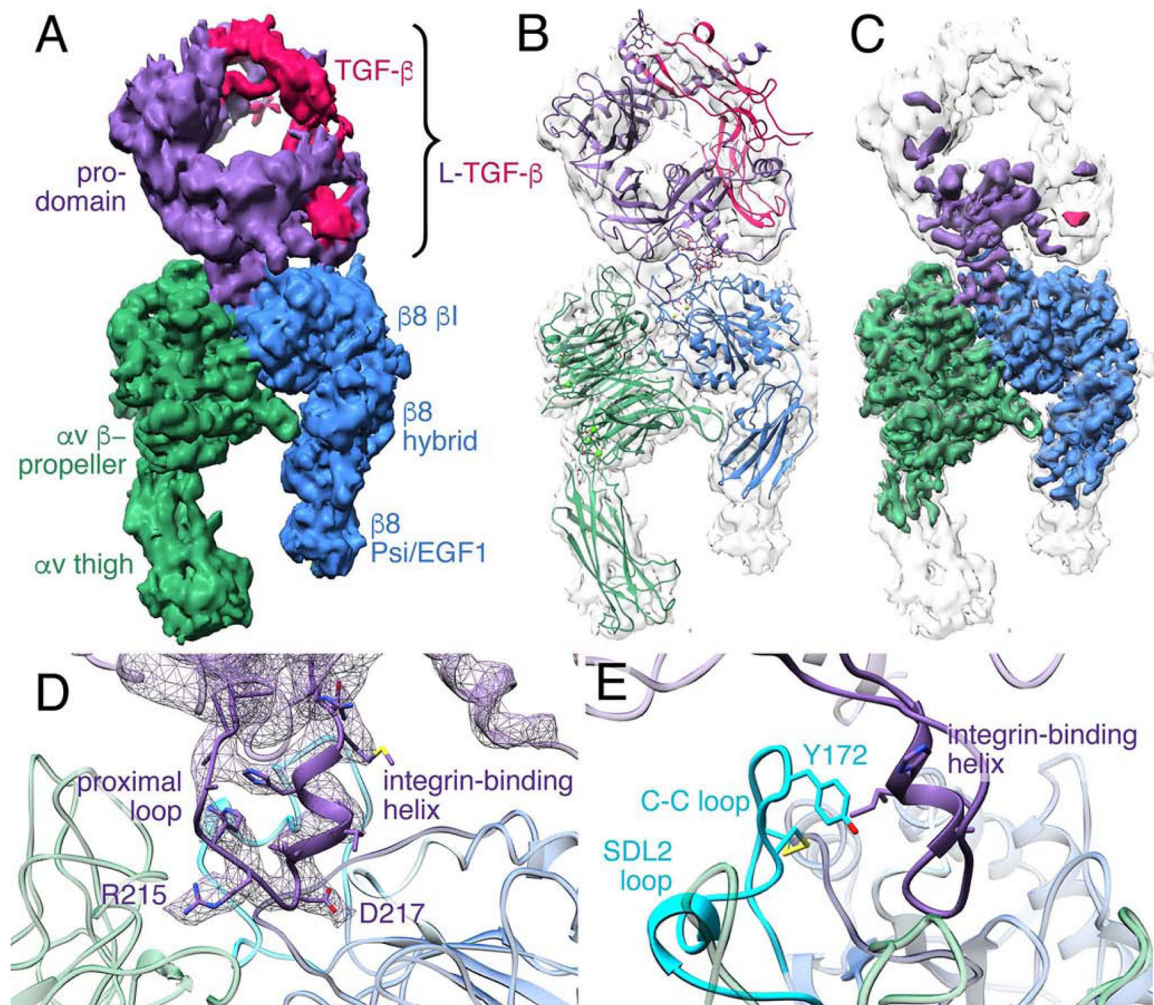


Fig. 1. The $\alpha\text{v}\beta\text{8}$ integrin ectodomain bound to L-TGF- β1

(A) Cryo-EM density map of $\alpha\text{v}\beta\text{8}$ integrin ectodomain with L-TGF- β1 bound. The map is displayed as unsharpened and at a low-threshold. The color code is as follows: integrin αv -subunit is green, integrin β8 -subunit is blue, prodomain of L-TGF- β1 is purple, and mature TGF- β1 is red. (B) Ribbon diagram of $\alpha\text{v}\beta\text{8}$ /L-TGF- β1 displayed within the density map shown in (A). (C) The sharpened cryo-EM map is shown in color, superimposed with the unsharpened map in semi-transparent white. (D) A ribbon diagram shows a close-up view of the binding interface between $\alpha\text{v}\beta\text{8}$ and L-TGF- β1 . The EM density of the integrin-binding motif of L-TGF- β1 , including proximal loop, RGD motif, and integrin-binding helix is shown in purple mesh. (E) Ribbon diagram showing the C-C loop within the SDL2 loop of the β8 subunit (cyan) relative to the L-TGF- β1 integrin-binding helix. Related information is in Fig. S1, S3.

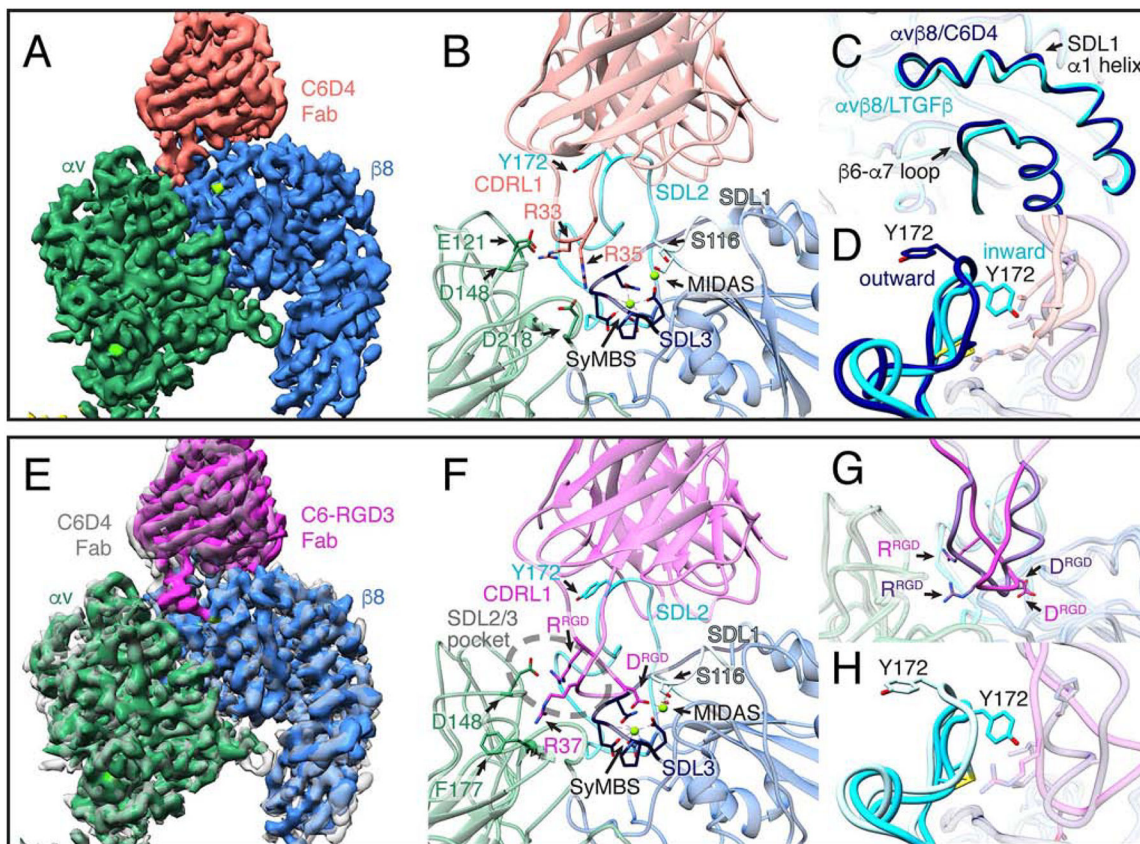


Fig. 2. Mechanisms of action of inhibitory C6D4 and C6-RGD3 Fabs

(A) A close-up view of the cryo-EM density map of the $\alpha v\beta 8$ /C6D4 complex. The full structure is shown in Fig. S2. The $\alpha v\beta 8$ ligand-binding cleft is fully occupied by C6D4, demonstrating a steric mode of inhibition. Color code: αv is dark green, $\beta 8$ is blue, and cations are bright green. (B) Close-up ribbon diagram of the $\alpha v\beta 8$ /C6D4 interface showing the position of the C6D4 Fab (coral) on the $\alpha v\beta 8$ binding-cleft, highlighting interactions between the Fab CDR_{L1} loop R33 interacting with αv E121 and αv D148, and Fab R35 with αv D218, respectively (for all interactions see Fig. 3, S1, S3 and S4). Additional color code: SyMBS and MIDAS is green, SDL1 is light blue, SDL2 is cyan, SDL3 is dark blue. (C-D) Comparisons are shown between $\alpha v\beta 8$ /C6D4 and $\alpha v\beta 8$ /L-TGF- $\beta 1$ with key changes occurring in $\beta 8$ upon ligand binding noted. Color code: unliganded C6D4 structure is dark blue, liganded L-TGF- β structure is cyan. There are very subtle ligand-induced changes in the SDL1 $\alpha 1$ helix (~ 1 Å at the tip) and the $\beta 6$ - $\alpha 7$ loop (C). Y172 of the SDL2 loop is positioned outward relative to the ligand binding site in complex with C6D4 but moves inwards upon ligand binding (D). (E) Cryo-EM density map of $\alpha v\beta 8$ (same colors as in A) in complex with the C6-RGD3 ligand-mimetic Fab (magenta) superimposed on the $\alpha v\beta 8$ /C6D4 map (semi-transparent gray). Aligning the αv -subunit β -propeller domain of the two maps shows a slight shift in the angle of C6-RGD3 Fab binding. (F) Close-up ribbon diagram of the $\alpha v\beta 8$ /C6-RGD3 binding interface shows the alternative RGD binding mode with C6-RGD3 Fab. Key contacts are: Arg^{RGD} interacting with αv D148 at the edge of the $\beta 8$ proximal SDL2/3 pocket (dashed ellipse) and C6-RGD3 Fab Asp^{RGD} interacting with the MIDAS cation. C6-RGD3 Fab R37 (³⁰-LGRGDLGRL⁻³⁸) interacts with αv F177 (for all

interactions see Fig. 3, S2–4). Integrin color codes are as in B, with C6-RGD3 in magenta. (G) Superimposition of the C6-RGD3 ligand-mimetic RGD loop (magenta) and the native L-TGF- β loop (purple) shows that C6-RGD3 binds in an alternative binding conformation. (H) $\beta 8$ Y172 in the $\alpha v\beta 8$ /C6-RGD3 complex (light blue) is in the outward position compared to the $\alpha v\beta 8$ /LTGFb complex (cyan). Related information is in Fig. S2–4.

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SDL1			SDL2			SDL3		
C6-RGD3			L ₃ L ₂ H ₃			H ₃ H ₃ H ₁ H ₁		
C6D4			L ₂ L ₃ H ₂ H ₁			H ₁		
β8	112	DVS AS MHNN	β8	152	KTVSPYISIH-PERIH NCSDY N--LDCMPH	β8	201	RQKISGN IDTPE
β6	123	DLSAS M DDD	β6	163	KPVSPFVKTT-PEEIANPC SSIP --YFCLPTF	β6	212	NQKISANIDTPE
β3	119	DLSYS M KDD	β3	159	KPVSPYMYIS PE ALENPCYDMK--TTCLPMF	β3	211	NQSVSRNRDAPE
β5	112	DLSLS M KDD	β5	162	KDISPFSYTA-PRYQTNPCIGYKLF P NCVPSF	β5	213	KQRVSRNRDAPE
β1	130	DLSYS M KDD	β1	170	KTVMPISTT-PAKLRN P CTSE--QNCTTPF	β1	218	KQRISGNLDSPE
β2	112	DLSYS M LDD	β2	152	KTVLFPV N TH-PDKLRN P C N KE--KECQPPF	β2	201	KQLISGNLDAPE
β7	140	DLSYS M KDD	β7	181	KTVLFPVSTV-PSKLRH P CPTRL--ERCQSPF	β7	229	RQSVSGNLD S PE
β4	108	DFS NS MSDD	β4	150	KVSV P Q T DMR-PEK L KE F WP-----NSDPPF	β4	197	GERISGNLDAPE
		* *						*
		* *						*
		* *						*
		= = = =			= = = =			= = = =

Fig. 3. L-TGF-β and inhibitory antibodies contact multiple overlapping residues in β8 SDL loops
 Sequence alignments for integrin b subunits are shown in decreasing order of homology to β8. Each SDL region is indicated above the sequence. Each CDR V_H and V_L loop of C6-RGD3 and C6D4 is shown above the specific amino acids which it contacts. The β8 SDL residues which contact L-TGF-β as well as C6D4 or C6-RGD3 are shown in green letters. The β8 SDL residues which form the hydrophobic patch that interact with the integrin binding helix of L-TGF-β1 are shown in orange letters. The residues that are mutated in Fig. 5 are shown in red letters. The cation binding sites are indicated below the sequence by colored asterisks with MIDAS in blue, ADMIDAS in magenta, and SyMBS in black. Conserved residues are indicated (=). Related information is in Table S1–4.

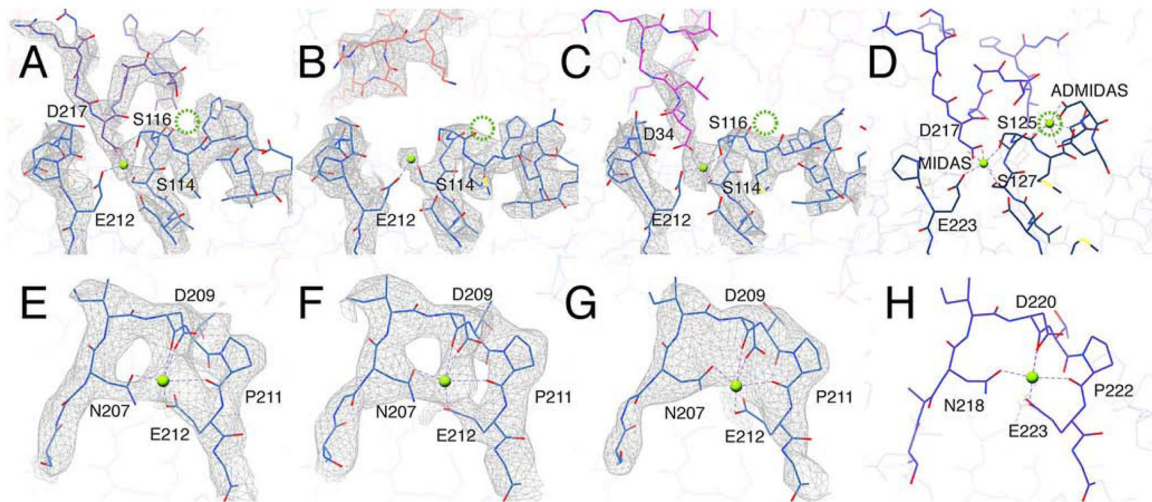


Fig. 4. The ADMIDAS cation is not present in $\alpha v \beta 8$

(A-C) The MIDAS cation binding site formed in $\alpha v \beta 8$ integrin (blue) bound with: L-TGF- $\beta 1$ (purple, A) C6D4 (coral, B) and C6-RGD3 (magenta, C). In all three structures there is clear density for the MIDAS cation (bright green). No density for the ADMIDAS cation is observed (the expected position for ADMIDAS is noted by the dotted green circle). Shown are the cation-coordinating residues of the integrin b subunit or the Asp of RGD of L-TGF- β (D217) or C6-RGD3 (D34). (D) The same binding interface in the $\alpha v \beta 6$ /LTGF β crystal structure (5FFO (Dong et al., 2017)). (E-H) Maps and models of the SyMBS cation in $\alpha v \beta 8$ /L-TGF- β (E), $\alpha v \beta 8$ /C6D4 (F), $\alpha v \beta 8$ /C6-RGD3 (G), and $\alpha v \beta 6$ /LTGF β crystal structure (5FFO) (H). All panels are color coded as in Fig. 2. Related information is in Fig. S3.

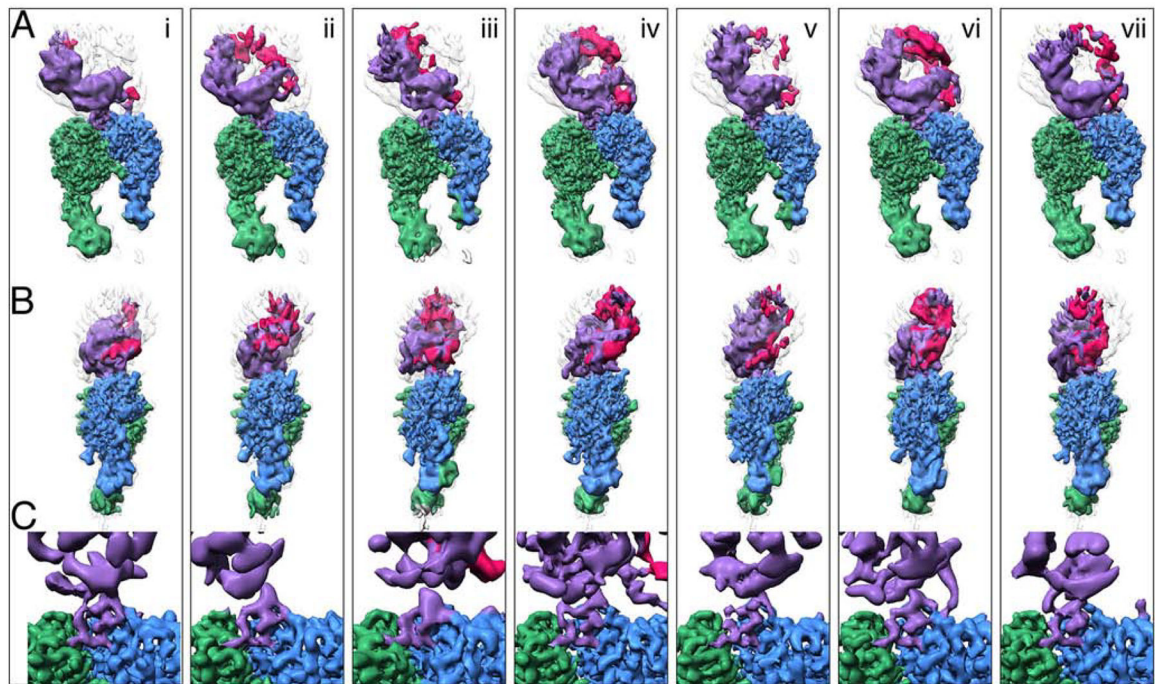


Fig. 5. L-TGF- β is flexible when bound to $\alpha v\beta 8$

Seven structures (i-vii) illustrate the conformational variability of the $\alpha v\beta 8$ /LTGF β complex. All structures are aligned to each other using the αv -subunit b-propeller domain. Each vertical panel shows two views (A, and B) of the same structure rotated 90° around the z-axis and a close-up of the integrin-ligand binding interface with the L-TGF- β integrin-binding helix (C), which is formed in all structures. Subclass (iv) is shown in Figs. 1, 2 and 4. Color coding is the same as for Fig. 2. For (A-B) the primary conformation is shown in colors, as defined in Fig. 1A, and all six other conformations are shown as semi-transparent to provide a reference for L-TGF- β 's range of motion. From left to right, the percentage of particles that went into each class are as follows: 14%, 14%, 14%, 19%, 16%, 11%, 12%. Related information is in Fig. S5.

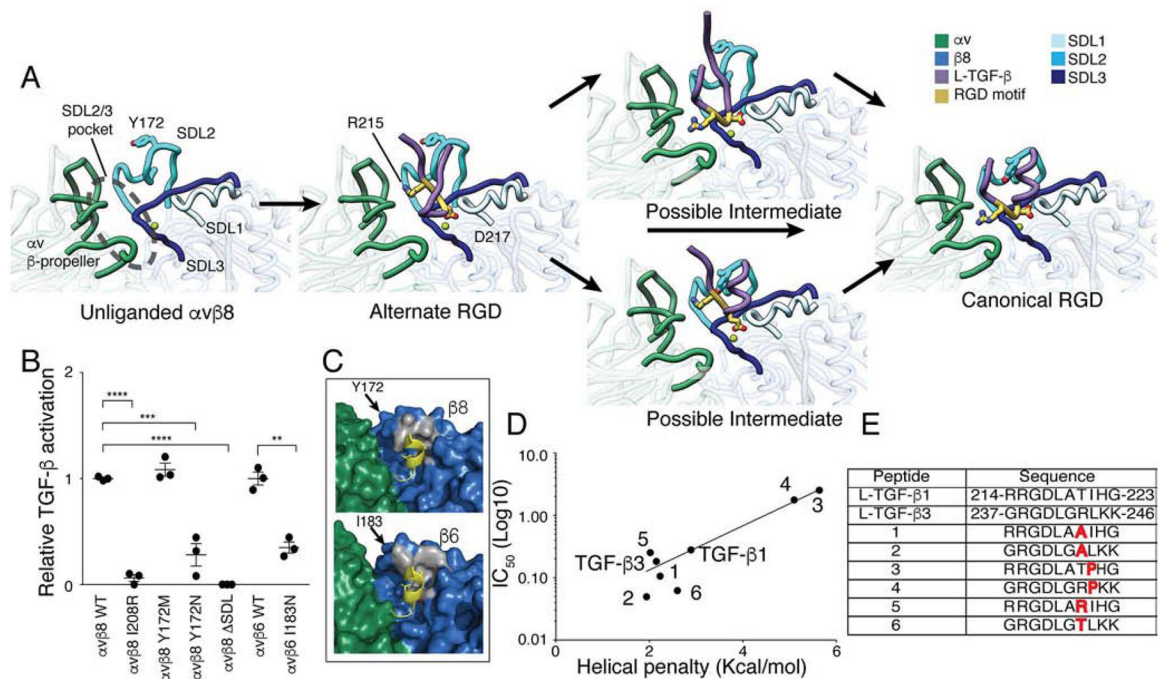


Fig. 6. Structure based modeling of L-TGF- β binding to α v β 8

(A) A stepwise model of L-TGF- β binding to α v β 8. Our structures were used to define unliganded (α v β 8/C6D4), alternate (α v β 8/C6-RGD3) and canonical RGD binding modes (α v β 8/L-TGF- β). The hypothesized transitions between these binding states suggest possible intermediates where the integrin binding helix either forms before or after engaging the canonical RGD position (color coding as in Fig. 1). (B) TGF- β activation assays test the function of β 8- and β 6-subunit SDL mutants. Wild-type (WT) α v β 8 or α v β 6 or mutant receptors (β 8: I208R, Y172M, Y182N, entire SDL2 deletion mutant (-SDL), or β 6: I183N) were stably expressed in CHO cells, sorted for uniform expression, confirmed for proper folding using C6D4 or anti- β 6 antibodies (Fig. S6) and tested for ability to support TGF- β activation with TMLC reporter cells. Activation relative to α v β 8 WT expressing CHO cells is shown. (C) Shown is surface rendering of the b-subunit hydrophobic patch (grey) of α v β 8 (upper) and α v β 6 (lower) and the relationship of β 8 Y172 and β 6 I183 with the L-TGF- β integrin-binding helix (yellow sticks and cartoon). (D) Inhibition of L-TGF- β 1 binding to α v β 8 by TGF- β 1 and TGF- β 3 integrin-binding motif peptides and Ala and Pro mutants was measured. Shown is the inhibitory concentration (IC_{50}) plotted against helical penalty score (Pace and Scholtz, 1998). (E) Sequences of peptides used in (D) are shown. ** $p=0.01$, *** $p=0.001$, **** $p=0.0001$ by unpaired two-sided Student's t test. Related information is in Fig. S6.

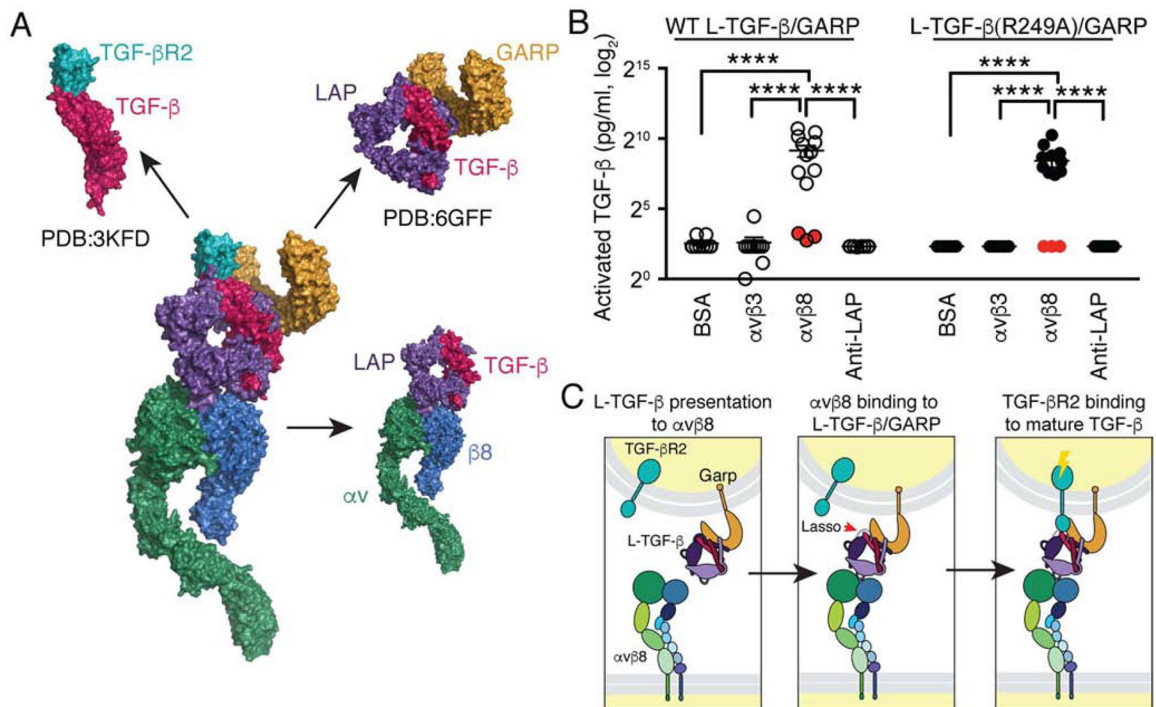


Fig. 7. A structural model of the $\alpha v\beta 8$ /L-TGF- β /GARP/TGF- β R2 complex predicts that releasing of mature TGF- β is not required for $\alpha v\beta 8$ -mediated TGF- β activation

(A) Surface representation of a model of the putative complex derived from our $\alpha v\beta 8$ /L-TGF- β 1 structures combined with the crystal structures of the GARP/L-TGF- β 1 complex (PDB: 6GFF (Lienart et al., 2018)), and the TGF- β /TGF- β R2 complex (PDB: 3KFD (Radaev et al., 2010)) is shown. Color code: integrin $\alpha v\beta 8$ (αv -green, $\beta 8$ -blue), L-TGF- β 1 (LAP-purple, TGF- β -red), GARP-orange and TGF- β R2-teal. (B) WT L-TGF- β 1/GARP or L-TGF- β 1(R249A)/GARP expressed on the surface of TGF- β reporter cells (TMLC) show efficient TGF- β activation when plated on immobilized $\alpha v\beta 8$. TMLC are stably transfected with a highly specific TGF- β responsive promoter driving a truncated plasminogen activator inhibitor type I (PAI-1) promoter, which contains SMAD-binding elements driving a luciferase reporter cassette (Abe, et al, 1994). TGF- β activation is then measured as luciferase activity. A R249A mutation in TGF- β prevents proteolytic cleavage by furin of mature TGF- β from LAP during intracellular processing (Shi, et al, 2011). TMLC expressing L-TGF- β 1 (R249A)/GARP cannot produce diffusible active TGF- β and thus report only TGF- β activated within the L-TGF- β 1 complex (Fig. S7). Minimal or undetectable levels of TGF- β were released into the supernatants from WT L-TGF- β 1/GARP or L-TGF- β 1 (R249A)/GARP TMLC cells when plated on $\alpha v\beta 8$ or control substrates (supernatants were removed and assayed separately using TMLC cells, filled red circles). Luciferase activities are normalized and converted to amount of active TGF- β (pg/ml), as described (Annes et al., 2003). The y-axis is in \log_2 scale. **** $p < 0.0001$ by one-way ANOVA and Tukey's multiple comparisons test. (C) Model of TGF- β mediated TGF- β activation is shown. Activation occurs in three steps: Left panel: Initially, $\alpha v\beta 8$ in an extended-closed conformation surveys the environment for cells presenting L-TGF- β on their cell surface by co-expression of adaptor molecules such as GARP. Middle panel:

Binding of one cell expressing $\alpha v\beta 8$ to a L-TGF- β expressing cell increases the flexibility of the latency lasso of TGF- β (red arrow) which exposes the active domain on one fingertip of the mature TGF- β homodimer. Right panel: TGF- β R2 expressed by the same cell presenting L-TGF- β binds to the exposed active domain of mature TGF- β , which initiates the TGF- β signaling cascade. Related information is in Fig. S7.

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