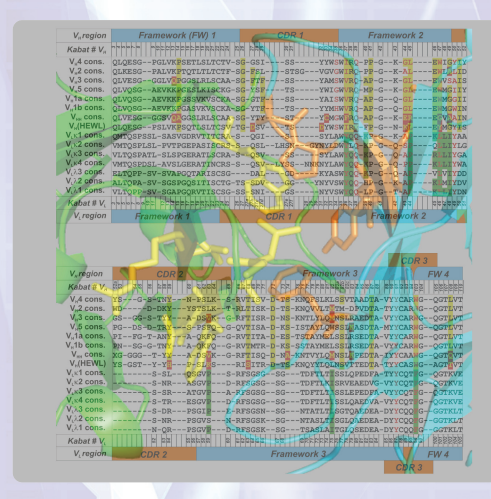


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Residue Covariations in Antibody Domains



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Conserved amino acid networks involved in antibody variable domain interactions

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ABSTRACT

Engineered antibodies are a large and growing class of protein therapeutics comprising both marketed products and many molecules in clinical trials in various disease indications. We investigated naturally conserved networks of amino acids that support antibody $V_{\rm H}$ and $V_{\rm L}$ function, with the goal of generating information to assist in the engineering of robust antibody or antibody-like therapeutics. We generated a large and diverse sequence alignment of V-class Ig-folds, of which V_H and V_L domains are family members. To identify conserved amino acid networks, covariations between residues at all possible position pairs were quantified as correlation coefficients (φ-values). We provide rosters of the key conserved amino acid pairs in antibody $V_{\rm H}$ and $V_{\rm I}$ domains, for reference and use by the antibody research community. The majority of the most strongly conserved amino acid pairs in V_H and $V_{\rm L}$ are at or adjacent to the $V_{\rm H}$ - $V_{\rm L}$ interface suggesting that the ability to heterodimerize is a constraining feature of antibody evolution. For the V_H domain, but not the V_L domain, residue pairs at the variable-constant domain interface $(V_{\rm H}-C_{\rm H}1$ interface) are also strongly conserved. The same network of conserved V_H positions involved in interactions with both the V_L and $C_{\rm H}1$ domains is found in camelid $V_{\rm HH}$ domains, which have evolved to lack interactions with $V_{\rm L}$ and C_H1 domains in their mature structures; however, the amino acids at these positions are different, reflecting their different function. Overall, the data describe naturally occurring amino acid networks in antibody Fy regions that can be referenced when designing antibodies or antibody-like fragments with the goal of improving their biophysical properties.

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Key words: immunoglobulin variable domain; Ig-fold; V-class; covariation; antibody engineering.

INTRODUCTION

Antibodies are useful targeted therapeutics because of their ability to bind specific ligands with high affinity and specificity. Antibody variable domains ($V_{\rm H}$ in the heavy chain, $V_{\rm L}$ in the light chain), which provide the binding capability, may be purposely engineered to impart desired antigen recognition or binding affinity properties. Some designs have implemented recombinant production of isolated $V_{\rm H}$ - $V_{\rm L}$ domains (Fv region), providing researchers with more design flexibility than standard antibody therapeutics (e.g., the expression of the Fv region as a single polypeptide chain or "scFv" 1-5). However, removal of the V_H - V_L domains from the quarternary structure of an antibody can lead to stability and solubility problems. Several mechanisms have been proposed to account for the generally poor biophysical behavior of scFvs and related designs, and include the intrinsic instability of the isolated domains, the weak affinity between $V_{\rm H}$ and $V_{\rm L}$ domains, and the absence of possibly stabilizing interactions with the antibody constant domains. 6 An understanding of the specific amino acids that mediate interactions between the $V_{
m H}$ and $V_{
m L}$ domains, and between the variable and constant domains, would enable improved designs of antibodies and antibody-like proteins.

Antibody variable domains are part of the immunoglobulin domain or "Ig-fold" superfamily. The Ig-fold superfamily is a large group of structurally related protein domains commonly found in mammalian cell surface proteins or in soluble extracellular signaling proteins. Ig-fold domains consist of two β -sheets, each arranged in a "Greek-key" topology, that are packed tightly against one another and are generally supported by an intradomain disulfide bond. Depending on the number of strands in each β -sheet and the loop connections between the strands, the superfamily can be divided into several subfamilies including the C-, I-, and V-classes. Antibody variable domains are V-class Ig-folds and their constant domains are C-class Ig-folds (Fig. 1). Ig-fold or Ig-fold-like domains are also present in cell adhesion proteins, integrins, allergens, T-cell

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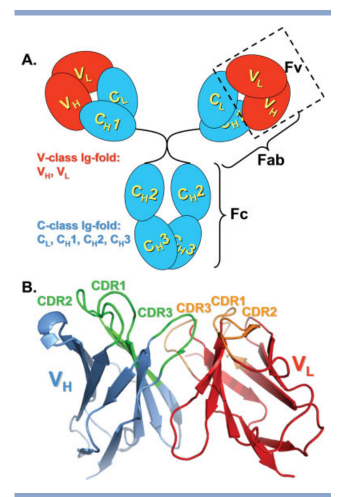


Figure 1

Diagrams of an immunoglobulin and its Fv domain. A: Schematic diagram of an IgG antibody. The variable domains that compose the antigen-binding or Fv-region are shown in red and the constant domains are shown in blue. The variable domains are V-class Ig-folds, whereas the constant domains are C-class Ig-folds, which are highly similar to V-class Ig-folds, but lack two additional β-strands commonly found in V-class structures. B: Ribbon diagram of an antibody Fv-region consisting of a variable domain from the immunoglobulin heavy chain (V_H-blue) and a variable domain from the immunoglobulin light chain (V_L -red). The complementarity determining regions (CDRs) of the V_H (shown in green) and the $V_{\rm L}$ (shown in orange) comprise the antigen-binding site.

receptors, major histocompatibility complexes, immunoglobulin receptors, and many other protein families with diverse functions.

The past decade has seen a significant increase in the number of publicly available Ig-fold sequences. Large databases of antibody variable domain and T-cell hypervariable domain Ig-fold sequences have been compiled.9 Information from these databases has been instrumental in antibody humanization, affinity maturation, and the stabilization of single chain Fv (scFv) or other antibody constructs.^{5,10-13} Antibody sequence databases generally influence antibody design by enabling frequency analyses at single amino acid positions (i.e., consensus modeling) that may be used for generating rational designs. 12,14–16 Recent studies with other protein domain superfamilies have extended sequence-based approaches by examining how amino acid pairs or networks may be conserved within subsets of a protein superfamily with related function or across diverse members of a protein superfamily. Such amino acid networks may define important structural or functional features of these protein domains. 17-19 These approaches, sometimes referred to as "covariation analyses," track whether the presence (or absence) of a particular amino acid at one position correlates with the presence (or absence) of another amino acid at a second position within a multiple sequence alignment. Although covariation analyses have been performed on several protein families (including SH3 domains, WW-domains, TPR-motifs, GPCRs, serine proteases, globins, viral coat proteins, and others $^{20-23}$), very little has been described concerning covariation analyses of Ig-folds. The paucity of Ig-fold covariation data may stem from several factors, one being that large collections of Ig-fold sequences were, until recently, limited primarily to antibody sequences, particularly human and murine.^{24,25} Also, accurate alignment of diverse members of large proteins (>100 amino acids) like Ig-fold domains is challenging and misalignments can limit the validity of covariation data.²³

Here we describe the application of covariation analyses to a high-quality, 3D-structure-based alignment of diverse V-class Ig-fold sequences. A diverse V-class Ig-fold sequence alignment was constructed, and covariations were quantified as correlation coefficients (ϕ -values²³) for every amino acid pair (i.e., every residue combination found at all possible pairs of positions) in the alignment. The results serve as a rich repository of amino acid interactions conserved throughout Ig-fold evolution. The data reveal conserved residue networks that may support interactions between the $V_{\rm H}$ and $V_{\rm L}$ domains. The data also reveal that V_H domain networks involved in interactions with $V_{\rm L}$ domains are co-conserved with residue networks observed at the V_H-C_H1 junction suggesting that these two functional areas have coevolved to support the overall quarternary antibody structure.

METHODS

Creation of structure-based Ig-fold alignments

Structures of Ig-fold proteins or Ig-fold domains from multidomain proteins were gathered from the ASTRAL database, 26,27 which contains domain structures matching the Structural Classification of Proteins (SCOP, Version 1.69) hierarchy. SCOP classifies the Ig-fold as a member of the "Beta Proteins, Immunoglobulin-like βsandwich fold, Immunoglobulins" superfamily. 28 PDB

files of the V-class Ig-folds were downloaded using customized shell scripts. Each Ig-fold structure was inspected visually using Swissprot DeepView; sequences were removed from the study if they were erroneously categorized, incomplete (either missing residues due to a lack of electron density or domain swapped²⁹), redundant (i.e., those with identical sequences), or obviously did not conform to the β-sandwich Ig-fold topology. Sequences of aberrant length (>2-times the standard deviation about the mean V-class length, 112.0 ± 10.6 residues) were also removed. 702 structures were aligned using the Secondary Structure Matching (SSM) assisted implementation in the Schrödinger Prime structalign program. 30-32 Schrödinger Prime was used to generate structure-based V-class sequence alignments based on the proximity of each C_{α} atom subsequent to an all-to-all structure alignment, which minimized the average distance between all structural pairs. The alignment was most accurate in the regular β-strand regions and less accurate in the connecting loop regions because of variable loop lengths and structures.

Generation of a diverse V-class Ig-fold sequence alignment

A custom Hidden Markov Model (HMM) of the Vclass Ig-folds was built from the structure-based sequence alignments. The HMM was created with the HMMER software package (version 1.8), using the "hmmbuild" and "hmmcalibrate" functions.33 The HMM was used to find potential V-class Ig-fold sequences in the NR-database maintained at NCBI using the "hmmsearch" function. The output of this function ranked the hit sequences by their scores relative to our custom V-class HMM, and sequences with scores above a recommended threshold were retained as candidate members of the V-class dataset. The output also provided the number of "hits" per sequence (i.e., the number of Ig-folds within a contiguous gene sequence) and the exact residue positions of the hits. For sequences containing one or more candidate V-class sequences, the relevant subsequences were extracted from the full NR sequence using a custom Java executable. As an additional test to confirm that each sequence pulled from NR using our V-class HMMs belonged to the V-class Ig-fold subfamily, the custom shell script "pfamverify" using the HMM tool "hmmpfam" was applied to each Ig-fold candidate sequence.³⁴ Ig-clan HMMs (including V-, I-, C1-, C2-, and less specific Ig HMMs) were downloaded from the PFAM website. Sequences that scored lower with the PFAM V-class HMM than with other PFAM Ig-fold HMMs, and sequences whose score with the PFAM Vclass HMM lay below recommended cutoffs (TC1 defined at the PFAM website) were removed. Thus, Vclass Ig-fold sequences were retained only if their PFAM scores validated their Ig-fold class assignments. The Ig-

fold sequences extracted from NR were aligned by our structure-based V-class HMM using the "hmmalign" function in the HMMER package. The resulting V-class dataset contained 48,696 sequences including those from both the SCOP 3D protein database and NR.

The resulting sequence collection was biased toward well-studied Ig-fold-containing proteins (i.e., human and murine V-class sequences frequently deposited in NR). To reduce the over-representation of these sequences, we developed a heuristic algorithm that eliminated sequences based on identity cut-off criteria. In brief, percent identities were calculated for all sequence pairs. Sequences were grouped into bins representing their maximum percent identity with any other sequence (i.e., 99% bin, 98% bin, 97% bin, etc.). Sequences within each bin were then ranked according to decreasing nongap residue count, giving better ranks to sequences with fewer gaps. In each bin, sequences with an equal number of nongap residues were ranked by Henikoff weights to filter out more common sequence types while preserving rare sequences with the goal of increasing diversity within the final datasets.³⁵ An identity cutoff of 80% was used for V-class sequences. This left 2786 sequences, each with less than 80% identity to all other sequences, in the *V*-class dataset.

The resulting multiple sequence alignment contained many positions populated by gaps (>50% gaps for most sequences). To eliminate this problem, columns that were not match states in the HMM were removed. This resulted in 144 remaining columns for our custom Vclass alignment. Still, 354 sequences contained >40% gaps. These sequences, which were generally incomplete, were removed from the alignment. The final V-class Igfold dataset contained 2432 sequences. Virtually all the V-class sequences in the dataset were naturally occurring (nonengineered).

Correlation coefficient (ϕ -value) calculation

Covariation between amino acid pairs in multiple sequence alignments were calculated as correlation coefficients (φ-values), as described previously.²³ The calculations were encoded into a Java executable and run with Java Runtime Engine (JRE) version 1.4.2. φ-values were defined as

$$\frac{(x_{i}y_{j} \times \overline{x}_{i}\overline{y}_{j}) - (x_{i}\overline{y}_{j} \times \overline{x}_{i}y_{j})}{\sqrt{(x_{i}y_{j} + \overline{x}_{i}y_{j}) \times (x_{i}\overline{y}_{j} + \overline{x}_{i}\overline{y}_{j}) \times (x_{i}y_{j} + x_{i}\overline{y}_{j}) \times (\overline{x}_{i}y_{j} + \overline{x}_{i}\overline{y}_{j})}},$$
(1)

where $x_i y_j$ is the number of times amino acids of type "x" or "y" are found in the same sequence at positions iand j, respectively, $\overline{x}_i \overline{y}_j$ is the number of times both amino acids are absent from the same sequence, $x_i \overline{y}_i$ is the number of times x is found present while y is absent,

and $\overline{x}_i y_i$ is the number of times x is absent while y is present. This equation can be rewritten as:

$$\phi(x_i y_j) = \frac{(a \times d) - (b \times c)}{\sqrt{efgh}},$$
(2)

where a to h are given by the contingency table:

	X_i	\overline{X}_i	Total
y_j	а	b	е
\overline{y}_i	С	d	f
Total	g	h	

and $a = x_i y_j$, $b = \overline{x}_i y_j$, $c = x_i \overline{y}_j$, $d = \overline{x}_i \overline{y}_j$, e = a + b, f = c + d, g = a + c, and h = b + d. Particular residue pairs (specific combinations of residues at specific positions) were not considered unless they were observed in the alignments a minimum of 10 times.

Statistics

Statistical significance of the φ-values was evaluated with a chi-square (χ^2) test, using Bonferroni-corrected Pvalues to adjust for multiple testing.³⁶

The χ^2 test is often used to evaluate the significance of values observed in contingency tables of two dichotomous variables, such as the contingency table above. The equation for this use of χ^2 can be written as

$$\chi^2 = \sum \left[\frac{\left(o_k - e_k \right)^2}{e_k} \right],\tag{3}$$

where o_k stands for the observed frequency and e_k stands for the expected frequency in one cell of the table. χ^2 is calculated by taking the sum of the squared and normalized differences between the observed and expected frequencies over all the cells. When expected frequencies are unknown, they can be estimated from observed frequencies and the equation becomes

$$\chi^2 = \frac{N \times (ad - bc)^2}{efgh},\tag{4}$$

with a to h representing the same values as in Eq. (2)above, and N standing for the total number of samples. Comparing Eqs. (2) and (4), it is evident that

$$\chi^2(x_iy_j) = \phi(x_iy_j)^2 \times N$$

This relationship between χ^2 and φ is useful because of the rich information available about the χ^2 statistic, including tables of P-values for χ^2 with specified degrees of freedom (df). However, before proceeding to use this relationship, we performed random simulations to confirm that it held for our dataset. We took our V-class sequence alignment, performed repeated (tens of thousands) random shuffling of the residues at each position (so that the residue frequencies at each position remained unchanged, but the correlations across positions were randomized), and computed ϕ -values for each randomization. We then calculated the probabilities of observing specific strong covariations by chance directly from these random simulations. In all cases examined, we found close agreement between the probabilities observed in the simulations and those calculated from χ^2 .

Having validated the use of χ^2 to determine significance in our dataset, we converted our ϕ -values to χ^2 using Eq. (4), and used standard χ^2 tables (df = 1) to find P-values. φ-values were calculated for the 186,171 amino acid pairwise combinations that occurred at least 10 times in our V-class alignments. To correct for multiple testing, we used a Bonferroni-corrected P-value³⁶ as our criterion for significance, striving for significance at the true P < 0.0001 level. For positive correlations, this corresponded to φ-values >0.1237. Use of the Bonferroni correction along with this strict P criterion gave a very conservative list of amino acid pairs with significant ϕ values, and greatly reduced our chances of finding false positives. Even with this conservative approach, 13,796 significant positive correlations (see Results) were observed.

RESULTS

Alignment quality and diversity

Information about protein 3D structures can significantly improve the quality of multiple sequence alignments.³⁷ As described in the Methods, we compiled 3D structures of V-class Ig-folds from SCOP and generated a structure-based multiple sequence alignment of these Vclass domain sequences. A custom HMM was built from this structure-based alignment and used to align additional Ig-fold sequences from the NR sequence database. Here we discuss the quality and the diversity of the resulting alignment.

The quality of the alignment guided by our structurebased HMM was evaluated by examining whether the HMM properly aligned sequences that, though disparate in residue identity, are known to form the same part of an Ig-fold 3D structure. As expected, residues that make up the β -strands of both V_H and V_L domains were well aligned, whereas alignments of residues in the loop regions, whose structures are more variable, often contained many gaps. We also looked to see whether the HMM properly aligned the consensus sequences of antibody $V_{\rm H}$ and $V_{\rm L}$ chains, in the $V_{\rm H}$ - $V_{\rm L}$ interface region. The heterodimeric structure of the interface is highly symmetrical (both $V_{\rm H}$ and $V_{\rm L}$ domains use the same face of their Ig-fold to create the heterodimer). Thus, residues buried in the interface should align in 3D, despite differences between the amino acids of $V_{\rm H}$ and $V_{\rm L}$ Ig-folds at these positions. We used a 1.8 Å crystal structure of an antibody Fab from our lab (Jordan et al., manuscript in preparation) to determine the residues of both $V_{\rm H}$ and $V_{\rm L}$ that bury surface area at the interface using the program MOLMOL. ³⁸ The $V_{\rm H}$ and $V_{\rm L}$ residue positions buried at the interface mapped to the same residue positions within the multiple sequence alignment, even though the amino acid identities at these positions vary.

Covariation analyses are most successful when applied to sequence datasets that are highly diverse. 23,39-41 From a practical standpoint, φ-value correlation coefficients increase when there are many instances of both the presence and absence of a conserved amino acid pair across a sequence alignment (see numerators of Eqs. 1 and 2). Highly diverse sequence sets are more likely to contain sequences both with and without the pair than are datasets of highly related sequences. As our goal was to investigate conserved amino acid networks in V_H and $V_{\rm L}$ domains, it was therefore important that our V-class Ig-fold dataset contains a background of Ig-fold family members that are not evolutionarily constrained to perform the same function as $V_{\rm H}$ and $V_{\rm L}$ domains. Additionally, covariation signals pertaining to polar interactions have been shown to be stronger in datasets with moderate to high evolutionary distances between sequences.⁴⁰ Protein interfaces, in which Ig-folds frequently appear, often rely more heavily on polar interactions than do protein cores, ⁴² providing another reason for generating a V-class Ig-fold sequence dataset that was highly diverse. The unfiltered sequence set from NCBI contained ~50,000 sequences highly biased toward immunoglobulin variable domains. An 80% identity filter was used to reduce bias toward over-represented V-class families, thus promoting diversity. After filtering, the dataset contained 2432 V-class sequences. Members of the V-class dataset could be divided into three functional categories: (1) 50% were immunoglobulin variable genes (including both $V_{\rm H}$ and $V_{\rm L}$ antibody domains); (2) 16% were T-Cell Receptor V-class genes; and (3) 34% were Vclass genes derived from diverse functional families, each of which comprised less than 5% of the V-class sequences. The sequences were derived from species ranging from cartilaginous fish to primates. There was a bias toward human immunoglobulin variable domain sequences with 574 of the total 2432 V-class sequences being human $V_{\rm H}$ (484 of 993 $V_{\rm H}$) or human $V_{\rm L}$ (90 of 187 $V_{\rm L}$). Examples of other species contributing $V_{\rm H}$ and $V_{\rm L}$ sequences to the V-class database include mouse (44), cow (16), camel (174), llama (83), macaque (17), and chicken (9). Despite the bias toward human $V_{\rm H}$ and $V_{\rm L}$, the average sequence identities within $V_{\rm H}$ and $V_{\rm L}$ subgroups were low – 41 \pm 1% and 29 \pm 1%, respectively. The distribution of germline $V_{\rm H}$ and $V_{\rm L}$ sequences passing the 80% identity filter roughly matched the naturally observed distribution of variable domain sequences⁴³ suggesting that variable gene subclasses were similarly diverse and fairly represented in the sequence dataset. Positional entropy calculations using the final V-class dataset demonstrate the much higher positional diversity with the V-class dataset compared with antibody $V_{\rm H}$ or $V_{\rm L}$ datasets that may be used for consensus analyses 15,44 (Supp. Info. Fig. 1).

Correlation coefficients (6-values) between residues of V-class Ig-folds

We adopted a previously described method, the use of φ-value correlation coefficients, for quantifying covariations of residue pairs within sequence alignments.²³ Figure 2 shows the number and distribution of ϕ -values calculated for amino acid pairs that were observed ≥ 10 times within the V-class Ig-fold alignment. Positive φvalues represent positive correlations (the presence of one amino acid at one site in the alignment is correlated with the presence of another amino acid at another site). As ϕ -values move from 0 to +1, the strength of the correlation between the two amino acids increases. Negative φvalues represent negative correlations (the presence of one amino acid at one site in the alignment is correlated with the *absence* of another amino acid at a second site), which become stronger as ϕ -values move toward -1.0. Our statistical analyses (see Methods) showed that ϕ -values greater than 0.1237 were significant (P-values < 0.0001). This conservative statistical estimate revealed 13,796 significant positive covariations. However, statistical significance does not entirely indicate the strength of

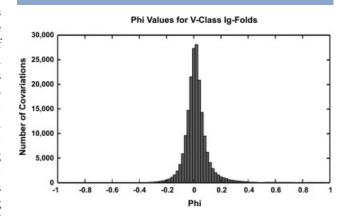


Figure 2

Distribution of ϕ -values calculated for the V-class alignment. There are 4,118,400 (20 \times 20 \times $\sum_{n=1}^{144} n - 1$) possible amino acid pairings within the V-class sequences. Of these possible pairings, 1,098,890 actually exist within the sequence database (i.e., some amino acids pairings are not observed across columns of the alignment). The histogram shows the distribution of φ-values from the 186,171 pairings that occur at least 10 times. The 13,796 φ-values greater than 0.1237 were considered statistically significant, using a conservative statistical approach (see text).

Table I Antibody V_H and V_L (Kappa) Amino Acid Pairs with the Strongest Covariations (ϕ -values)

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S7-G8 0.62 037-S67 0.47 A V63-Q81 0.62 037-K39 0.46 A-B E6-L18 0.60 037-P44 0.46 A-B G26-W47 0.60 B P44-P59 0.46 A G44-W47 0.60 A-B G57-S67 0.46 C G44-L45 0.60 A-B P59-S63 0.45 C E6-G8 0.59 Y36-L46 0.44 A-B G8-T87 0.59 B 037-G68 0.44 A	
S7-G8 0.62 0.37-S67 0.47 A V63-Q81 0.62 0.37-K39 0.46 A-B E6-L18 0.60 0.37-P44 0.46 A-B G26-W47 0.60 B P44-P59 0.46 A G44-W47 0.60 A-B G57-S67 0.46 C G44-L45 0.60 A-B P59-S63 0.45 C E6-G8 0.59 Y36-L46 0.44 A-B G8-T87 0.59 B 0.37-G68 0.44 A	
E6-L18 0.60 0.37-P44 0.46 A-B G26-W47 0.60 B P44-P59 0.46 A G44-W47 0.60 A-B G57-S67 0.46 C G44-L45 0.60 A-B P59-S63 0.45 C E6-G8 0.59 Y36-L46 0.44 A-B G8-T87 0.59 B 0.37-G68 0.44 A	
G26-W47 0.60 B P44-P59 0.46 A G44-W47 0.60 A-B G57-S67 0.46 G44-L45 0.60 A-B P59-S63 0.45 E6-G8 0.59 Y36-L46 0.44 A-B G8-T87 0.59 B Q37-G68 0.44 A	
G44-W47 0.60 A-B G57-S67 0.46 G44-L45 0.60 A-B P59-S63 0.45 E6-G8 0.59 Y36-L46 0.44 A-B G8-T87 0.59 B 0.37-G68 0.44 A	
G44-L45 0.60 A-B P59-S63 0.45 E6-G8 0.59 Y36-L46 0.44 A-B G8-T87 0.59 B 0.37-G68 0.44 A	
E6-G8 0.59 Y36-L46 0.44 A-B G8-T87 0.59 B 0.37-G68 0.44 A	
G8-T87 0.59 B 0.37-G68 0.44 A	
0.81–M82 0.59 P44–I75 0.44 A	
=*: ···*=	
W103-V109 0.59 A B P44-G57 0.43 A	
G8-L18 0.58 Q6-P8 0.42	
G106–T107 0.58 Y36–F98 0.42 A–B	
W103–Q105 0.58 A–B Q37–I48 0.42 A–B	
G8-G26 0.57 0.37-S63 0.42 A	
G26–T87 0.57 B 148–175 0.42 A	
T87–W103 0.57 B A P44–S67 0.41 A	
G106–V109 0.57 B G64–I75 0.41	
P14–W47 0.56 B A P8–Y36 0.40 B	
G26–E46 0.56 B T5–Q37 0.40 B	
R38-E46 0.56 A-B 037-R54 0.40 A	
E46–W47 0.56 A–B P59–I75 0.40	
E46–T87 0.56 A B P44–F98 0.39 A–B	
V37–L45 0.54 A–B P44–S63 0.39 A	
G8-G10 0.53 B 148-S63 0.39 A	

The two columns labeled "Top VH [VL] covarying amino acids" list the amino acids in the format A-B, and provide the residue codes and Kabat positions. Entries in the columns " $V_H - V_L$ Interface," " $V_H - C_H 1$ domain interface," and " $V_L - C_L$ domain interface identify amino acids (A, B, or both of each pair) near the specified interface.

the covariations. After careful evaluation of the data, we designated ϕ -values between 0.25 and 0.5 as moderate covariations, and ϕ -values greater than 0.5 as strong covariations. Of the 4.1 million possible amino acid combinations within the V-class Ig-fold alignment, 3212 (0.078%) had ϕ -values ≥ 0.25 and 133 (0.003%) had ϕ values ≥ 0.5 .

As validation of our covariation analysis, we examined the data in two ways to confirm that expected patterns were present in the results. First, we investigated the relationship between φ-values and distance between amino acid pairs in 3D space. Previous studies have shown that strongly covarying amino acid pairs often involve positions that are near each other in 3D structures, although the trend has invariably been reported as weak.^{23,40} To see if the same pattern was present in our data, we plotted the ϕ -values ≥ 0.3 against the distance between the amino acid pairs in 3D space using our Fab crystal structure. As reported by others, we found a weak but significant relationship between φ-value and the 3D proximity of the pair members (data not shown). Second, we investigated whether our covariation data recapitulated an amino acid network known to exist within a particular subset of V-class Ig-folds. The example we chose was a set of five residues (residues 6–10, Kabat numbering⁴⁵) at the N-terminus of human/murine IgG V_H domains. These residues adopt different backbone conformations depending on the presence of specific amino acid pairs. 46 Four conformations exist, depending on whether glutamic acid or glutamine is present at $V_{\rm H}$ position 6. Q6 is not well-conserved within $V_{\rm H}$ domains (it is also found commonly in V_L domains) and does not have significant covariations. However, E6 is highly conserved in variable heavy chains ($V_{\rm H}$ 2, $V_{\rm H}$ 3, and $V_{\rm H}$ 4 subclasses in particular). E6 correlations with residues 7–10 yield some of the highest ϕ -values of the covariation dataset (S7 = 0.51; G8 = 0.59; G9 = 0.65; and G10 = 0.53; Table I). These high correlations among residues 6-10 are consistent with the known involvement of these residues in determining the N-terminal B-strand conformation of IgG V_H domains. High φ-values were also found among other positions known to be structurally important including V_H subfamily-dependent core positions 18, 63, 67, and 82 that have been described previously.⁴/

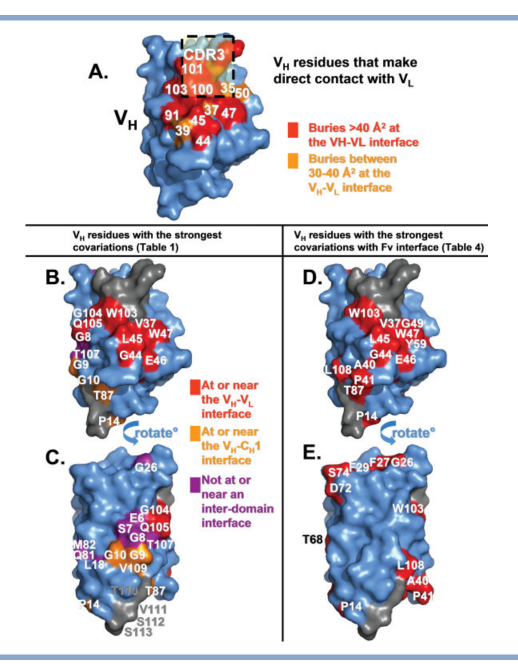


Figure 3

Covariations mapped to surface representations of an antibody $V_{\rm H}$ domain derived from an in-house Fab structure. (A) Surface representation of a $V_{\rm H}$ domain. Residues that bury >40 Å² at the Fv interface are shown in red and those that bury between 30 and 40 Å² are shown in orange. In (B-E), residues colored grey (CDR3 residues as well as four residues at the C-terminus) were not match states in the HMM-derived V-class alignment and were not evaluated in this study. (B,C) VH residues from amino acid pairs with the highest ϕ -values from Table I were mapped to the $V_{\rm H}$ surface: red, proximal to the $V_{\rm H}$ - $V_{\rm L}$ interface; orange, proximal to the $V_{\rm H}$ - $C_{\rm H}$ 1 interface; and purple, distant from the two interfaces. (D,E) $V_{
m H}$ residues from Table IV that display multiple covariations (ϕ -value >0.25) with $V_{
m H}$ - $V_{
m L}$ interface residues with greater than average ϕ -values are mapped onto the V_H surface in red. Residues from Tables I and IV that are completely buried in the interior of the structure are not shown.

Covariation results broadly applied to V_{H} and V_{L} domains

In this section, we describe patterns evident on examining the strongest covariations found within $V_{\rm H}$ and $V_{\rm L}$ domains. The $V_{\rm H}$ and $V_{\rm L}$ amino acid pairs with the highest φ-values are listed in Table I. The locations of the residues involved in the strongest covariations from Table I were mapped onto our in-house Fab 3D structure $[V_H,$ Fig. 3(B,C); V_L , Fig. 4(B,C)]. Interestingly, most of the residues contributing to the strongest covariations were found at or very near to the V_H-V_L interface [Table I, red in Figs. 3(B,C) and 4(B,C)], indicating a conserved amino acid network supporting this interface. $V_{\rm H}$

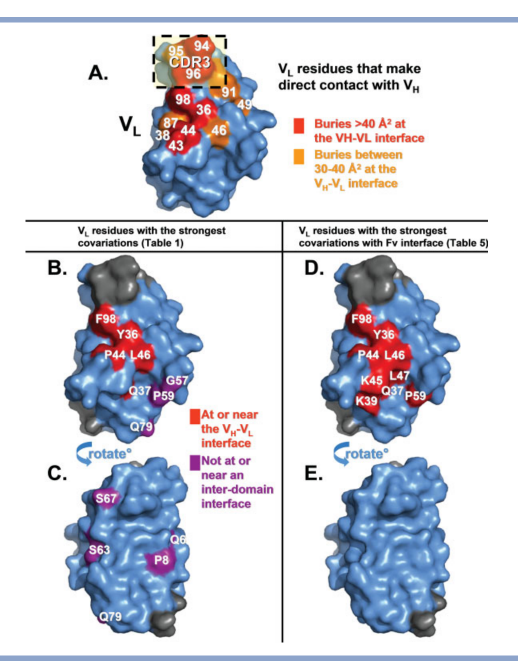


Figure 4

Covariations mapped to surface representations of an antibody V_L domain derived from an in-house Fab structure. A: Surface representation of a $V_{\rm L}$ domain. Residues that bury >40 Å² at the Fv interface are shown in red and those that bury between 30 and 40 Å² are shown in orange. In (B-E), residues colored grey (CDR3 residues as well as four residues at the C-terminus) were not match states in the HMM-derived V-class alignment and were not evaluated in this study. (B,C) $V_{\rm L}$ residues from amino acid pairs with the highest ϕ -values from Table 1 were mapped to the $V_{\rm L}$ surface: red, proximal to the $V_{\rm H^-}V_{\rm L}$ interface; orange, proximal to the $V_{\rm L^-}C_{\rm L}$ interface; and purple, distant from the two interfaces. (D,E) $V_{\rm L}$ residues from Table V that display multiple covariations (ϕ -value >0.25) with V_H - V_L interface residues with greater than average ϕ -values are mapped onto the V_L surface in red. Residues from Tables I and V that are completely buried in the interior of the structure are not shown.

domains also appear to conserve an amino acid network near the variable-constant domain (V_H-C_H1) interface [Table I, orange in Fig. 3(B,C)]. This latter network, however, is not observed in V_L domains (Table I). Other strongly conserved residue pairs involved the N-terminal region of $V_{\rm H}$ (residues 6–10), and a few buried hydrophobic residues known to be highly subtype dependent (both described earlier).^{23,47}

For an alternative overview, we also compiled tables of $V_{\rm H}$ or $V_{\rm L}$ domain residues that had the most covariations (ϕ -values \geq 0.25) with other amino acids in their respective domains (V_H, Table II; V_L, Table III), regardless of

Table II Top 40 V_H Amino Acid Positions with the Most Covariations (ϕ -value > 0.25) with Other V_H Residues

Amino		#Links	A	#Interface	A
acid	Kabat#	#LINKS all	Avg. φ-value	#Interrace links	Avg. φ-value to interface
G	10	74	0.39	3	0.34
G	8	74	0.35	4	0.37
T	87	71	0.38	6	0.44
W	103	69	0.36	6	0.36
M	82	69	0.39	1	0.41
G	26	67	0.37	6	0.46
Υ	59	66	0.36	5	0.41
V	63	64	0.38	1	0.33
Q	81	63	0.36	2	0.31
W	47	61	0.37	6	0.54
E	46	61	0.36	6	0.41
R	19	60	0.36	1	0.33
L	18	60	0.38	3	0.36
<u> </u>	69	56	0.34	5	0.32
T	68	56	0.33	4	0.38
G	49	56	0.33	5	0.43
E	6	56	0.37	1	0.39
I	51	55	0.35	5	0.38
Y	79	54	0.34	3	0.38
S	62	54	0.36	3	0.36
G	16	54	0.34	1	0.27
D	72	52	0.34	5	0.39
A	40	52	0.35	4	0.40
G	65	50	0.33	3	0.32
V	37	50	0.35	5	0.54
R	38	49	0.34	5	0.39
S	17	48	0.33	2	0.30
S	7	47	0.34	4	0.30
K	43	46	0.34	2	0.36
A	24	46	0.33	3	0.33
F	27	45	0.32	5	0.35
L	4	45	0.32	3	0.29
S	21	44	0.33	2	0.32
V	109	44	0.32	2	0.46
F	29	43	0.33	5	0.41
Q	105	42	0.32	3	0.38
R	71	39	0.32	1	0.28
S	25	39	0.32	3	0.38
L	82c	38	0.32	2	0.33
K	75	38	0.32	1	0.29

Residues that bury surface area at the Fv interface are highlighted in black rows. Residues immediately adjacent in primary sequence to interface residues are in grey rows.

the covariation strengths. Residues with the most covariations do not map to any single region of V_H (Supp. Info. Fig. 2) or V_L (Supp. Fig. 3), which is not surprising given the inclusion of many residues from weakly conserved networks. However, this analysis revealed an abundance of conserved networks evident with less stringent constraints on significant φ-values. We did note that residues involved in the most covariations mapped predominately to the β-sheet regions. Some covariations involve amino acids in the loop regions, but these are seen less frequently and likely reflect poorer alignment statistics in the loop regions, rather than a lack of conserved networks in the loop regions.

Analysis of the interface between antibody V_H and V_L domains

To further investigate $V_{\rm H}$ and $V_{\rm L}$ residues that may play a role in heavy and light chain association, we examined which $V_{\rm H}$ or $V_{\rm L}$ amino acids covary with amino acids that make direct contacts across chains at the V_H - V_L interface. Framework V_H and V_L positions that bury surface area at the $V_{
m H}\!-\!V_{
m L}$ interface are in $V_{
m H}$ 35, 37, 39, 44, 45, 47, 50, 91, and 103; and in V_L 36, 38, 43, 44, 46, 49, 87, and 98. These positions are mapped onto the surfaces of V_H and V_L in Figures 3(A) and 4(A), respectively, and onto a V-class sequence alignment (using our in-house V-class HMM) in Figure 5 (red letters in framework regions). The CDR3 loops of both $V_{\rm H}$

Table III Top 40 V_L Amino Acid Positions with the Most Covariations (ϕ -value > 0.25) with Other V_L Residues

Amino acid	Kabat#	#Links all	Avg. φ-value	#Interface links	Avg. φ-value to interface
G	64	47	0.4	6	
G	64 57	47	0.4	6	0.36 0.33
S	22	44 44	0.34	0	0.33
V	104	44	0.33	0	0
v P	59	43	0.35	6	0.35
Q	100	43	0.33	0	0.33
0	37	41	0.32	6	0.35
P	44	40	0.33	6	0.35
S	65	36	0.31	0	0.00
S	67	36	0.35	5	0.33
G	68	35	0.32	4	0.28
Ĭ	75	35	0.32	4	0.36
İ	48	34	0.33	4	0.35
Υ	36	33	0.33	6	0.37
R	54	30	0.33	3	0.31
Р	15	29	0.35	0	0
K	39	28	0.32	4	0.3
S	63	28	0.33	3	0.32
T	5	27	0.32	2	0.31
Q	79	25	0.31	3	0.3
P	8	23	0.31	4	0.32
Q	89	23	0.31	2	0.28
S	10	21	0.31	1	0.32
S	56	21	0.31	2	0.28
1	21	20	0.29	2	0.28
G	66	19	0.31	0	0
A	43	18	0.29	3	0.31
T	74	18	0.3	2	0.27
S	14	17	0.33	1	0.36
F	62	17	0.31	0	0
T	72	17	0.3	1	0.25
L F	46	15	0.3	4	0.32
F	98	14	0.33	4	0.36
0	6	13	0.3	1	0.39
0	42	13	0.28	0	0
K	45	13	0.29	3	0.28
Y V	49	13	0.29	2	0.27
	58	11	0.3	2	0.25
D	70	11	0.3	0	0
Α	84	11	0.31	0	0

Residues that bury surface area at the Fv interface are highlighted in black rows. Residues immediately adjacent in primary sequence to interface residues are in grev rows.

V _⊬ region	Framework (FW) 1	CDR 1	Framework 2
Kabat # V,,	wanne e5155466765575	27 28 282 22 Z	22222222
V _H 4 cons.	QLQESGPGLVKPSETLSLTC	TV-SG-GSISSYYWS	WIRQ-PP-GK-GLEWIGYIY
V _H 2 cons.	QLKESGPALVKPTQTLTLTC	TF-S <mark>G-F</mark> SLSTSGVGVG	WIRQ-PP-GK-ALEWLELID
V _H 3 cons.	QLVESGGGLV PGGSLRLSO	CAA-S <mark>G-FTFSSYAMS</mark>	W <mark>VRQ-AP</mark> -GK- <mark>GL</mark> EWVSAIS
V _H 5 cons.	QLVQSGAEVKKPGESLKISC	KG-S <mark>G-YSFTSYWIG</mark>	W <mark>VRQ-MP</mark> -GK- <mark>GL</mark> EWMGIIY
V _H 1a cons.	QLVQSGAEVKKPGSSVKVSC	KA-S <mark>G</mark> -GT <mark>F</mark> SSYAIS	W <mark>VRQ-AP</mark> -GQ- <mark>GLEWMG</mark> GII
V _H 1b cons.	QLVQSGAEVKKPGASVKVSO	KA-S <mark>G</mark> -YT <mark>F</mark> TSYYMH	WVRQ-AP-GQ-GLEWMGWIN
V _{HI} cons.	QLVESGGGSV GGSLRLSC	AA-S <mark>G-YTYSTY</mark>	Wiro-ap-gk-ireevain
V _H (HEWL)	QLQESGPSLVKPSQTLSLTC	SV-T <mark>G-</mark> SVTSTWWS	WIRK-FP-GN-KLEYMGYIS
V _L K1 cons.	QMTQSPSSL-SASVGDRVTITO	RA-SQGISYLA	WYQQ-KP-GK-AFKLLIYAA
V _L K2 cons.	VMTQSPLSL-PVTPGEPASISO	RS-SQSLLHSNGYNYLD	WYLQ-KP-GQ-SPQLLIYLG
V _L K3 cons.	VLTQSPATL-SLSPGERATLSO	RA-SQSVSSSYLA	WYQQ-KP-GQ-APRLLIYGA
V _L K4 cons.	VMTQSPDSL-AVSLGERATING	RS-SQSVLYSSNNKNYLA	WYQQ-KP-GQ-PPKLLIYWA
V _L λ3 cons.	ELTOPP-SV-SVAPGOTARISO	SGDALGDKYAS	WYQQ-KP-GQ-APVLVIYDD
V _L λ2 cons.	ALTOPA-SV-SGSPGQSITISO	TG-TS-SDVGGYNYVS	WYQQ-HP-GK-APKIMIYDV
V _L λ1 cons.	VLTQPP-SV-SGAPGQRVTISC	SG-SS-SNIGSNYVS	WYQQ-LP-GT-APKLLIYDN
Kabat # V	warehous states	24 27 27 27 27 27 27 27 27 27 27 27 27 27	2000 00 00 00 00 00 00 00 00 00 00 00 00
V₁ region	Framework 1	CDR 1	Framework 2

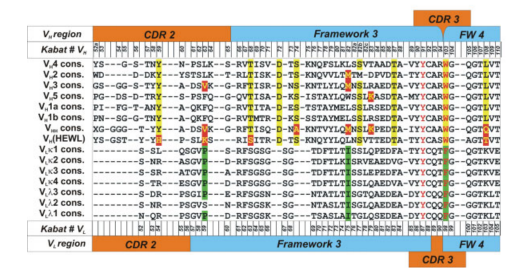


Figure 5

Sequence alignments of $V_{\rm H}$ and $V_{\rm L}$ sequences using an in-house V-class Ig-fold HMM. The panel includes representative camelid $V_{\rm HH}$ sequences and a soluble anti-HEWL $V_{\rm H}$ sequence for comparison with the consensus $V_{\rm H}$ domains. 48 Residues colored red are those that bury a significant amount of surface area at the interface between V_H and V_L. Residue positions highlighted in yellow (V_H) or green (V_L) strongly covary with many residues that bury surface area at the Fv interface (from Tables IV and V). Camelid $V_{\rm HH}$ and soluble anti-HEWL $V_{\rm H}$ residues colored yellow and highlighted in red are involved in similar residue position networks, but with different amino acids from classical $V_{\rm H}$ domains at those positions (from Table VI). VH subclass consensus residues that match the camelid or anti-HEWL residues at those positions are identically colored and highlighted. Only residue positions that were match states in the in-house, structure-based HMM are listed in the alignment. Positions of the framework and CDR regions are shown above and below the V_H and V_L sequences, respectively.

and $V_{\rm L}$ also bury surface area between the two domains to form a continuous antigen-binding surface. However, the HMM profile eliminated the CDR3 loops of both the $V_{\rm H}$ and $V_{\rm L}$ domains from the alignments due to the inability to define consistent CDR3 profiles. The absence of CDR3 data were deemed unimportant, as few strong intradomain covariations would be predicted to arise from the highly variable CDR3 loops.

The $V_{\rm H}$ and $V_{\rm L}$ amino acids with the most covariations (ϕ -value ≥ 0.25) to the interface residues above are listed in Table IV $(V_{\rm H})$ and Table V $(V_{\rm L})$. The entries in these tables are sorted by (1) the difference between the entry's average ϕ -value with interface residues versus its overall average φ-value and (2) the number of the entry's covariations with interface residues. Amino acids near the top of the tables are perceived to have a greater role in supporting the V_H - V_L interface. The residues from Tables IV and V are highlighted in the sequence alignment in Figure 5 (yellow or green highlights for VH or V_L, respectively) and have been mapped to the surfaces of the $V_{\rm H}$ and $V_{\rm L}$ domains [$V_{\rm H}$, Fig. 3(D,E); $V_{\rm L}$, Fig. 4(D,E)]. Several interface residues themselves rank highly

Table IV $V_{\rm H}$ Residues with Multiple Covariations (ϕ -value > 0.25) with $V_{\rm H}$ Residues that Bury Surface Area at the Fv Interface

Amino acid	Kabat#	#Links w/ interface residues ^a	Avg. φ-value w/interface residues ^a	#Links w/all positions ^a	Avg. φ-value w/all positions ^a	$\Delta \phi$ -value (interface-all)	Interface
V(I)	37	5(3)	0.54 (0.30)	50 (19)	0.35 (0.34)	0.19	Х
W	47	6	0.54	61	0.37	0.17	Χ
L	45	5	0.53	33	0.36	0.17	Χ
G	44	4	0.52	31	0.35	0.17	Χ
Р	14	6	0.46	29	0.35	0.11	C _H 1
G	49	5	0.43	56	0.33	0.10	X - 1
G	26	6	0.46	67	0.37	0.09	
F	29	5	0.41	43	0.33	0.08	
S	74	5	0.39	26	0.32	0.07	
T	87	6	0.44	71	0.38	0.06	C _H 1
E	46	6	0.41	61	0.36	0.05	X - 1
Υ	59	5	0.41	66	0.36	0.05	
R	38	5	0.39	49	0.34	0.05	X + 1
D	72	5	0.39	52	0.34	0.05	
A	40	4	0.40	52	0.35	0.05	X + 1
T	68	4	0.38	56	0.33	0.05	
Р	41	6	0.33	32	0.30	0.03	
I	51	5	0.38	55	0.35	0.03	X + 1
F	27	5	0.35	45	0.32	0.03	
L	108	5	0.34	12	0.31	0.03	C _H 1
S	82b	4	0.32	24	0.29	0.03	
W	103	6	0.36	69	0.36	0.00	Χ
Nonspecific							
Υ	32	5	0.30	15	0.29	0.01	
G	55	5	0.30	29	0.30	0.00	
I	69	5	0.32	56	0.34	-0.02	
S	7	4	0.30	47	0.34	-0.04	
G	8	4	0.35	74	0.39	-0.04	
L	11	4	0.30	30	0.31	-0.01	
L	11	4	0.30	30	0.31	-0.01	
0	39	3	0.27	12	0.29	-0.02	Х

Residues are sorted based on the difference between their average φ-value with interface residues versus their average φ-value with all positions within the V-class alignment. Residues that bury surface area at the interface are highlighted in black and marked with an X in the final column. Residues that are adjacent in primary sequence to interface residues are highlighted in dark grey and marked with an $X\pm 1$ in the final column. Residues at the C_H1 interface are in light grey rows. "Non-specific" residues (bottom of table) are those falling below an arbitrary cutoff above which residues appear to have strong, specific connections with interface residues. This cutoff was chosen based on a Δφ-value ([average φ-value with interface residues] - [average overall φ-value]) ≤0.01. W103 was grouped with the specific interface residues because it is an interface residue.

as do many of the residues adjacent in primary sequence. W47_{VH}, which incidentally is the $V_{\rm H}$ framework residue that buries the second highest amount of surface area at the interface, appears to be the central node of the V_{H} side of the interface network based on the number and strength of its covariations with other interface residues, even though it is not at the center of the residues that make direct contact within the V_H-V_L interface [Fig. 3(D)]. Y36_{VL} and P44_{VL} appear to be the central nodes of the V_L-side of the interface network based on the same criteria [Fig. 4(D)]. P14_{VH}, T87_{VH}, and L108_{VH}, all near the V_H-C_H1 interface, also covary strongly with the V_H interface residues, suggesting that the maintenance of both the Fv and V_H-C_H1 interfaces are co-conserved traits. In contrast, covariations were weak for V_L residues in the proximity of the V_L – C_L interface.

Four V_H residues, V37, G44, L45, and W47, form a patch on the surface of $V_{\rm H}$ that interacts with $V_{\rm L}$ [Fig.

3(A–C)]. Each of these four residues has 30 or more φvalues >0.25 with other $V_{\rm H}$ residues; however, the ϕ -values between these four residues are collectively the strongest observed for each of these residues, with an average φ-value of 0.6 (see Tables I and IV). The side chains do not pack directly against one another or appear to interact strongly, suggesting that the residues covary for functional reasons—in this case enabling immunoglobulin heavy chain V_H domains to interact with immunoglobulin light chain V_L domains. In this context, the sidechains of W47, L45, and V37 together form a roughly flat hydrophobic surface that matches well with residues P44, F87, and F98 of $V_{\rm L}$ (Fig. 6). A fifth $V_{\rm H}$ residue— W103, one of the only other $V_{\rm H}$ residues burying surface area at the interface with V_L —also covaries with the four $V_{\rm H}$ residues, though with weaker ϕ -values averaging 0.38.

Two other V_H residues—R38 and E46—strongly covary with all five of the V_H residues discussed above, with

^aMinimum φ-value cutoff of 0.25.

Table V $V_{\rm L}$ Residues with Multiple Covariations (ϕ -value > 0.25) with $V_{\rm L}$ Residues that Bury Surface Area at the Fv Interface

Amino acid	Kabat#	#Links w/ interface residues	Avg. φ-value w/interface residues ^a	#Links w/all positions	Avg. φ-value w/all positions ^a	$\Delta \varphi ext{-value}$ (interface-all)	Interface
Υ	36	6	0.37	33	0.33	0.04	Х
l	75	4	0.36	35	0.32	0.04	
L	47	2	0.33	9	0.29	0.04	X + 1
F	98	4	0.36	14	0.33	0.03	Х
P	44	6	0.35	40	0.33	0.02	Χ
Ĺ	46	4	0.32	15	0.30	0.02	Χ
P	59	6	0.35	43	0.35	0.00	
Q	37	6	0.36	41	0.37	-0.01	X + 1
	48	4	0.32	34	0.33	-0.01	X + 1
K	39	4	0.29	28	0.32	-0.03	X + 1
K	45	3	0.28	13	0.39	-0.11	X - 1
Nonspecific							
G	57	6	0.33	44	0.34	-0.01	
S	67	5	0.33	36	0.35	-0.02	
K(R)	103	3 (1)	0.32 (0.27)	4 (1)	0.34 (0.27)	-0.02	
P	8	4	0.32	23	0.31	-0.01	
G	64	4	0.28	35	0.32	-0.04	
G	68	4	0.29	35	0.32	-0.03	
D	85	4	0.32	36	0.36	-0.04	
R	54	3	0.31	30	0.33	-0.02	
S	63	3	0.32	28	0.33	-0.01	
Q	79	3	0.30	25	0.31	-0.01	
S	56	2	0.28	21	0.31	-0.03	

Residues are sorted based on the difference between their average ϕ -value with interface residues versus their average ϕ -value with all positions within the V-class alignment. Residues that bury surface area at the interface are highlighted in black and marked with an X in the final column. Residues that are adjacent in primary sequence to interface residues are highlighted in grey and marked with an $X\pm 1$ in the final column. "Non-specific" residues (bottom of table) are those falling below an arbitrary cutoff above which residues appear to have strong, specific connections with interface residues. This cutoff was chosen based on a Δφ-value ([average φ-value with interface residues] - [average overall ϕ -value]) \leq 0.00. Q37, I48, K39, and K45 were grouped with the specific interface residues because they are adjacent in primary sequence to interface residues.

average φ-values of 0.42 and 0.47, respectively. The φvalue between R38 and E46 is also strong [$\phi = 0.56$, Table I, Fig. 3(A-C)]. R38 is almost completely buried by E46 in the interior of $V_{\rm H}$ and its guanidinyl group forms a specific salt bridge with E46's carboxyl group (Fig. 6). The charge burial of R38 is supported by other interactions with D90 and K/Q43. The E46 side chain does not contact any V_H residues at the V_H-V_L interface; thus, E46 is likely important for creating optimal surface topology and perhaps an electrostatic component important for $V_{\rm L}$ binding.

On the V_L side of the interface, the most strongly covarying interface residues are Y36 and P44 [the positional equivalents of V_H residues V37 and L45—Table I, Fig. 4(A–C)] with average ϕ -values with other V_L interface residues of 0.43 and 0.42, respectively. Also covarying with these two residues are Q37, A43, L46, and F98 (positional equivalents of $V_{\rm H}$ R38, G44, W47, and W103, Fig. 5), but with lower average φ-values (0.36, 0.31, 0.33, and 0.35, respectively). The φ-values between residues within the V_L domains were lower on average than those between V_H residues; this can be explained by the number of V_L sequences in the alignment being smaller (half

the number of the V_H sequences) and more heterogeneous (containing both V_{κ} and V_{λ} domains). Similar to what was observed for V_H interface residues, a cluster of V_L residues—Y36, A43, P44, L46 and F98—do not pack directly against one another, but instead combine to form the $V_{\rm H}$ binding surface. Unlike $V_{\rm H}$ residues R38 and E46 that form a salt bridge with one another, $V_{\rm L}$ positions 37 and 45 only weakly covary with one-another (Q37 and K45 have a ϕ -value = 0.34, Fig. 5). In general, the $V_{\rm L}$ residues important for $V_{\rm H}$ -binding are well conserved for both kappa and lambda light chain variable domains.

The strongly correlated residues at the Fv interface of both $V_{\rm H}$ and $V_{\rm L}$ thus appear to form conserved networks that enable recognition between the domains (Fig. 6). The interaction surface is fairly flat between the two domains. V_H_L45 and W103 insert themselves into a small groove created by V_L residues Y36, P44, L46, F87, and F98. The small size of V_L_P44 helps create the groove into which V_H_L45 and V_H_W103 intrude. In addition, the relatively small $V_{\rm H}$ V37 side chain creates a cavity on the hydrophobic surface of V_H into which the side chain of V_L F98 inserts (Fig. 6).

^aMinimum φ-value cutoff of 0.25

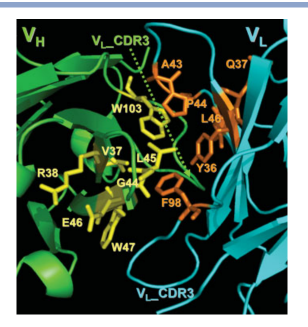


Figure 6

Structural view of the $V_{\rm H}$ and $V_{\rm L}$ residues from Tables IV and V. The polypeptide backbone of the $V_{\rm H}$ domain (green) and $V_{\rm L}$ domain (blue) are depicted using a cartoon ribbon diagram. VH residues V37, R38, G44, L45, W47, and W103 are displayed in the stick format in yellow. V_L residues Y36, Q37, A43, P44, L46, and F98 are displayed in the stick format in orange.

Comparison of conserved residues networks of V_H domains and camelid V_{HH} domains lacking light chain interactions

Although most antibody V_H sequences associate with light chain $V_{\rm L}$ s, a subset of camelid variable heavy chain domains, denoted V_{HH} domains, are expressed naturally and function in the absence of both a light chain and a $C_{\rm H}1$ domain. ⁴⁹ $V_{\rm HH}$ domains are also substantially more soluble than $V_{\rm H}$ domains. The discovery of these simple and soluble V_H-like domains has had an enormous impact on antibody engineering because they represent potentially more facile reagents for protein design and discovery than traditional antibodies (which require combinations of heavy and light chains for function, stability, and solubility⁵⁰). We therefore investigated whether conserved residue networks differ between standard $V_{\rm H}$ and camelid $V_{\rm HH}$ domains. We expected to observe such differences at the positions in $V_{\rm H}$ that serve to support the V_H-V_L interface. Our results revealed 32 significant covariations involving identical positions within $V_{\rm H}$ and $V_{\rm HH}$ domains; however, the 32 covarying pairs contained different amino acids for $V_{\rm H}$ versus $V_{\rm HH}$ domains (Table VI). Among these contrasting residues are a tetrad of amino acids that have been previously reported to differentiate $V_{\rm H}$ from $V_{\rm HH}$ domains: V37F, G44E, L45R, and W47G. Substitution of this tetrad of camelid amino acids into V_H domains does not entirely impart them with the solubility of $V_{\rm HH}$ domains; CDR3 composition and other factors have also been shown to be important, 52-56 Our covariation results reveal additional framework residue positions, outside the tetrad described earlier, that naturally distinguish $V_{\rm H}$ from $V_{\rm HH}$ domains. These residues are at positions 13, 14, 33, 49, 63, 74, 82, 83, and 108. Solubilizing mutations at residues 74 and 108 have been reported.⁵⁷ Most of these positions are involved in networks surrounding the V_H - V_L or V_H - C_H 1 interfaces (Table VI, Fig. 3), as expected. A consensus camelid $V_{\rm HH}$ sequence derived from the ${\sim}50$ diverse camelid sequences in the V-class alignment is included in Figure 5 to highlight the positions of these observed differences between $V_{\rm H}$ and $V_{\rm HH}$ domains.

A natural human V_H raised against hen egg-white lysozyme was also demonstrated to be soluble in a monomeric form, similar to camelid domains (although the domain presumably maintains its ability to associate with $V_{\rm L}$). 52,58 This independently soluble anti-HEWL $V_{\rm H}$ domain contains yet another set of nonstandard V_H amino acids. Many of these residues are involved in the Fv interaction network and one is proximal to the V_H-C_H1

Contrasting Features of V_H and Camelid V_{HH} Domains Based on Covariation Analyses

		Camelid $V_{ m HH}$	
$V_{\rm H}$ linked pair	φ-value	linked pair	φ-value
G44-L45	0.61	E44-R45	0.57
L45-L108	0.29	R45-Q108	0.57
V(I)37-L45	0.54	F37-R45	0.50
P14-V37	0.40	A14-F37	0.50
L45-W47	0.70	R45-G47	0.46
V37-W47	0.65	F37-G47	0.44
_	_	C33 ^a –G47	0.44
_	_	A14-Q108	0.44
P14-G44	0.35	A14-E44	0.43
G44-W47	0.61	E44-G47	0.42
K13 ^b –L45	0.31	Q13-R45	0.42
V37-G44	0.53	F37-E44	0.40
_	_	C33 ^a –R45	0.40
L82 ^b -L45	0.29	M82-R45	0.38
V37-L108	0.32	F37-Q108	0.37
G49-L45	0.46	A49-R45	0.36
W47-L108	0.35	G47-Q108	0.36
L63 ^b -L45	0.30	V63-R45	0.36
_	_	C33 ^a –F37	0.36
P14-W47	0.48	A14-G47	0.35
_	_	C33 ^a –E44	0.35
K13 ^b -G44	0.30	Q13-E44	0.34
_	_	Q13-F37	0.32
S74-L45	0.45	A74-R45	0.30
L82 ^b -G44	0.29	M82-E44	0.27
S74-L108	0.25	A74-Q108	0.27
_	_	K83-E44	0.27
K13 ^b -W47	0.36	Q13-G47	0.26
_	_	V63-E44	0.26
S74-G44	0.38	A74-E44	0.25
S74-V37	0.33	_	_

^aC33 often makes a disulfide with CDR3 in camelid V_{HH} domains to stabilize

 $^{{}^{\}rm b}V_{\rm H}3$ consensus matches the $V_{\rm HH}$ consensus amino acids at these positions.

interface: D27, D32, K39, K44, Y47, H59, K63, S68, and T108 (Fig. 5). Together with the $V_{\rm HH}$ results, it appears that multiple and independent amino acid networks may impart solubility to $V_{\rm H}$ and $V_{\rm HH}$ domains.

DISCUSSION

Despite an enormous amount of research involving antibodies and antibody-like therapeutics, very little use has been made of covariation analyses to investigate functional features of antibody domains. A study by Altschuh and coworkers^{59,60} investigated covariations across murine and human germline $V_{\rm H}$ or $V_{\rm L}$ sequences, with the goal of defining positions within each germline subclass that use mutually exclusive framework amino acid pairs to influence the structural conformations of CDR loops. Our study had a different goal: to use covariation analyses for determining naturally occurring amino acid networks, in antibody variable domains, that are generally important for antibody structure and function. To this end, we generated covariation data using a larger and more diverse set of V-class Ig-fold

Our results show that the most strongly conserved amino acid networks in antibody $V_{\rm H}$ and $V_{\rm L}$ domains are found at the interface between $V_{\rm H}$ and $V_{\rm L}$, suggesting that preservation of this interface is a factor influencing antibody evolution. Interestingly, a small network of amino acids near the V_H-C_H1 interface is also highly conserved. However, this network is not observed for residues near the V_L - C_L interface. Biophysical studies with light chains in isolation have shown that the $V_{\rm L}$ and $C_{\rm L}$ domains do not influence the unfolding kinetics or thermodynamics of one another, suggesting that the interaction between the two domains is weak. 61,62 Alternately, Fabs (consisting of V_H , $C_H 1$, V_L , and C_L domains) often show concerted unfolding reactions.⁴⁴ A viable explanation for the concerted unfolding reactions of Fabs compared with light chains in isolation is that the $C_{\rm H}1$ and $C_{\rm L}$ domains act as ideal linkers for the $V_{\rm H}$ and $V_{\rm L}$ domains and vice versa. 62 It may also be that stronger interactions between $V_{\rm H}$ and $C_{\rm H}1$, compared with $V_{\rm L}$ and C_L, promote cooperative unfolding of all four Fab domains.44 The covariation data described in this report demonstrate that several V_H residues at the V_H - C_H 1 junction are involved in a strongly conserved network of amino acids and suggest that Variable-Constant domain interactions may be more important for immunoglobulin heavy chains than for immunoglobulin light chains.

CONCLUSIONS

In summary, we performed covariation analyses using a large, high-quality, and diverse alignment of V-class Igfold sequences. The data were used to discover antibody variable domain amino acid networks that are evolutionarily conserved. Mapping the most highly conserved $V_{\rm H}$ and $V_{\rm L}$ networks to their structures revealed that the networks cluster near the V_H-V_L interface and near the V_H-C_H1 interface, demonstrating the importance of preserving these interfaces during the evolution of antibody sequences.

These covariation data serve as a powerful tool for antibody (and Ig-fold domain) engineering. Insights from covariation analysis have improved our ability to rationally design more stable scFvs (Supp. Fig. 4). Most scFvs contain the majority of the residues within the $V_{\rm H}$ and V_L conserved networks revealed by the covariation data. Our initial approach for stabilizing scFvs has been to find gaps or holes within these existing networks that can be rectified by mutagenesis. Many of these changes have improved the thermal unfolding midpoint $(T_{\rm M})$ of scFv V_H or V_L domains by 1-12°C (Miller et al., manuscript in preparation). Stabilization of scFvs has enabled their use as building blocks that can be appended to fulllength IgG molecules to produce stable bispecific IgGlike antibodies for consideration in clinical applications.63

The covariation data described here may also be useful for engineering other aspects of V-class Ig-fold proteins, such as soluble $V_{\rm H}$ domains with conserved $V_{\rm HH}$ amino acid networks. The approach can be extended to other Ig-fold domains, such as C- and I-class, to identify amino acid networks supporting structure and function of other immunoglobulin or cell-surface receptor domains.

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