

Mutational landscape of antibody variable domains reveals a switch modulating the interdomain conformational dynamics and antigen binding

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Somatic mutations within the antibody variable domains are critical to the immense capacity of the immune repertoire. Here, via a deep mutational scan, we dissect how mutations at all positions of the variable domains of a high-affinity anti-VEGF antibody G6.31 impact its antigen-binding function. The resulting mutational landscape demonstrates that large portions of antibody variable domain positions are open to mutation, and that beneficial mutations can be found throughout the variable domains. We determine the role of one antigen-distal light chain position 83, demonstrating that mutation at this site optimizes both antigen affinity and thermostability by modulating the interdomain conformational dynamics of the antigen-binding fragment. Furthermore, by analyzing a large number of human antibody sequences and structures, we demonstrate that somatic mutations occur frequently at position 83, with corresponding domain conformations observed for G6.31. Therefore, the modulation of interdomain dynamics represents an important mechanism during antibody maturation in vivo.

antibody | affinity maturation | immunology | deep mutational scanning | conformational dynamics

Unlike T-cell receptors, B-cell-derived antibodies accumulate somatic mutations in the antigen-binding variable domains of heavy chain (HC) and light chain (LC), in both the complementarity-determining regions (CDRs) and the framework regions (FWRs) that scaffold the CDRs (1, 2). Together with the V(D)J gene recombination, somatic hypermutation (SHM) adds to the gene diversity of antibody variable domains (V_H and V_L) necessary for efficient immune recognition (3).

Deciphering the molecular mechanism of SHMs in improving antibodies is critical for understanding the evolution of the immune repertoire. It is well recognized that mutations at the CDRs or FWR structurally adjacent to the CDRs can modulate and optimize the antigen-binding interface (4–7), and several studies have used saturated mutagenesis of CDR position to explore the effect of various mutations on affinity and specificity (8–11).

The role of SHM in antigen-distal FWR is less well understood, however. Previous studies have suggested that the antigen-distal FWR contributes primarily to the variable domain structural integrity; thus, mutations at these positions would be either detrimental to or neutral for antigen-binding function (12–14). Other authors have characterized the potential of antigen-distal framework mutation as beneficial for antigen-binding function. For example, framework mutations can compensate for the destabilizing effect of mutations at CDRs needed for antigen binding (15). In other cases, non-CDR SHMs have been shown to be required for the neutralization activity of broadly neutralizing anti-HIV antibodies (16).

Thus far, the roles of SHM have been studied primarily by examining the functional consequences of reverting the somatic mutation of selected antibodies back to the germline sequences (15–21). Although these studies have demonstrated that antibodies

depend on somatic mutations to achieve high-affinity antigen binding, the approach is limited to the small set of mutations present in a given antibody.

Here we took a systematic approach to assessing the mutability of the whole variable domain for maintaining or improving folding stability and antigen binding. We used a well-optimized anti-vascular endothelial cell growth factor (VEGF), G6.31, with high affinity ($K_d = 0.4$ nM) and excellent thermostability [fragment antigen-binding (Fab) melting temperature (T_m) = 84 °C]. G6.31 derives its HC and LC variable domains from the frequently used human germline V families V_H3 and V_{K1} , respectively, and exhibits the canonical variable domain backbone structure (22–24), and thus is a good representative of human antibodies.

The present study had two aims: first, to map the mutational landscape of the two variable domains to delineate the potential effects of all possible single mutations, and second, to identify novel molecular mechanisms of antigen distal framework mutations for improving antibody function. A high-affinity antibody such as G6.31 likely has well-optimized CDRs, thus providing an opportunity to identify beneficial antigen-distal mutations.

Significance

On encountering antigens, antibody's variable domains evolve and mature through multiple rounds of somatic mutation. By surveying the effects of all possible single mutations of an antibody's variable domains on folding stability and antigen binding, we showed that even for a high-affinity antibody such as the one used in our study, beneficial mutations can be found both near and far from the antigen-binding site. Furthermore, our mutational scan revealed an antigen-distal framework position working as a structural switch whereby a mutation can allow the variable domains to sample different conformational spaces and thus potentially different binding functions. This understanding of the mechanism of somatic mutation in human antibodies illustrates how antibodies efficiently use somatic mutation for evolving diversity in immune recognition.

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We generated a phage display library of G6.31 Fab variants with all possible single mutations throughout the variable HC (V_H library) and LC (V_L library) domains. We compared the frequency of each mutation in the library before and after selection for binding to its various binding partners, including protein A, protein L, and VEGF, using deep sequencing (25, 26). We found that a large number of single mutations in the framework outside the structural core and the V_H - V_L interface are well tolerated. We identified mutations at positions distal to CDRs that can further improve the antigen-binding affinity and/or thermostability of this well-optimized

antibody, and examined the molecular mechanism of mutations at LC-83 in modulating Fab stability and antigen binding. We found that approximately 25 Å away from the antigen-binding site, the mutation at LC-83 can alter the predominant orientation of the constant domains (C_L and C_{H1}) to the variable domains V_L and C_L (the Fab elbow angle), as well as the orientations of the two variable domains toward each other (the torsion angle between V_L and V_H). We also found that somatic mutations at LC-83 are common among human antibodies of the V_{Kappa1} germline origin. Our work suggests that antigen-distal framework SHMs occurring during antibody maturation can

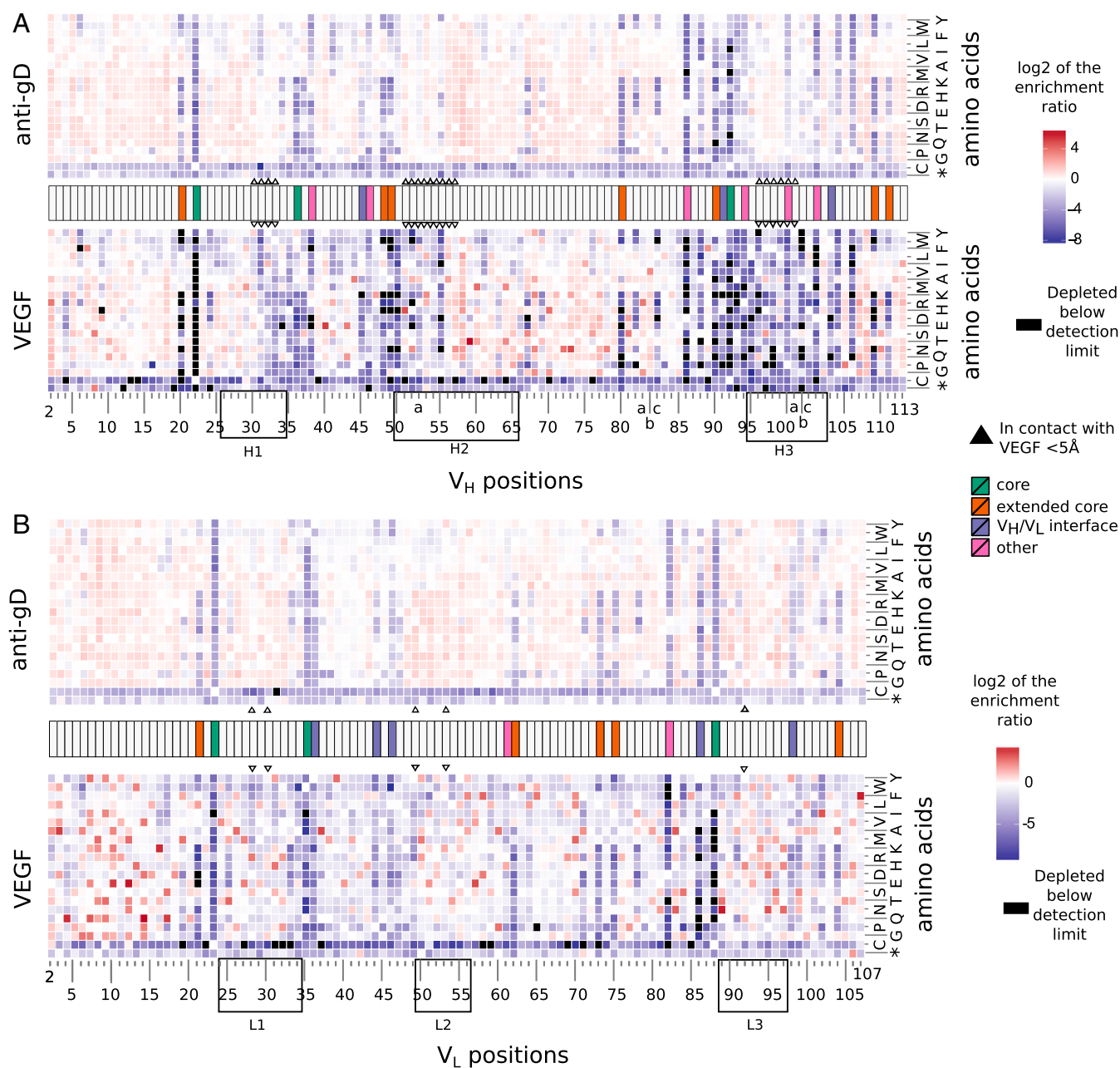


Fig. 1. Mutational landscape of V_H and V_L . The change of the abundance of each mutation in the V_H (A) and V_L (B) during selection for stable expression using anti-gD tag or for binding antigen VEGF (Fig. S1A) is calculated as \log_2ER and depicted in the heat maps with a scale of red (enriched) to blue (depleted). All mutations were accounted for in the preselection libraries. Mutations in black are those that are absent after selection indicating strong depletion. Highly conserved positions of V_H or V_L are colored based on their structural role, including hydrophobic core (colored green), extended core (orange), V_H - V_L interface (purple), or other (pink; hydrogen-bonded cluster) (Figs. S1–S4). CDR positions are boxed. Residues that contact VEGF ($\leq 5\text{\AA}$ in distance based on the G6/VEGF complex structure; PDB ID code 2FJG) are marked with arrowheads.

improve antibody properties by optimizing interdomain structural dynamics.

Results and Discussion

High Tolerance to Single Mutations in V_L and V_H for Stable Fab Assembly. We displayed G6.31 Fab on the surface of M13 phage by coexpressing the LC that is C-terminally fused to a 14-mer peptide expression tag gD and the HC up to the C_{H1} domain, which is then fused to the C-terminal domain of M13 p3 minor coat protein (Fig. 1 and Fig. S1) (24). We generated a single-site saturated mutagenesis library of both variable domains. The V_L library and V_H library mutated V_L positions 2–107 and V_H positions 2–113 (Kabat numbering), respectively, one residue at a time, to all 20 amino acids. The first position was omitted to avoid the effect of mutation on signal peptide processing. To detect the impact of mutation on functional expression and fold stability, we panned both libraries against an anti-gD antibody, protein A, or protein L, which binds the LC tag, the side of V_H , the side of V_L , respectively, to enrich for well-expressed Fab variants (Fig. 1A and Figs. S1 and S2). We previously showed that detection of the gD tag on the LC indicates stably assembled Fab with properly folded V_L and V_H , because only the HC is physically linked to the phage coat (27). The sequences from the selected and unselected library pools were determined using Illumina MiSeq pair-end reads spanning ~300 base pairs in each direction, as described previously (26).

From the 10^5 – 10^6 reads per sample, we verified that all expected mutations were found in the unselected pools, and further filtered the sequences to reduce biases from PCR or sequencing (Methods and Table S1). We then calculated the enrichment ratio (ER) for each mutation by dividing the frequency in the selected pools by the frequency in the unselected pools, as described previously (25, 26). The ER of a particular mutation likely aggregates several effects, which include the impact of the mutation on functional expression, the stability of the fold as well as the binding affinity of the ligands in selection.

The ERs of all single mutations in V_H library and V_L library from panning with three different ligands (anti-gD, protein A, and protein L) yielded similar and consistent results demonstrating the validity of the approach (Figs. S1–S3). The ERs of mutations in V_H library from anti-gD and protein L panning were closely correlated ($r^2 = 0.98$). The ERs from the protein A and protein L-panned V_H were also correlated ($r^2 = 0.611$) with the differential enrichment attributed to the subset of V_H positions binding to protein A (28). A similar pattern was seen for the V_L library panning data.

We chose the ER dataset from anti-gD panning to represent folding selection, and used a heat map to display the patterns of enrichment (red) or depletion (blue) for each mutation (Fig. 1 and Figs. S1 and S2). Enrichment indicates that the mutation is likely beneficial for the selected function (stable expression), whereas depletion indicates unfavorable mutations. Overall, the vast majority of positions appear to be tolerant to either the full range of mutations or a set of conserved mutations, because mutations at these sites do not exhibit overt depletion (Fig. 1 and Figs. S1, S2, and S4).

Consistent with what is known about the folding of the antibody variable domain (7, 13, 14, 29, 30), we found a set of mainly structurally buried framework positions resisting mutation (Fig. S4). These “conserved” framework positions include the hydrophobic core and V_H – V_L interface residues, as well as the residues involved in a hydrogen bond network at the bottom half of the variable domains. Mutations at some of the hydrogen bond network positions, like LC-R61 and LC-D82, have been shown to induce aggregation and fibril formation in LCs (31). Furthermore, a few CDR positions (e.g., R94, P100, D101 in the CDR-H3) also exhibited low tolerance to mutation, suggesting their role in overall stability. R94 and D101 are known to be important

for the “kinked” structure of CDR-H3 (32), whereas HC-P100 may be structurally important for the V_H – V_L interface as it packs against LC-Y49 in the G6 structure.

Antigen-Distal Framework Mutations That Improve G6.31 Affinity. The V_L and V_H libraries were also selected against VEGF to obtain an overview of the effects of single substitutions on antigen binding (Fig. 1). Compared with the selection for stable expression using anti-gD, protein A, or protein L, greater stringency selection was used to enrich clones that not only bind VEGF, but bind with high affinity. Thus, the extents of enrichment/depletion obtained were more prominent. We first note that except for the CDR positions expected to be important for VEGF interaction, the ER profile was strikingly similar to that observed from the selection for functional folding and expression (Fig. 1). This is consistent with the fact that stable expression is a prerequisite for antigen binding. Along with confirming the mutational landscape for stable expression, the prominent ERs obtained from VEGF panning showed that VEGF binding for G6.31 is a HC CDR-centric event (Fig. 1), consistent with the finding that HC CDRs account for most of the structural contact with VEGF (22). Collectively, the ERs showed a bimodal distribution (Fig. 2). Whereas most mutations appear to have a neutral or negative impact on binding or folding, many potentially beneficial mutations are located in CDRs as well as in nonconserved regions of the framework. Similar bimodal distributions have been observed in deep mutagenesis scanning experiments, suggesting that this is a frequent fitness landscape for protein interactions (33).

We next expressed and purified several Fab variants based on the ERs obtained, which should reasonably classify the mutation

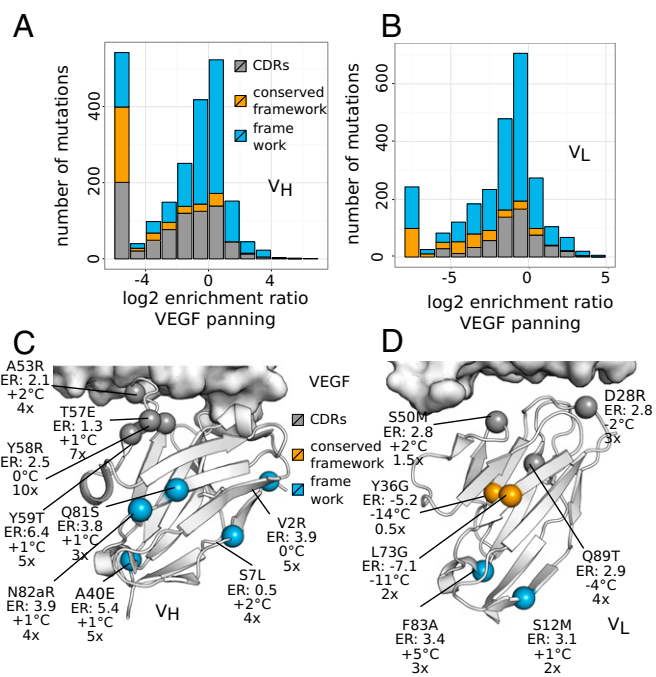


Fig. 2. Identification of VEGF affinity-improving mutations in G6.31. Shown is the distribution of log₂ERs obtained from VEGF panning of the V_H library (A) and the V_L library (B). Mutations that are depleted beyond the detection limit are set to the maximum observed depletion. Mutations are color-coded according to their location in the CDRs (gray), in the conserved framework as identified in Fig. 1 and Fig. S4 (yellow), or in the rest of the framework (blue). (C and D) Using the same color scheme, selected mutations are mapped on the G6/VEGF complex structure (PDB ID code 2FJG), and the change in T_m (°C) and the fold improvement in antigen-binding affinity (K_d) compared with G6.31 Fab are shown.

as favorable or unfavorable for ligand binding and/or thermostability (Fig. S5A and B), and measured their affinity for VEGF using Biacore and their thermostability using differential scanning calorimetry (DSF). As expected, two examples of highly depleted mutations in the conserved framework (LC-Y36G and LC-L73G) exhibited greatly reduced thermostability (with a drop in T_m of 14 °C and 11 °C, respectively) but with intact antigen-binding affinity (0.5- and 2-fold improvement). Thus, depletion of such mutants is an effect of unstable folding, which clearly dominates over the selection for VEGF binding.

To identify antigen-distal mutations that improve the antibody, we mapped 16 mainly beneficial mutations (9 in V_H and 7 in V_L) on the G6/VEGF complex structure (Fig. 2C and D). Whereas two CDR mutations (HC-T57E and HC-Y58R) resulted in the largest increase in VEGF binding affinity (7- to 10-fold), many antigen-distal framework mutations also produced a significant increase in affinity (2- to 5-fold). We found LC-F83 particularly interesting, because it locates 25 Å away from the antigen near the V_L - C_L interface. The F83A mutant exhibited not only the greatest increase in thermostability of all examined mutations (T_m +5 °C), but also a threefold increase in VEGF-binding affinity, predominantly due to a ~twofold improvement in the rate of dissociation (off-rate), which suggests an optimized interface. Transferring the F83A mutation to another antibody (34) of the same framework as G6.31 similarly improved thermostability (T_m +6.6 °C) and binding affinity (2.8-fold, to K_D of 1.23 nM), demonstrating that this result is not specific to G6.31 (Fig. S5C).

LC-F83 Conformations Correlate with the Fab Elbow Angle and the V_L - C_L Interface in G6 Structures. To understand how the antigen-distal mutation LC-F83A can have such a strong impact on antigen binding and stability, we first looked at the potential structural role of the LC-F83A mutation. Because G6.31 is a variant of G6 by four substitutions in CDR-L3, we used structures of G6 in the VEGF bound ($G6_{VEGF}$) and free form ($G6_{free}$) that we previously reported as a model for G6.31 (22). Notably, the $G6_{free}$ crystal structure contains 12 Fab molecules in the asymmetric unit, which can be clustered into two groups based on the Fab elbow angle (Fig. 3A), that is, the orientation of the constant domains to the variable domains (35, 36).

The particular class of LC in a given antibody is known to exert a dominant influence on distribution of Fab elbow angles (35). Lambda LC antibodies adopt a wide range of angles, whereas kappa LCs show a bimodal distribution peaking at ~140° and

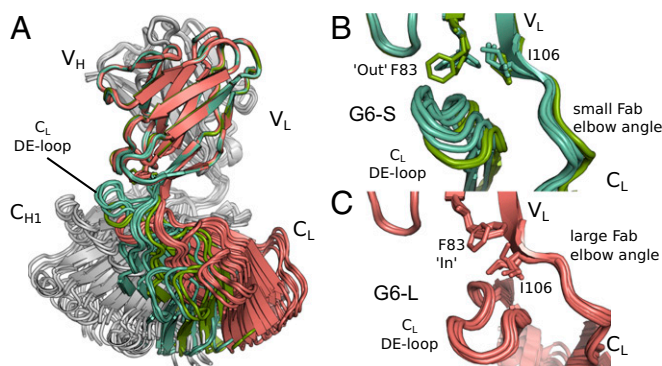


Fig. 3. Association of G6 Fab elbow angles with LC-F83 conformation. (A) Fab elbow angles of the 12 molecules present in the G6 Fab crystal structure asymmetric unit (PDB ID code 2FJF). LCs of the molecules exhibiting a small elbow angle (G6-S) are colored in green, and those with a large elbow angle (G6-L) are in red. HCs are in gray. (B and C) Detailed views of the V_L - C_L interface and the side chain conformation of LC-F83 and LC-I106 in G6-S molecules (green) or G6-L (red). The β -strand E, the helix α -1, and the loop connecting both elements are removed for clarity.

~175° (35). G6 behaves like a typical human kappa LC antibody. Six $G6_{free}$ molecules (from the asymmetric unit) have small elbow angles (143–155°) which, interestingly, are associated with a tightly packed V_L - C_L interface (comprising the DE loop of the C_L domain packed against α 1 of V_L ; $304 \pm 48 \text{ \AA}^2$), whereas the other six $G6_{free}$ molecules have a large elbow angle (170–187°) with a less tightly packed V_L - C_L interface ($234 \pm 29 \text{ \AA}^2$) (Fig. 3B and C). Moreover, the side chain conformation of LC-F83 correlates with the G6 Fab elbow angle. In the large elbow angle structures, the LC-F83 side chain is tightly held in a hydrophobic pocket (the “in” conformation) in the distal capping region of the V_L domain (Fig. 3C). In the small elbow angle structures, the LC-F83 side chain is not held in the hydrophobic pocket (the “out” conformation) and adopts rotamers that are partly solvent-exposed (Fig. 3B). Although LC-F83 is not part of the Fab elbow, its rotamers are associated with changes in the elbow residue LC-I106. In structures where LC-F83 is in the “out” conformation, the side chain of LC-I106 moves “up” to occupy the hydrophobic pocket, whereas when LC-F83 is in the “in” conformation, the side chain of LC-I106 moves “down.” In contrast to LC-F83, the movement of LC-I106 is not limited to the side chain, but also involves changes in the local protein backbone (LC-105 Φ -angle, LC-106 Ψ -angle) that determine the elbow angles as observed. In summary, the crystal structures of G6 indicate that change in the G6 Fab elbow angle is associated with the conformational states of LC-F83.

Molecular Dynamics Simulations Indicate the Effect of LC-83 Mutation on the G6 Elbow Conformation.

To gain further insight into the effects of the LC-F83A mutation found on our structural analysis, we turned to molecular dynamics (MD) simulations. The effect of the LC-F83A mutation was simulated in two different structural backgrounds, a G6 Fab crystal structure with a large elbow angle, G6-L [G6 HC V and LC U in $G6_{free}$; Protein Data Bank (PDB) ID code 2FJF], or a G6 Fab crystal structure with a small elbow angle, G6-S (G6 chains B and A; PDB ID code 2FJF). Both structures, together with the two structures modeled with an LC-F83A mutation (G6-L-83A and G6-S-83A), were simulated in water for 100 ns. All molecules except G6-L-83A showed a slight reduction of the elbow angle of the respective crystal structure in the first 25 ns of the simulation; however, after the initial 25 ns, the elbow angles of G6-L, G6-S, and G6-S-83A remained stable. For G6-L, the F83 stayed in the “in” conformation, and the elbow angle remained large (~161°), whereas G6-S maintained the “out” F83 conformation and a small elbow angle (~135°). G6-S-83A maintained the small elbow angle distribution (Fig. 4A and B and Fig. S6A). In contrast to the other three structures, G6-L-83A showed a quick transition from a large elbow angle to a smaller elbow angle (~140°) and a larger V_L - C_L interface (Fig. 4A and B and Fig. S6A), a conformation that is maintained during the remainder of the simulation.

Meta-Analysis of Human Antibody Structures Showing a Correlation Between LC-83 Conformation and V_L - C_L Packing.

We next wished to determine whether the coupling of the side chain conformation of LC-83 with the Fab elbow conformation is a common feature in human antibodies. The most frequently used human kappa germline families, IGKV1 and IGKV3 (IMGT database) (37), carry phenylalanine at position 83, whereas the other, less frequently used germlines carry mostly valine (IGKV2 and IGKV4) or alanine (IGKV5 and IGKV6). The isoleucine at position LC-106 is conserved in all five human IGKJ germline genes. We examined the conformation of LC-F83 of 319 V_{kappa} -containing human Fab crystal structures in the PDB and divided these structures into two groups: LC-F83 in an “in” conformation (dihedral χ 1 angle between -50° and -100°) and LC-F83 in an “out” conformation (dihedral χ 1 angle mostly between 50° and 180°) (Fig. 5A). Indeed, the two groups of LC-F83 rotamers correlate with changes in the Fab elbow

region at position LC-106 (LC-105 Φ -angle, LC-106 ψ -angle) in the vast majority of structures (Fig. 5B). Furthermore, the structures with a large elbow angle have a smaller, less tightly packed V_L - C_L interface, whereas structures with a small elbow angle have a large V_L - C_L interface (Fig. 5C).

Given that the LC-F83A mutation in G6.31 resulted in greater thermostability and greater affinity toward the antigen, we examined the elbow angle of nine human V_{kappa} -containing Fab struc-

tures carrying LC-A83 and LC-I106 available in the PDB. We showed that those structures mirror the properties of the LC-F83 “out” confirmation structures with a small elbow angle and a large V_L - C_L interface (Fig. 5B and C). We determined that for eight of the nine antibodies, LC-A83 is a result of somatic mutations. The remaining antibody LC is from IGKV6, which carries an alanine as a germline at position LC-83. In summary, we have demonstrated that the bimodal distribution in the Fab elbow angle observed in

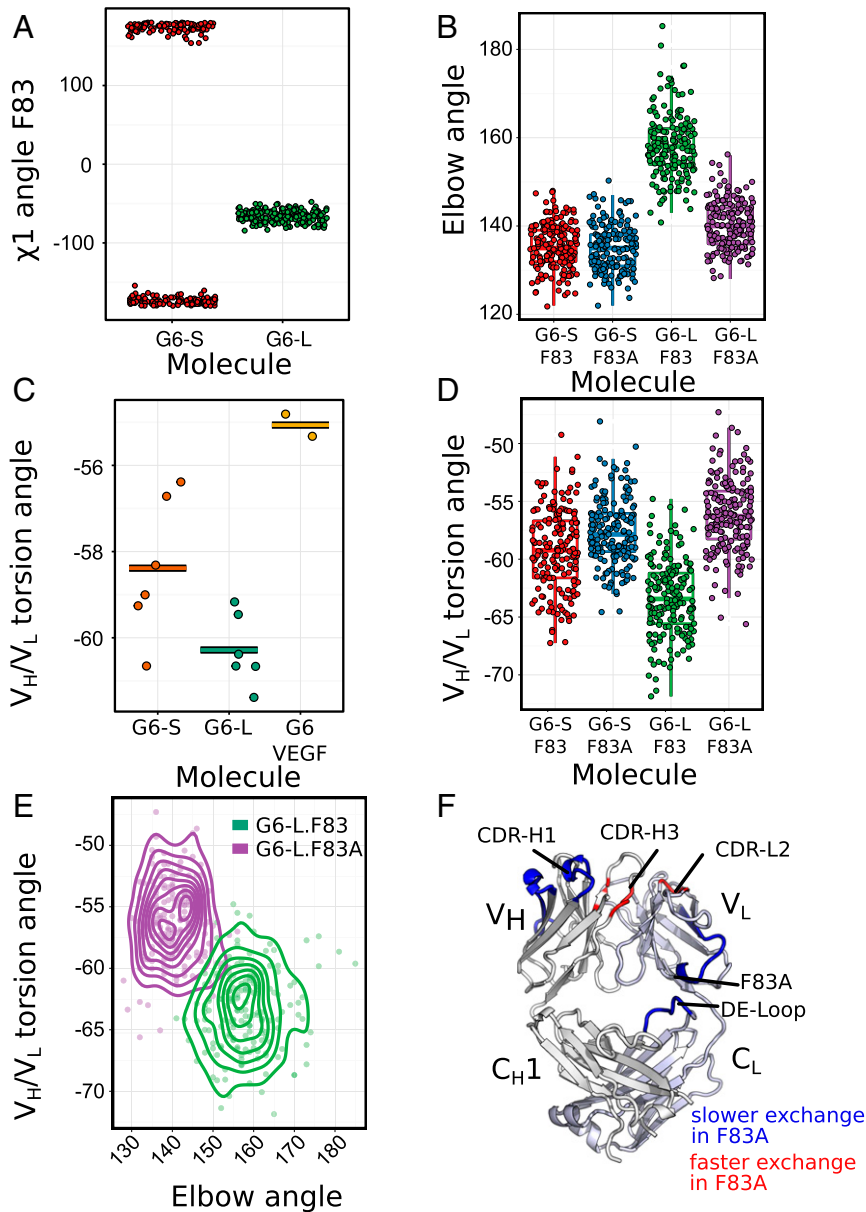


Fig. 4. Mutations at LC-F83 regulate G6 conformation and domain dynamics. (A) Distribution of the F83 side chain conformation (χ_1 angle) of G6-S (small elbow angle) or G6-L (large elbow angle) of the $G6_{\text{free}}$ (PDB ID code 2FJF) during a 100-ns MD simulation, showing little transition. The χ_1 angle of the G6-S F83 was artificially split because it was plotted in a range of -180° to $+180^\circ$. (B, D, and E) Data from the last 75 ns of the 100-ns MD simulations. (B) Distribution of the Fab elbow angle for the molecules G6-S, G6-S-83A, G6-L, and G6-L-83A. All samples are significantly different ($P < 0.001$) except G6-S and G6-S-83A, as determined by ANOVA with Tukey’s honest significance difference (HSD) test. (C) V_H - V_L torsion angle for the six molecules of $G6_{\text{free}}$ structures with F83 in an “out” conformation (orange) and for the six molecules with $G6_{\text{free}}$ with F83 in an “in” conformation (green), as well as the V_H - V_L torsion angle of the VEGF-bound G6 structures (yellow). (D) V_H - V_L torsion angle distribution of the same molecules obtained from the same MD simulation. All samples except G6-S-83A and G6-L-83A are significantly different ($P < 0.001$) by the HSD test. (E) Contour plot showing the overall Fab interdomain dynamics. The elbow angle vs. the V_H - V_L torsion angle is plotted for selected time points of the MD simulation using a G6 structure with LC-F83 in an “in” conformation (G6-L; red) and the same structure with an LC-F83A mutation (G6-L-83A; green) as starting points. Two structures occupy distinct regions in the conformational landscape. (F) In HX-MS experiments comparing G6.31 F83A and G6.31 Fab, the positions that exchange faster (red) or slower (blue) are depicted on the cartoon representation of the $G6_{\text{free}}$ Fab structure (PDB ID code 2FJF) (Fig. S6).

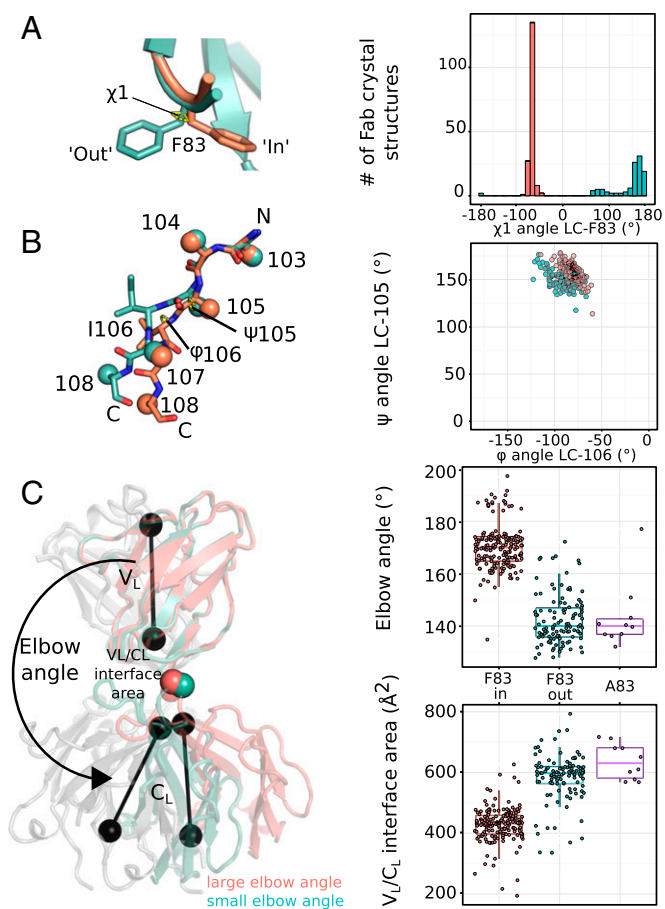


Fig. 5. LC-83 conformation correlates with the elbow angles in human antibody structures. (A) The bimodal distribution of F83 side chain χ_1 angle rotamers of 319 V_{κ} -containing human antibody structures from PDB with Phe at the position 83 in an "in" (red) or an "out" (green) conformation. (B) The 319 structures categorized according to their F83 conformation as in A exhibit a distinct distribution of elbow angle, as represented by the backbone conformation at position 105 (ψ angle) and position 106 (ϕ angle). (C, *Left*) Illustration of how elbow angles are determined for G6 Fab structures (in cartoon representations) with LC-F83 in an "in" (orange) or an "out" (green) conformation. The black dumbbells represent the pseudo-twofold symmetry axis through the variable domains or the constant domains used to calculate Fab elbow angles. The red and green spheres represent the border residue (LC-107) between the V_L and C_L domains. (C, *Right*) Elbow angle or V_L-C_L interface area of the V_{κ} containing human Fab crystal structures from the PDB with LC-83 with an "in" or "out" conformation, with nine V_{κ} Fab structures that harbor LC-A83 singled out for comparison.

human kappa LC antibody structures (35) is associated with conformations of the phenylalanine side chain at position LC-83. Furthermore, the type of amino acid present at position LC-83 indeed influences the range of the elbow angle and, consequently, the V_L-C_L interface area.

Therefore, our meta-analysis of human antibody structures, along with the MD simulations, strongly suggest that the greater thermostability of G6.31_{LC-F83A} compared with G6.31 can be explained by an increase in the V_L-C_L interface interaction area through the decrease in the Fab elbow angle, which likely further stabilizes the fold of the four domains composing the Fab.

Elbow Angle Dynamics of the G6 Fab Influence the Antigen-Binding Interface by Modulating the V_H-V_L Torsion Angle. Whereas the change in Fab elbow angle explains the increase in thermostability of the LC-F83A variant, the effect of the mutation on antigen-binding affinity cannot be explained. We hypothesize that the LC-F83A

mutation may influence the orientation of the V_H and V_L domains toward each other (i.e., the V_H-V_L torsion angle) either indirectly via the change in Fab elbow angle or directly because LC-83 sits close to the V_H-V_L interface. The V_H-V_L interface has been shown to have a strong influence on antigen-binding affinity (38, 39). Furthermore, the V_H-V_L torsion angle and flexibility of the V_H-V_L interface have been reported to vary substantially between the antigen-free and the antigen-bound forms (40, 41).

Indeed, the crystal structures of G6 show significant differences in the V_H-V_L torsion angle between the G6_{free} and the G6_{VEGF}. The mean V_H-V_L angle was -60° for the six G6_{free} molecules with a large elbow angle and -58° for the six G6_{free} molecules with a small elbow. The V_H-V_L torsion angle of G6_{VEGF} (PDB ID code 2FJG) was -55° , closer to that of the G6_{free} with small elbow angle (Fig. 4C).

To build confidence that the observed torsion angles are not an artifact of crystallization, we turned again to MD simulation. We found that the median V_H-V_L torsion angle was -62° for G6-L (with a large elbow angle) and -59° for G6-S (with a small elbow angle), consistent with that seen in G6_{free} crystal structures (Fig. 4D). Moreover, the two mutant Fabs (G6-L-83A and G6-S-83A) exhibited an even smaller V_H-V_L angle (both -57°) (Fig. 4D). The dynamic distribution of the V_H-V_L torsion angle and Fab elbow angle for G6-L and G6-L-83A (Fig. 4E) clearly indicates that the LC-F83A mutation modulates the interdomain conformations. The analysis suggests that, through its influence on the V_H-V_L torsion angle, the LC-83 mutation improves G6-VEGF binding affinity likely by optimizing the interface complementarity with VEGF or reducing the entropy cost for VEGF binding.

Hydrogen-Deuterium Exchange Mass Spectrometry Confirms the Effect of F83A Mutation in the V_L-C_L Interface and the Antigen-Binding Regions.

To assess how the LC-F83A mutation affects interdomain dynamics of the G6.31 Fab, we used hydrogen-deuterium exchange mass spectrometry (HX-MS) to measure the exchange rate of backbone amide hydrogen atoms in D₂O by following the increase in mass as backbone protons exchange with deuterium over time. The exchange rate of each backbone amide proton reflects the aggregate structural energetics of all H-bonding interactions with that specific exchange site (42). Increased structural stability or reduced conformational fluctuation generally causes a relative reduction in the observed exchange rate of nearby residues, and the converse is true as well.

Comparing exchange rates between G6.31 and G6.31_{F83A} Fab revealed both short-range and long-range consequences of the mutation on structural energetics (Fig. 4F and Fig. S6B). The significant reduction in the exchange rate near position 83 in G6.31_{F83A} is consistent with the prediction that V_L-C_L interface packing increases on mutation. Long-range allosteric effects of the mutation are manifested near the antigen-binding site. The CDR loop H1 exchanges more slowly in G6.31_{F83A}, whereas the sections of the CDR-H3 and CDR-L2 known to be a part of the V_H-V_L interface (7, 43, 44) exchange more rapidly.

The results obtained by the MD simulation and HX-MS are in good agreement, and thus both methods independently confirm the impact of this antigen-distal mutation at LC-83 on Fab domain orientation and dynamics.

LC-83 and LC-106 Are Frequently Mutated in Human Antibodies. We next investigated whether mutation at the position LC-83 occurs frequently during affinity maturation in vivo among human antibodies of the same germline family. We used two approaches to examine the distribution of SHMs in LC sequences originating from IGKV1 (V_{κ} 1) germlines. In the first approach, we collated the SHMs of 4,623 unique human IGKV1 LC sequences available in several publicly available databases (Fig. 6A, *Upper*). In the second approach, we amplified IGKV1 sequences from

We note that the F83S mutation reportedly increased the affinity of a $V_{\text{kappa}}1$ -containing anti-estradiol antibody (45).

In summary, LC-83 and LC-106 are frequently mutated in human antibodies. Given the influence of these two positions on affinity and stability, optimization of the overall Fab domain dynamics may be a widely used molecular mechanism during antibody affinity maturation in vivo.

Conclusions

In this work, we mapped the entire mutational landscape of the V_{H} and V_{L} domains of a human antibody by single-site saturated mutagenesis, showing the impact of mutation at each position on antibody stability and antigen binding. Using a similar approach, Traxlymeyer et al. (46) recently reported that residues of the HC constant domains CH2 and CH3 are generally conserved and resist mutation for proper folding, in stark contrast to the high mutational tolerance of the numerous variable domain positions of G6.31 (Figs 1 and 2 and Figs. S1 and S3). This suggests that antibody constant domains are highly evolved and optimized for folding and stability, whereas variable domains can be mutated with excellent potential to further improve stability and optimize binding function. The comprehensive mutational landscape of the variable domain suggests a road map for the somatic mutation process to enhance antibody function and also guide the engineering and optimization of antibodies needed for therapeutic development.

To search for novel mechanisms of somatic mutation that affect antigen-binding function, we focused on beneficial mutations at framework positions distal to the antigen. We report how mutation at residue 83 of G6.31 LC can modulate interdomain structural dynamics of Fab molecules through interaction with elbow residue LC-I106. The interplay of these two residues is associated not only with the dynamics of Fab elbow angle, but also with the interaction between V_{L} and C_{L} and between V_{L} and V_{H} (Figs. 3–5). We further demonstrate the high frequency of mutations at LC-F83 in human antibodies of the IGKV1 germline source, often to smaller residues (Fig. 6), and show that these mutations are associated with similar changes in the interdomain conformations as for G6.31.

Our analysis of anti-VEGF antibody G6.31 indicates several specific roles for the residue in LC position 83 during in vivo affinity maturation. First, LC-F83 mutations have an impact on antigen binding by influencing the orientation of V_{H} and V_{L} (Fig. 6), and thus the topography of the CDRs. Second, F83 mutation to the smaller side chain (A, V, or S) induces a more compact Fab conformation and increases structural stability. Third, LC-F83 mutation can affect the biological activity of an antibody–antigen interaction through its modification of the Fab elbow angle and interdomain dynamics (47–49). Furthermore, B-cell signaling is sensitive to small conformational changes in the B-cell receptor (50). Modulation of the Fab elbow angle dynamics may have a favorable effect on B-cell receptor signaling (51).

Our study provides insight into the active role of framework positions in modulating antibody affinity and stability. The CDRs typically provide the main direct contacts with antigens, but—as highlighted by our analysis—affinity also depends on framework residues, even when the FWR positions are structurally far away from the antigen. Our investigation into the role of mutations at LC position 83 has unraveled the complex relationship between a framework mutation and the conformational dynamics of a Fab molecule, emphasizing the importance of conformational dynamics in antibody–antigen interaction.

Materials and Methods

Full Variable Domain NNK-Walk Library Design, Generation, and Phage Panning. V_{H} residues 2–113 and V_{L} residues 2–107 of G6.31 Fab in a phagemid construct, as described previously (26) (Fig. S1), were subjected to randomization by oligonucleotide-directed mutagenesis (52). Between 10 and 12 sequential

residues were mutagenized in a single reaction using a mixture of oligonucleotides, each with an NNK codon at one position targeted for randomization. NNK encodes for all 20 amino acids ($n = G, A, T, \text{ and } C; K = G \text{ and } T$, in equal proportions). The template for each mutagenesis reaction was generated by introducing stop codons (TAA) at all of the positions targeted for randomization in each reaction. The distribution of NNK vs. wild-type (WT) codons at a given position depended on the size of the randomized segment; for example, for the V_{L} domain, 106 positions (amino acid position 2–107) were randomized. Thus, for a given position in the library, 99.1% of reads carried the WT codon, whereas only 0.9% of the reads carried mutations resulting from the NNK randomization. The V_{H} and V_{L} libraries were generated separately by electroporating the pool of DNA products from mutagenesis reactions into *Escherichia coli* XL1 cells, yielding $\sim 10^9$ transformants as described previously (24).

Phage Panning Selection. To select for stably expressed clones, the V_{H} and V_{L} libraries were incubated with protein A, protein L, or anti-gD immobilized on ELISA wells for 2 h, followed by washing away of the unbound phage. Bound phage was eluted in 100 mM HCl for 20 min, then neutralized with 1/10 volume of 1 M Tris pH 11.0 and used to infect *E. coli* for amplification, followed by one round or two rounds of plate-based selection. VEGF-binding clones were selected by incubating the phage display libraries with 5, 0.5, and 0.1 nM biotinylated VEGF in solution in the successive rounds of selection with increasing stringency, as described previously (53). Bound clones were captured on ELISA wells coated with neutravidin, washed, eluted, and amplified as above.

Illumina Sequencing of G6.31 Deep Mutational Scanning Libraries. For deep sequencing, phagemid DNA was isolated from *E. coli* XL1 cells from either the unselected or the selected G6.31 NNK library and used for a limited-cycle PCR-based amplification of V_{L} and V_{H} regions using Phusion polymerase (New England Biolabs). PCR products were purified to generate libraries using the TruSeq Nano DNA library preparation kit (Illumina). Multiplexed adapter-ligated libraries with unique barcodes were sequenced on an Illumina MiSeq sequencing system for 300-bp paired-end sequencing.

Deep Mutational Scanning Data Analysis. After merging of paired-end reads using FLASH (54), data analysis was performed using the statistical programming language R (55) and the ShortRead package (56) (Table S1). The first step in quality control was to filter out sequences that did not carry the respective V_{H} and V_{L} barcodes. In a second step, the flanking regions of the V_{H} and V_{L} domains were identified for each merged read and checked for the correct lengths to remove the reads with indels (insertion and deletion). To further correct for sequencing errors, non-NNK mutations were converted back to the WT base, and reads that contained two or more NNK mutations were filtered out. Calculating the frequency of all mutations at each randomized position generated position weight matrices. ERs for all mutations were calculated by dividing the frequency of a given mutation at a given position in the sorted sample by the frequency of the same mutation in the unsorted sample, as described previously (25, 26). Mutagenesis of one segment of V_{H} (position 14–25) in the original V_{H} library resulted in errors which led to low expression of the respective segment. This result was likely related to a mutational error in the original stop template used in the mutagenesis reaction for this V_{H} segment 14–25, which affected expression of the variants. This error was discovered only after sequencing and data analysis. Therefore, a second V_{H} library was generated with correction in the 14–25 segment (by making a new stop template vector), and phage selection was repeated. Outside of the 14–25 regions, data from the two V_{H} libraries were generally correlated with each other, and data from the second library were used in all of the selections for folding and expression (anti-gD antibody, protein A, and protein L). The VEGF selection data of the corrected segment from the second library were merged with the sequence data of the original library after calculation of ERs. To adjust for different sequencing depths, the dataset of the second V_{H} library was adjusted by obtaining a random sample of sequences of sizes matched to the dataset obtained from the first V_{H} library (Table S1).

Antibody Expression and Characterization. V_{H} and V_{L} sequences of selected variants were cloned into a mammalian Fab vector for expression and purification with a protein G column. Surface plasmon resonance measurements with a BIACore T200 instrument was used for affinity determination as described previously (24, 26). In brief, single cycle injection was used for kinetic measurement. VEGF (8–109) was coupled to Series S sensor chip CM5 (GE Healthcare) to achieve an R_{max} value of ~ 50 response units. Serial dilutions of Fab in HBS-P buffer (0.01 M HEPES pH 7.4, 0.15 M NaCl, and 0.005%

Tween 20) (0.08–50 nM) were injected. The VEGF-binding responses were corrected by subtracting responses from a blank flow cell. Association rates (k_{on}), dissociation rates (k_{off}), and K_d (k_{off}/k_{on}) were calculated using a simple one-to-one Langmuir binding model. For T_m , we used DSF, which monitors thermal unfolding of proteins in the presence of fluorescent SYPRO orange dye (Invitrogen). The diluted dye (1:20; 1 μ L) was added into 24 μ L of Fab protein (~100 μ g/mL). The fluorescence intensity during the temperature increase from 20 °C to 100 °C was plotted, and T_m , the inflection point of the transition curve, was calculated using the Boltzmann equation (57).

Structural Meta-Analysis. Human antibody structures were obtained from the structural antibody database SABDab (58). Additional structure filtering and manipulation were performed using Bio3D (59). The interface area between constant and variable domains was calculated using a local instance of CCP4-Pisa (60). ABangle (40) was used to characterize the V_H - V_L interaction. Pymol and a publicly available script (www.pymolwiki.org/index.php/Elbow_angle) were used to calculate the elbow angle.

MD Simulations. Before the MD simulations were performed, segments in the constant domains of G6 (HC130-145, HC220-223, and LC211-213) that had not been resolved in the G6 crystal structure were merged from a high-resolution crystal structure (4hh9.DC) into the G6 structure. Mutations at the LC-83 position were introduced using Pymol. The AMBER12 simulation package (61) with the ff99SB force field was used in all minimization, equilibration, and production stages. All structures were solvated in a truncated octahedron box using the TIP3P water mode with a 10-Å buffer of solvent between the solute's furthest dimensions in each direction to avoid interaction with self-images. Each system was neutralized by adding Cl counterions using a Coulomb potential on a 1-Å grid. (All initial systems were of net positive charge.) The system (solute and solvent) was minimized in six stages with a harmonic restraint on the solute slowly decreasing from 40 to 0 kcal/mol/Å², with each stage over 100 steps of steepest descent. Following the energy minimization, the system temperature was increased from 150 K to 300 K over 50 psec at 2-fsec integration steps. Bonds involving hydrogens were constrained by the SHAKE algorithm. Production data of 100 ns were collected at the NPT ensemble at 3-fsec time steps and recorded every 15 psec. The particle mesh Ewald method was used to calculate the full electrostatic energy of a periodic box. The obtained simulated structures were analyzed using VMD (62), Bio3d, CCP4-Pisa, Pymol, and ABangle.

HX-MS. G6.31 WT and F83A mutant Fab (45 μ M) were diluted 15-fold into a 20 mM histidine-acetate buffer at pH 7.0 and 50 mM NaCl with >90% D₂O content to begin the labeling reaction at 20 °C. At six logarithmically spaced time intervals spanning 30 s to 1,000 min, the labeling reaction was quenched by pH reduction (pH 2.5) and the addition of 2 M guanidinium chloride and 0.25 M Tris(2-carboxyethyl)phosphine before injection onto a cold online system (63). In brief, quenched samples were first passed through an immobilized pepsin column (2.1 \times 30 mm; Applied Biosystems) at 0 °C for proteolysis and then bound to a trap column (Acquity Vanguard C8) for desalting before being separated by reverse-phase chromatography

(Acquity UPLC BEH C18; 1.7 μ m particle size; 1.0 \times 50 mm) and introduced into the mass spectrometer (Thermo Orbitrap Elite; 120 kHz resolution at m/z 400) for measurement of deuterium content. Chromatographic mobile phases were prepared as described previously (64) with a pH of 2.25 to maximize deuterium recovery, which averaged 82% by measurement of fully deuterated controls. Mutant and WT experiments were interleaved, and randomized time points were collected in triplicate.

This process resulted in 152 unique peptides, consistently identified by the ExMS program (65) for both WT and mutant samples, providing sequence coverage of roughly 95%. The analysis of deuterium content involved the use of previously described custom Python scripts (64, 66, 67). Significant differences in deuterium uptake levels between WT and mutant were identified by Student's *t* test as having a *P* value < 0.05, as described previously (68).

Somatic Mutations in Human Antibodies. This analysis was performed for antibody sequences obtained from publicly available sources. Duplicated sequences were removed, and the closest germline genes were assigned for the respective V-segments using IGBlast (69). Sequences to which a mouse or rat germline was assigned were removed from the dataset. Sequences were Kabat-numbered (70) and somatic mutations in the framework of the V-segment were identified. Mutations in the CDR regions (as defined by the Kabat numbering scheme) were not considered in our analysis.

In a second experiment, the RNA from peripheral blood lymphocytes, tonsil, spleen, and bone marrow of more than 1,000 individuals was obtained from commercial source (Clontech, Amsbio, and Biochain) either as total RNA or poly(A)-positive RNA. The RNA was first transcribed into cDNA using the SMARTer RACE cDNA Amplification Kit (Clontech). IGKV-specific DNA was amplified using 5' RACE (Advantage 2 PCR Kit; Clontech) with a primer annealing the 5' region of the IGKC region (5' CATCAGATGGCGGAA-GATGAAGACAGATGGTGC 3'), whereas IGKV1 segments were enriched in a second PCR step using the following primers: forward, 5'-GCCATCCA-GATGACCCAGTCTCC-3'; reverse, 5'-GGCTGCACCATCTGTCTTC-3'. The PCR product was circularly (and hence repeatedly) sequenced using Pacific Bioscience RSII (EA Genomic Services). A total of 994.8 Mbp were obtained. A consensus of each circular sequencing read was created to improve the fidelity of sequencing using Pacific Bioscience's SMRT Analysis 2.3 software package. The germline annotation of the consensus sequences was performed using VDJFasta (71). Sequences were Kabat-numbered, and somatic mutations in the framework of the V-segment were identified. A total of 19,034 sequences were annotated as IGKV1; among these, 3,796 contained at least three FWRs and two CDRs and were included to generate the SHM distributions.

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