Stability engineering of anti-EGFR scFv antibodies by rational design of a lambda-to-kappa swap of the VL framework using a structure-guided approach

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Abbreviations: scFv, single-chain variable fragment; Ig, immunoglobulin; VH, variable heavy; VL, variable light; CDR, complementarity-determining region; EGFR, epidermal growth factor receptor; PBS, phosphate-buffered saline; BSA, bovine serum albumin

Phage-display technology facilitates rapid selection of antigen-specific single-chain variable fragment (scFv) antibodies from large recombinant libraries. ScFv antibodies, composed of a VH and VL domain, are readily engineered into multimeric formats for the development of diagnostics and targeted therapies. However, the recombinant nature of the selection strategy can result in VH and VL domains with sub-optimal biophysical properties, such as reduced thermodynamic stability and enhanced aggregation propensity, which lead to poor production and limited application. We found that the C10 anti-epidermal growth factor receptor (EGFR) scFv, and its affinity mutant, P2224, exhibit weak production from E. coli. Interestingly, these scFv contain a fusion of lambda3 and lambda1 V-region (LV3 and LV1) genes, most likely the result of a PCR aberration during library construction. To enhance the biophysical properties of these scFvs, we utilized a structure-based approach to replace and redesign the pre-existing framework of the VL domain to one that best pairs with the existing VH. We describe a method to exchange lambda sequences with a more stable kappa3 framework (KV3) within the VL domain that incorporates the original lambda DE-loop. The resulting scFvs, C10KV3_LV1DE and P2224KV3_LV1DE, are more thermodynamically stable and easier to produce from bacterial culture. Additionally, C10KV3_LV1DE and P2224KV3_LV1DE retain binding affinity to EGFR, suggesting that such a dramatic framework swap does not significantly affect scFv binding. We provide here a novel strategy for redesigning the light chain of problematic scFvs to enhance their stability and therapeutic applicability.

Introduction

Antibody therapeutics represent a growing class of molecules entering clinical trials.1 Single-chain variable fragment (scFv) antibodies are composed of the variable heavy (VH) and variable light (VL) domains of an immunoglobulin (Ig) tethered by a peptide linker.2 scFvs are highly versatile in their use in basic research, diagnostics, and therapy, and they are easily engineered into higher order formats such as bispecific (including bispecific T-cell engagers) or trispecific antibodies, chimeric antigen receptors, or IgG.3-6 Identification of target-specific scFvs by various display technologies that utilize a genetic library of VH and VL gene segments is rapid and cost-efficient in comparison to conventional immunization-based strategies.7 While these antibodies are highly target specific, they are often limited in their stability and aggregation propensity, which negatively affect expression, purification, concentration, immunogenicity, and function.8-12 This is due in part to the lack of quality control mechanisms that are usually provided by an antibody-producing B cell, and results in selection of scFvs with suboptimal structural stability.13 The impaired stability may be the result of a lack of stabilizing Fab constant regions, incompatible VH/VL pairings, as well as PCR-induced aberrations such as missense mutations and gene fusions. Although the structural integrity of these antibodies may be sufficient for their use in basic research, their biophysical properties must be optimized if they are to be incorporated into clinical diagnostics and therapeutic agents.

The ErbB tyrosine kinases are often over-expressed and the key drivers of tumor promotion in multiple cancer types, particularly head and neck, lung, breast, and gastric cancers.14 This makes ErbB members particularly ideal targets for therapeutic antibody blockade. For example, the monoclonal antibodies...
cteuzumab and panitumumab block ErbB1/epidermal growth factor receptor (EGFR) signaling and are approved for use in head and neck and colorectal cancer,\textsuperscript{15-17} while trastuzumab and pertuzumab bind to domains IV and II of ErbB2, respectively, to block ErbB2 activity and are approved for breast and gastric cancers.\textsuperscript{18}

The anti-EGFR scFv, C10, was previously identified from a phage antibody library and selected for its ability to promote EGFR internalization, a valuable feature that can be utilized for antibody drug delivery.\textsuperscript{19} However, C10 scFvs and one of its affinity-matured variants, P2224, tend to aggregate.\textsuperscript{20} Because of their favorable biological properties, it would be desirable to engineer more stable variants of these antibodies prior to further stages of drug development. We examined the C10 framework sequence and determined that the V\textsubscript{L} domain is likely to be a fusion between \& \lambda3 (LV3) and \& \lambda1 (LV1) V-region genes, and that the sequence is probably a byproduct of PCR amplification during library construction. Many affinity-matured derivatives of C10 have been generated and all contain this hybrid light chain. Based on the non-native nature of this framework sequence, we propose that the V\textsubscript{L} \& \lambda3-\& \lambda1 fused framework negatively affects the overall biophysical characteristics of the C10 scFv and its derivatives. Additionally, we anticipate that the V\textsubscript{L} \& \lambda3-\& \lambda1 framework will contribute to suboptimal behavior when engineered into higher order antibody formats.

The mechanisms controlling antibody production, thermostability, and aggregation resistance are not completely understood and various methods have been employed to enhance the biophysical properties of antibodies, including stability engineering of the hydrophobic core, surface-exposed residues, and the V\textsubscript{H}:V\textsubscript{L} domain interface.\textsuperscript{1,8,21-23} Here, we describe a structure-based redesign strategy in combination with our previously described complementarity-determining region (CDR) clustering scheme\textsuperscript{24} to graft the V\textsubscript{L} CDRs of C10 and P2224 onto a more stable V\textsubscript{L} \& \kappa3 (KV3) framework. This is the first demonstration that a structure-guided redesign involving a \& \lambda to \& \kappa switch successfully improves the stability of an scFv while maintaining comparable binding affinity.

**Results**

**Selection of the Vk3 framework, C10 homology modeling, and redesign**

One potential indicator of scFv stability and propensity to aggregate is total protein yields during production.\textsuperscript{25} Under the production strategies employed in these studies, increased expression temperatures appeared to negatively affect yields of the parental scFv proteins. Yields of 2.5 mg/L (range: 2.3 – 2.6) and 0.9 mg/L (range: 0.4 – 1.5) of purified C10 scFv protein were obtained when expressions were carried out at 25 and 30 °C, respectively (Supplementary Table 1). Despite differing by only a single framework residue and 3 residues in the V\textsubscript{H} CDRs, the yields of the affinity matured P2224 were dramatically lower than those obtained with C10 at both temperatures; yields of 0.6 mg/L (range: 0.3 – 1.2) and 0.12 mg/L (range: 0.07 – 0.18) of culture were obtained at 25 and 30 °C, respectively (Fig. 1 and Supplementary Table 1). This suggested that the P2224 scFv was, like C10, unstable at higher temperatures and that the affinity maturation process further destabilized P2224 relative to the parental molecule.

To determine what elements in the protein sequence and structure might lead to low stability, we compared the protein sequences of the parental C10 light and heavy chains to human germline V-region sequences provided by IMGT.\textsuperscript{26} The C10 heavy chain sequence (through the beginning of CDR H3) is 99% identical to the human IGHV1–69*01 amino acid sequence. The C10 light chain sequence showed the highest sequence identity (87%) to \& \lambda light chain V-region gene IGLV3–19*01 through the beginning of CDR L2 but only 66% after L2 (Fig. 2A). By contrast, C10L was 51% identical to \& \lambda light chain V-region gene IGLV1–44*01 through the beginning of L2 but 100% identical after L2, including the first 9 residues of L3. The DNA sequences also showed high similarity to IGLV3–19*01 through CDR L2 and high similarity to IGLV1–44*01 after L2. We concluded that the V\textsubscript{L} sequences of C10 and its derivatives are likely to contain a fusion product of V\textsubscript{L}3 (IGLV3–19*01) and V\textsubscript{L}1 (IGLV1–44*01) V-region genes. Since this fusion is not typically found in nature and V\textsubscript{L}3 (IGLV3–19*01) and V\textsubscript{L}1 (IGLV1–44*01) V-region genes. Since this fusion is not typically found in nature and V\textsubscript{L} domains tend to confer lower production yields from bacteria and a greater propensity to aggregate,\textsuperscript{27} it was likely that the hybrid V\textsubscript{L} sequence was the source of poor production.

To improve the production of C10 and P2224, we decided to swap the V\textsubscript{L} framework of C10 with a more stable V\textsubscript{L} framework that would also likely pair well with the IGHV1–69 domain of C10. Ewert et al. found that Vk3 frameworks are the most stable V\textsubscript{L} domains when studied in isolation from the V\textsubscript{H} domain.\textsuperscript{27} They also found that the combination of IGHV1–69 (their VH1a) with IGLV3 (their Vk3) produced the highest soluble yields of all the combinations they tested, except for the VH1b/Vk3 combination (closest to IMGT IGHV1–2/IGKV3D-7), which was slightly higher.\textsuperscript{27} More recently, Tiller et al. have determined the usage frequency of each germline V region in the human antibody repertoire as well as the pairing frequency of V\textsubscript{H} and V\textsubscript{L} domains.\textsuperscript{28} Of all V\textsubscript{L} domains, they found IGVK3–20 to be the most common, followed by IGKV1–39. For pairing with IGHV1–69, these 2 domains were the most common and approximately equal (12 and 14 cases, respectively) followed by IGKV3–11 (8 cases), IGKV3–15 (7 cases), and IGKV1–5 (5 cases). They also find the highest expression levels with IGHV1–69 to be IGKV1–5, IGKV3–11, IGKV3–20, and IGKV3–15, all roughly equal. We recently assigned germline V regions to all structures in the Protein Data Bank (PDB)\textsuperscript{29} and analyzed the pairing frequency of human V\textsubscript{H} and V\textsubscript{L} domains in antibodies of known structure. In a non-redundant set of antibodies, the most common pairing with IGHV1–69 was IGKV3–20 (7 cases), followed by IGKV1–39 (4 cases) and IGKV1–33 (4 cases). While the counts in these studies are not high, we chose the IGKV3–20 framework as a promising lead for producing a stable antibody with the IGHV1–69 domain of C10.

We chose the antibody X5 from PDB entry 1RHH\textsuperscript{30} as a template for grafting the C10 CDRs, since it has been well
and is the closest IGHV1–69 domain to C10 V\textsubscript{H} that is also bound to an IGKV3–20 domain. To guide our design process, we constructed homology models of C10 and C10 with an IGKV3–20 framework based on the sequence of the light chain of PDB entry 1RHH by grafting CDRs from templates with similar sequences onto suitable frameworks. The side chain conformations for the new sequences were optimized with the program SCWRL4 (see Materials and methods). A sequence alignment of C10 and C10KV3 is shown in Figure 2B. Parent and target structures were inspected visually for inconsistencies caused by the CDR grafts.

One noticeable problem from this structural inspection was a clash between the grafted \(\lambda_3\) CDR-L1 and the acceptor framework’s \(\kappa_3\)-typical \(d\)-\(e\) loop. The \(d\)-\(e\) loop (nomenclature according to ref.\textsuperscript{32}) is a framework loop of the FR3 region (residues 66–71 in Chothia numbering) between CDR2 and CDR3 that makes extensive contacts with CDR1. A close-up of the L1 and d-e loops of the unrefined models of C10 and C10KV3 is shown in Figure 3A. The \(d\)-\(e\) loop sequence motifs in \(\kappa\) and \(\lambda\) germline sequences are distinct, i.e., G[SPY]GT[DE][FY] and [KRN][SG][NTK][STA], respectively. In C10, the sequence of the \(d\)-\(e\) loop is KSGTSA, while in C10KV3 sequence is GSGTDF. In Figure 3A, the d-e loop of the C10KV3 model (blue) exhibits a large deviation from the C10 model (magenta).

We investigated if this clash was an artifact of the particular combination of CDR donor and acceptor framework structures or if this feature is more generally true for \(\lambda_3\)-to-\(\kappa_3\) grafts. We performed a structure alignment with the program THESEUS\textsuperscript{33} of a non-redundant set of 113 \(\kappa_3\) and \(\lambda_3\) light chain variable domains (each with a different CDR L1 sequence of length 11), and the result is shown in Figure 3B. When CDR L1 is 11 residues in length, there are 3 predominant clusters.\textsuperscript{29} The 2 largest are L1–11–1 and L1–11–2, consisting entirely of L1 CDRs from \(\kappa\) light chains.\textsuperscript{24} In both of these clusters, residue 71 of the d-e loop participates in a hydrophobic cluster of amino acid side chains consisting of residue 71 (Phe or Tyr) and residues 6 and 10 of the 11-amino acid L1 loop (usually Leu, Ile, and Val). This cluster of interactions is shown in Figure 3C. Residue 71 is Phe in nearly all L1–11–1 CDR structures and Tyr in L1–11–2 structures (almost all of which are mouse frameworks\textsuperscript{29}). The Tyr hydroxyl makes a hydrogen bond with the backbone of residue 7 of the L1 loop, flipping the conformation of residues 7 and 8.
By contrast, light chains with 11-amino acid L1 CDRs exist almost entirely in cluster L1–11–3, with a distinct sequence pattern compared to L1–11–1 and L1–11–2 CDRs in κ antibodies. In the PDB, these antibodies are all human IGLV3 (except for one hamster structure and one macaque structure) since other germlines (including human IGLV1) do not have L1 CDRs of length 11. The structures of L1–11–3 CDRs are quite different from L1–11–1 and L1–11–2, with residues 5 and 10 of the CDR pointing toward each other, inwards into the VL domain core, and participating in a hydrophobic cluster with the side chain of A71 and in some cases the hydrophobic portions of K/N/I66 of the d-e loop, as shown in Figure 3D. Computationally mutating A71 to Phe in κ antibodies results in severe steric conflicts with residues 5 and 10 of the L1 CDR (not shown), indicating that the conformation of L1–11–3 is not consistent with a Phe residue at position 71 in the d-e loop.

G66 of the d-e loop is completely conserved in human κ germline sequences, while this position is never Gly in human λ germline sequences. Glycine is able to access backbone conformations that residues with side chains are not able to achieve, particularly those with backbone dihedral $f > 0^\circ$. G66 has $f > 0^\circ$ (mean = 117°, std = 15°) in 792 or 99% of 807 (redundant) human κ domains in the PDB. As shown in Figures 3A and B, G66 allows the d-e loop to bend inward, toward the L1 loop. In contrast, the α-like R/N/I66 side chains at position 66 usually hydrogen bond to the backbone carbonyls of residues 5 and/or 8 of the L1 loop, stabilizing the α-like L1–11–3 conformation. The change in backbone conformation at position 66 is evident in Figures 3A and 3B.

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It is certainly possible that small adjustments in backbone and side-chain conformations could remove the clash shown in Figure 3A, resulting in a stable C10KV3 molecule. To investigate this, we utilized RosettaAntibody to build models of C10, C10KV3, and C10KV3 with a d-e loop with C10’s λ1 sequence (C10KV3_LV1DE). For comparison, we also built models of all-λ3 and all-λ1 variants of C10 (C10LV3 and C10LV1). The sequences of these constructs are given in Figure 2B. The initial model of C10KV3_LV1DE from RosettaAntibody utilized a κ-like structure of the d-e loop, because the program used a κ3 template. To produce a better model, we grafted a d-e loop with the sequence KSGTSA (as in C10) from the IGLV1–44*01 antibody in PDB entry 4GXV3 into the RosettaAntibody model of C10KV3_LV1DE. To obtain structural scores that might be correlated roughly with stability, we applied a refinement strategy in the program Rosetta that allows for changes in bond lengths and bond angles (“Cartesian minimization”) during extensive repacking of side chains with our backbone-dependent rotamer library and local optimization of both side-chain and main-chain dihedral angles. Each run consists of optimization moves in a random order so each resulting model (or decoy) is different. Starting from the RosettaAntibody models (including the d-e loop grafted C10KV3_LV1DE), we generated 250 decoys for each target with the Cartesiam minimization protocol. A box and whiskers
plot of these scores is shown in Figure 4A, demonstrating that
the distribution of scores for C10KV3_LV1DE is a little lower
than C10KV3, and that the λ3 constructs are predicted to be
more stable than the λ1 and λ3 variants and the original C10.
Notably, the refinement of C10KV3_LV1DE moved K66 of the
grafted d-e loop into a position with hydrogen bonds to the back-
bone carbonyl oxygens of residues 5 and 8 of CDR L1, shown in
Figure 4B. Consistent with the Rosetta results, we chose to con-
struct C10KV3_LV1DE and C10KV3 as well as the similar var-
iants of the affinity-matured antibody P2224, i.e.,
P2224KV3_LV1DE and P2224KV3.

The redesigned V_L CDR framework confers enhanced
production of C10 and P2224 scFvs

As described above, increasing the culture temperature from
25 to 30 °C during expression of C10 and P2224 resulted in an
apparent decrease in absolute yields, and led to our efforts to
enhance stability of the proteins. Although absolute yields of pro-
teins varied between production runs, trends related to yields
were observed. Despite the differences in amino acid composi-
tion and codon usage associated with the redesign process C10
and C10KV3_LV1DE were expressed and purified to approxi-
mately the same level (average: 2.5 vs 2.2 mg/L culture) when
experiments were performed at 25 °C (Fig. 1 and Supplemen-
tary Table 1). In contrast, C10KV3_LV1DE was purified to
approximately 2.5-fold higher levels than C10 at 30 °C, a tem-
perature that deleteriously affected expression of the parental
C10 scFv. Incorporating the redesigned KV3_LV1DE light
chain into P2224 had a similar effect on increasing yields. Yields
of P2224KV3_LV1DE, as compared to P2224, were increased
by approximately 2.5-fold when expressions were carried out at
25 °C. The effect of the redesigned P2224KV3_LV1DE was even more
dramatic when expressions were car-
ried out at 30 °C, with the yields of
P2224KV3_LV1DE being approxi-
mately 8-fold greater than those
obtained with P2224 (Fig. 1 and Sup-
plementary Table 1). In support of
our hypothesis that incorporation of
the λ1 d-e loop into
C10KV3_LV1DE would be a critical
structure determinant, the presence of
the κ3 d-e loop in C10KV3 decreased
production of the scFv by approxi-
mately 2.5 fold (Fig. 1B and E)

The C10KV3_LV1DE and
P2224KV3_LV1DE variants exhibit
enhanced thermostability

Since enhanced production of the
KV3_LV1DE-redesigns is a promising
indicator that the reformatted frame-
work provides greater structural stabil-
ity, we utilized differential scanning
fluorimetry (DSF) to examine the
thermostability of C10 and its derivatives. As shown in Figure 5, C10KV3_LV1DE and P2224KV3_LV1DE exhibit a greater derivative Tm (approx. 64°C and 62°C, respectively) than C10 and P2224 (approx. 62°C and 58°C, respectively). Although the Tm is enhanced by only 2–4°C, the difference between the parental and redesigned mutants is significant (Fig. 5C), and demonstrates that the κ3-based framework enhances the thermostability of these scFvs. Additionally, maintaining the original λ d-e loop of C10 proved to be critical in enhancing the thermostability of C10KV3_LV1DE and P2224KV3_LV1DE as evidenced by a 5–6°C decrease in Tm of C10KV3 and P2224KV3, which contain the κ3 d-e loop (Fig. 5B and D).

To further confirm that the framework mutants are indeed more thermodynamically stable, we analyzed C10, C10KV3_LV1DE, and P2224KV3_LV1DE by circular dichroism (CD) spectroscopy. Of note, the P2224 scFv was not analyzed in this assay because it consistently demonstrated a resistance to concentration above 1 μM, a property that has been remedied by the κ3 swap. For the rest, thermal melt scans were performed at a fixed wavelength of 218 nm to monitor β-sheet unfolding. Under this set of parameters, the ellipticity of the protein solution was recorded and C10KV3_LV1DE was deemed more thermodynamically stable (Tm = 69.1°C) than C10 (Tm = 65.6°C) due to enhanced melting temperature (Fig. 6). While the concentration of P2224 was too low to accurately calculate a Tm, P2224KV3_LV1DE also demonstrated higher thermostability (Tm = 68.3°C) than C10 (Fig. 6).

In addition to measuring the ellipticity, the dynode voltage, which can be used to calculate absorbance, was concurrently recorded to measure the heat-induced aggregation of the scFvs (Fig. 7A). Increasing absorbance is caused by variations in light scattering due to increasing particle size as a result of aggregation of the unfolded protein at elevated temperature.38 Interestingly, C10 and C10KV3_LV1DE appear to have different modes of aggregation (Fig. 7A). With increasing temperature up to 80°C, C10 demonstrates a steady increase in soluble protein aggregates, while C10KV3_LV1DE and P2224KV3_LV1DE exhibit aggregation at higher temperatures that results in precipitation of insoluble protein (Fig. 7A, bell shaped curve). This different mode of protein aggregation is also reflected in the CD-detected thermal unfolding curves (Fig. 6), where C10 and C10KV3_LV1DE both appear to irreversibly unfold, while their resulting spectral signatures are qualitatively different. After heating to 80°C and cooling back to 25°C, C10 does not completely regain its native structure. In fact, C10 appears to lose signal at 205 nm back to baseline while gaining a more negative signal at 218 nm, which may reflect an increase in β like-structure or aggregation. In
contrast, C10KV3_LV1DE and P2224KV3_LV1DE show little or no residual CD signal after cooling back to 25°C (Fig. 6A), indicating that both proteins are completely precipitated at 80°C (Fig. 6B). The fact that the k3 variants precipitate out of solution upon heating is not that surprising and is similar to the mode of heat-induced aggregation and precipitation of the single chain version of trastuzumab, 4D5, which is a VH3/Vκ1 antibody (Fig. 7). Overall, the measured increases in Tm for C10KV3_LV1DE and P2224KV3_LV1DE compared to C10 further reinforce the idea that the newly introduced k3 framework increases the inherent thermostability of C10 and provides support for the use of our structure-based strategy for antibody redesign.

Both C10 and C10KV3_LV1DE form large heat-induced aggregates

Since both C10 and C10KV3_LV1DE exhibit heat-induced irreversible unfolding, while the k3 variants form visible aggregates and C10 and P2224 do not, we examined whether the k3-redesign resulted in a change in the size and solubility of the aggregates. To do this, C10 and C10KV3_LV1DE were heated to 60°C for 10 min, cooled to room temperature, and analyzed for changes in hydrodynamic radius (or mass of an equivalent sphere) by dynamic light scattering. As shown in Figures 7B and C, the majority of C10 and C10KV3_LV1DE are predominantly monomeric at 25°C. Upon heating to 60°C, very large aggregates can be seen in both C10 and C10KV3_LV1DE preparations. This unfolding and aggregation appears to be irreversible since only large aggregates are observed when samples are cooled to 25°C and no detectable levels of monomer are present. It is interesting that solubility is maintained even for the large aggregates produced by C10. Overall the collective data suggests that the k3 redesign improves protein production, thermostability, and, although both C10 and the C10KV3_LV1DE will form large heat-induced products, the aggregated products are qualitatively different with C10, generating large soluble products, while the C10KV3_LV1DE, like 4D5, precipitates upon heat-induced unfolding and aggregation.

The k3 redesigns exhibit both similar binding affinities and cetuximab blocking ability as the parental k3-κ1 scFvs

Although the k3 variants are more stable, it is possible that the CDR grafting altered the scFv target binding by changing the orientation of the CDRs. Therefore, we compared the abilities of the parental and redesigned scFv to bind to EGFR+ cells. Under the conditions employed in our assays C10KV3_LV1DE bound weakly to EGFR+ cells with a Kd of approximately 3.8 μM. However, this binding was better than that detected for the parental C10, which failed to bind sufficiently to allow for determination of a Kd (Fig. 8A). As shown in Figure 8B, the Kd for P2224, P2224KV3, and P2224KV3_LV1DE were calculated to be 2.79 nM, 1.49 nM, and 1.62 nM, respectively, thus indicating that the k3-derived framework and d-e loop do not appear to alter affinity. Together these data demonstrate that the redesigned VL framework does not reduce the ability of the C10 and P2224 CDRs to bind cell surface EGFR.

Previous studies demonstrated that P2224 elicits therapeutic activity, at least in part, by binding to domain III of EGFR’s extracellular domain and competing with ligand (e.g., EGF) binding.20 Its epitope on domain III is sufficiently close to the epitope bound by cetuximab that P2224 can also compete with cetuximab for binding to EGFR.39 To confirm that the k3 redesign did not dramatically alter the binding epitope, we examined
whether P2224 and its κ3 derivatives retained the ability to block cetuximab binding. As demonstrated in Figure 8C, both P2224 and P2224CKV3_LV1DE block cetuximab binding in equivalent concentration-dependent manners. Collectively, these data demonstrate that our rational redesign of C10 and P2224 enhances protein production and thermostability without altering the antibodies’ abilities to bind their target antigen.

Discussion

Our previous work took advantage of the large diversity of crystal structures available in the PDB to expand upon the original findings of Chothia and allowed us to delineate a sequence/structure relationship between CDRs. Here, we demonstrate that combining this CDR clustering with an analysis of CDR/framework interactions correctly guides the stability engineering of the C10 series of anti-EGFR scFvs. Molecular modeling and extensive refinement with Rosetta of C10 and λ1, λ3, and κ3 variants indicated that the κ3 variants with and without the λ1 d-e loop were the most promising candidates for engineering a stable platform for the P2224 series of antibodies.

Many properties, including affinity, effector function, and pharmacokinetics, must be considered while engineering a therapeutic antibody. Display techniques coupled with advances in antibody engineering have led to rapid, robust methods for both isolating antibodies against targets of interest and tailoring their behavior. Significant effort has been devoted to understanding the effect of affinity on biological activity and developing methods to tailor affinity for desired purposes. This is exemplified by the ability of the C10 series of anti-EGFR scFv antibodies to block EGFR signaling and inhibit cell growth in a manner that Rut.

Figure 7. (A) Absorbance change derived from dynode voltage during thermal scan of 20 μM scFv at 218 nm derived from data in Figure 5. (B–C) Dynamic light scattering of 10 μM of C10 and C10KV3_LV1DE at various temperatures. (B) Averaged static molecular weight calculated over increasing temperature. (C) % mass graphed as a function of hydrodynamic radius. Data represents 20 scans of 30 secs (for a total of 10 min) at the set temperature. C10 and C10KV3_LV1DE were calculated to be >99% monomeric at 25°C. The plots are representative of at least 2 independent experiments.
correlates with increased binding affinity. These types of data, developed across a large array of antibody/antigen pairs, have led to incorporation of affinity maturation as a critical step in therapeutic antibody development. As observed with C10 and P2224, altering the amino acid sequence of an antibody with the defined goal of improving binding affinity can affect other biophysical properties, such as stability, expression level, and propensity to aggregate, which are also critical for moving antibodies toward clinical development. Although formulation can be used to overcome poor biophysical characteristics (e.g., cetuximab’s propensity to aggregate upon mechanical manipulation was addressed by a late-stage change in formulation), it is sometimes insufficient on its own. Therefore, steps to either identify an alternative lead with similar biologic activity and better biophysical characteristics or protein engineering approaches to develop more stable, aggregation-resistant variants of the lead agent, are undertaken early in the development process.

As mentioned above, data from the Plickthun group demonstrating the superior production and stability of VH1-Vk3 scFv and the subsequent work by Tiller and colleagues expanding this finding to IgG production and stability led us to generate CDR-grafted Vk3 versions of C10 and P2224. Additionally, Honegger et al. examined to what extent the choice of “hydrophobic core” affects stability compared with the influence of the CDR/framework pairing. To this end, they constructed and evaluated scFv comprising one of 3 HuCAL VH3 variants that differed in their hydrophobic “lower cores,” but had unchanged CDRs, “upper cores,” and exteriors, paired with the same stable and unchanged HuCAL consensus Vk3 chain. The authors demonstrated that switching from a VH3-Vk1 (such as hu4D5–8) to a VH3-Vk3 dramatically improved expression levels as well as thermostability. Of most relevance here, the authors concluded that intrinsic stability of a framework pairing is insufficient on its own to fully stabilize an scFv, highlighting that fit between the CDRs and framework is a major component of overall stability. This finding is consistent with a large body of literature focused on humanization of murine antibodies for the purpose of decreasing immunogenicity of therapeutic candidates (for review see ref. 52).

Humanization, pioneered by Winter and colleagues, takes advantage of the conserved nature of the antibody frameworks that allow for grafting of the murine CDRs onto a human acceptor framework. Strategies to select a human acceptor framework include using a well-behaved “fixed framework” or using the human germline gene that is most closely related to the parent murine antibody. Implicit in the humanization process is the need for the CDRs to retain their conformation, and thus the ability to bind antigen. As exemplified by generation of trastuzumab from mu4D5, framework residues within the parental murine antibody are often required to correctly orient the CDRs. Straight grafting of mu4D5 CDRs onto consensus VH3 and Vk1 frameworks to generate hu4D5–1 resulted in an 80-fold loss of binding activity and concomitant loss of anti-proliferative activity. Creation of hu4D5–8 through back-mutation of a series of VH (amino acids 71, 73, 78, 93) and VL (amino acid 56) framework residues and 2 CDR residues (CDR-H3 residue 102 and CDR-L2 residue 55), identified via molecular modeling, improved antigen binding 250-fold over hu4D5–1 and restored anti-proliferative activity. A lambda to kappa framework switch has been attempted previously for an anti-GCN4 intra-body. Wörn et al. started with an anti-GCN4 antibody consisting of variants of IMGT mouse germline V-regions IGHV2–6–7*02 (88% identity) and IGLV1*01 (97% identity). To graft the
mouse anti-GCN4 CDRs onto Vk and VH frameworks, Wörn et al. created two mutants: one labeled κ-graft and one labeled λ-graft. The κ-graft was produced by replacing the VH and VL CDR sequences of another scFv (“hybrid”), consisting of the VL domain of hu4D5 (94% identical to mouse IGKV6–17*01) and the VH domain of A48+++H2 (93% identical to mouse IGHV4–1*02), with the anti-GCN4 CDRs. In addition, the κ-graft contained the d-e loop of the original mouse λ anti-GCN4 antibody and the six residues following CDR H2 plus 2 additional back mutations in the heavy chain at positions 71 and 78. Thus this design is significantly more complicated than our P2224KV3_LV1DE design, which changes only the light chain CDRs and d-e loop and does not change the heavy chain at all. The anti-GCN4 κ-graft was significantly more stable than the original anti-GCN4 antibody, but lost binding by 3 orders of magnitude.

The “λ-graft” of Wörn et al. differed from the κ-graft by only 7 amino acids at the VH/VL interface, and these changes were generated to aid in proper orientation of the domains, which successfully enhanced solubility, expression, and structural stability. However, the l-graft still lost an order of magnitude in KD from the anti-GCN4 mouse antibody, suggesting that other residues within the κ framework were important for maintaining CDR orientation, and thus antibody affinity. Our design is therefore both simpler and more effective than the approach of Wörn et al.

As mentioned above, we hypothesized that the only change required to facilitate the grafting of the Vλ3-like CDRs was the retention of the framework d-e loop, an “upper core” change that was within the range considered necessary by Plückthun et al. for successful grafting. We stabilized the upper core of C10KV3 and P2224KV3 by back-mutating the d-e loop to the λ-like sequence found in the original C10 to produce C10KV3_LV1DE and P2224KV3_LV1DE. In this process, we actually removed the bulky aromatic F71 from the upper core of the Vk3 to make room for the probable λ-like conformation of the L1 loop, including interactions of A71 with L28 and A33, and added back the side chain of residue 66 (Gly in κ3 antibodies; Lys in C10), which affects the conformation and position of the whole d-e loop. We demonstrate that these changes enhance thermostability and expression while maintaining affinity. Interestingly, the d-e loop of the native Vk3 version in C10KV3 and P2224KV3 decreased thermal stability and production yields, consistent with the proposed role in stabilizing the Vλ domain, but did not affect antigen binding. While Rosetta predicted that both κ3 variants might produce more stable antibodies than C10, it was not able to distinguish C10KV3_LV1DE from C10KV3 from an energetic standpoint, reinforcing the utility of careful structural analysis.

An increased aggregation propensity for an antibody results in reduced biological activity, increased clearance, and decreased safety, since aggregate formation can lead to aberrant immunogenic reactions. Notably, although we improved upon thermostability, the mode of aggregation between the parental and redesigned mutants drastically changed. Based on CD spectra and dynamic light scattering, C10 and P2224 demonstrate temperature-dependent irreversible unfolding with enhanced β structure, which may be caused by increased formation of large, soluble oligomers. However, the κ3 variants exhibit thermal unfolding that is more characteristic of the hu4D5 scFv (VH3/VK1), which demonstrates irreversible unfolding and results in aggregate precipitation.

The mechanisms governing the different unfolding properties of C10 and C10KV3 are unclear at this time. Since we observed a gain in β-signature in CD spectra, it is possible C10 and P2224 are generating soluble pre-fibrillar structures. This property is demonstrated by Bence-Jones proteins, which are over-produced light chain byproducts of multiple myeloma cells that cause light chain amyloidosis leading to insoluble amyloid fiber accumulation and vital organ failure. The specific germline genes that are over-represented in Bence-Jones proteins are predominantly of λ origin (λ1, λ2, λ3, λ4 and κ1). However, it should be noted that large soluble aggregates are just as toxic to cells as precipitated aggregates. Whether C10 and P2224 are generating soluble pre-fibrillar structures that are remedied by the κ3 swap requires further investigation.

Overall, we present here a novel method for switching a problematic VI framework with more stable k3 sequences. Based on structural analysis, we made rational decisions to back-mutate the flanking framework to the conserved λ residues and demonstrate that the swap created mutants with enhanced production from E. coli, reduced resistance to concentration, increased thermostability, and altered. We anticipate that the strategy we have employed will be useful in many design applications governing scFv and multispecific antibodies and provide a tool to enhance the biophysical properties of these potential therapeutics early in the engineering and selection process.

Materials and Methods

Homology modeling of C10 and designed antibodies

Sequence alignments of C10 with human germline and PDB sequences were performed with PSI-BLAST. C10’s CDRs were grafted into the sequence of human κ3, λ3, and λ1 germline sequences by replacing the CDRs with the C10 sequences, according to the CDR definitions of North et al.,24 which are longer than the standard Chothia definitions.40 In our definitions, CDR1 and CDR3 both begin immediately after the cysteines of the disulfide bond and end just before the tryptophan or phenylalanine motifs immediately across from the cysteine, typically W[VQR]Q after L1 and H1 and [WFG][QG]G after L3 and H3. Our L2 definition is 3 residues longer than the Chothia definition on the C-terminus and our L2 definition is 2 residues longer on each end of the CDR.

We built a homology model of the C10 scFv, consisting of a human IGHV1–69 domain and of a likely human IGLV3–19/ IGLV1–44 fusion product, and for the C10 VL-CDRs grafted onto an IGKV3–20 framework (referred to as C10KV3). The C10 mutations giving rise to P2224, which are predominantly located in the VH1 heavy chain, were added to the models of C10 and C10KV3 to produce models of P2224 and P2224KV3,
respectively. Seeking to minimize the number of templates used in our C10 homology model, templates for framework and CDRs were not selected sequentially, but were considered simultaneously because we aimed to select framework templates that already contain CDRs in the correct conformations. We focused on CDRs H1 and L1 first, in conjunction with the framework choice. CDRs L2 and H2 were modeled directly on their donor framework loops as these loops in many cases preserve a common backbone structure and only display sequence variability. Choices for H3 and L3 were made together as we found an antibody structure with high homologies in both loops that provided a good model for the L3-H3 interaction.

We chose PDB entry 2G75 as the C10 wild-type template because it consists of an IGHV1/IGLV3 variable domain pair like the C10 antibody. As a λ3/λ1 fusion, the Vλ3 chain still contains approximately 2/3 of the λ3 light chain, so 2G75 represents this antibody well. The CDRs of 2G75 represent the most likely conformations of H1 (H1–13–10), H2 (H2–10–1), L1 (L1–11–3), and L2 (L2–8–1).24,29 The C10 model based on 2G75 requires grafting the L3-H3 coordinates from another structure, since the CDR3 lengths in 2G75 are not the same as those in C10. We chose PDB entry 2FB4,42 an IGHV3–30/IGLV1–44 framework, for both grafts. The grafted 2FB4 L3 backbone was left unchanged, while the grafted 2FB4 H3 was in part remodeled using the UCSF Chimera package and its interface to Modeler.63 In the partial remodeling of the H3, we preserved the disulfide-bonded loop in 2FB4 (sequence CSSASC) to model the same segment in C10 (sequence CSSTSC) and most of the interface with L3.

The H1, L1, H3, and L3 CDRs from the C10 model were grafted onto the structure of an IGHV1–69/IGKV3–20 antibody (PDB 1RHH) in order to model C10KV3. The acceptor H2 (H2–10–1) and L2 (L2–8–1) backbones were left unchanged because they belong to identical clusters for parent and acceptor frameworks, respectively. The side-chain packing of all models was optimized using the program SCWRL4.

We also modeled C10 and the C10LV1, C10LV3, and C10KV3 variants with the RosettaAntibody webserver65 with standard parameters but without the option of extensive modeling of H3. A model of C10KV3_LV1DE was constructed by manually grafting the d-e loop from PDB entry 4GXV with the same sequence as the d-e loop of C10 (KGSRTSA). Structures were analyzed and examined in Pymol (Schrödinger, Inc.). Clusters of the CDRs in the models were determined with our PyIgClassify website.29

The RosettaAntibody models were refined in a 2-stage procedure using the program Rosetta (version 3.6, release May 11, 2015). The first step was to idealize the bond lengths and bond angles of all of the models with the command line:

idealize.linuxgccrelease -s filename.pdb

The second step was to generate 250 decoys by running the mixed dihedral-angle/Cartesian-coordinate minimization protocol referred to as "dual-space relax,"36 starting from the idealized structures. The protocol runs 3 cycles of "fast relax" in dihedral angle space and 2 cycles of Cartesian minimization, which allows bond angles and bond lengths to change. The FastRelax protocol consists of 5 rounds of the following: multiplying the repulsive van der Waals parameters by a scale factor C (0 < C ≤ 1), several rounds of replacement of all side chains with random rotamers from our library66 with Metropolis criterion acceptance, and then continuous energy minimization of the backbone and side chains. The factor C is ramped up from 0.02 to 1.0 in each round. The lowest energy structure when C=1 is saved as a decoy, and passed to the next cycle of minimization. The dual-space relax command line is given here:

relax.linuxgccrelease -dual_space -non_ideal -shapovalov_lib_fixes_enable -nstruct 250 -s filename.pdb

The flag "-shapovalov_lib_fixes_enable" instructs Rosetta to use updated Ramachandran scoring functions (rama and P_aap), which we have recently developed based on kernel density estimates of the backbone conformations of the 20 amino acids.57

Cloning, expression, and purification of scFvs

The C10 and P2224 scFv genes, kindly provided by Dr. James Marks (University of California San Francisco), were previously described19,20,39 and the C10KV3_LV1DE coding sequence was constructed by gene synthesis. All three gene were cloned into the pSyn2 bacterial expression plasmid.66 To create P2224KV3_LV1DE, the P2224 affinity mutations were introduced into C10KV3_LV1DE by site-directed mutagenesis in a 2-step process using the Quikchange II site-directed mutagenesis kit and the following primers: FWDF P24 G113A 1RHH, REV P24 GI13A 1RHH, REV P22 ST GI 1RHH, and REV P22 ST GI 1RHH (Supplementary Table 2). The kappa DE-loop back-graft mutations (to generate C10KV3 and P2224KV3 were generated by conventional site-directed mutagenesis (Quikchange II) using the following primers: FWDF C10k3_LV1DE and REV C10k3_deGDF. (Supplementary Table 2).

For protein production, TG-1 cells were transformed with sequence-verified clones of pSyn2 scFv-expressing constructs and induced to produce soluble scFv under conditions of osmotic pressure as previously reported.67 In short, bacterial cultures grown to OD600 = 0.8 in 2 L flasks, were pelleted and resuspended in 0.5 L of 2XYT media containing 0.5 mM IPTG, 0.4 M sucrose, and 100 mg/ml carbenicillin and cultured for 16 hr at 25°C or 30°C, as appropriate. Following dialysis into Dulbecco’s phosphate-buffered saline (PBB), the soluble scFv fraction was recovered from the culture supernatant by IMAC affinity chromatography as previously described.68 Initial preparations were further analyzed by size-exclusion chromatography over a HiPrep 16/60 Sephacryl 100 column at a flow rate of 1 ml/min using an AKTA Prime+ (GE Healthcare). Resulting chromatographs were compared to known protein standards as previously described69 and deemed predominantly monomeric. Following IMAC purification, the preps were confirmed to be > 99% monomeric by dynamic light scattering (described below).
Circular dichroism, turbidity, and Tm measurements

Circular dichroism (CD) spectra, thermal unfolding curves, and turbidity (dynece voltage) were recorded using an Aviv 62DS spectropolarimeter using thermostatted cells with an optical path length of 1 mm. Ellipticity readings were time averaged for up to 10 sec at 60 points/s. The bandwidth was set to 2 nm and the dynece voltage was initially kept below 400 V at 25°C. CD spectra over the range from 195 to 250 nm were recorded on indicated scFv in 20 mM potassium phosphate at neutral pH.

To calculate derivative Tm by differential scanning fluorimetry, each scFv was analyzed at a concentration of 0.1 mg/mL in a 20 ml volume plated in quadruplicate in MicroAmp fast optical 96-well titer plates using the Protein Thermal Shift™ (Applied Biosystems) assay. PBS was used as a baseline control. Data were generated on an Applied Biosystems® 7500 Real-Time PCR System using continuous ramp mode at 0.5% ramp rate from 25°C through 95°C. Data were analyzed using the Protein Thermal Shift™ Software.

Dynamic light scattering

Dynamic light scattering experiments were performed on a DynaPro Molecular Sizing Instrument with Dynamics V6 data analysis software (Protein Solutions, Inc.) on 80 µl of a 10 µM solution of purified scFv in PBS. Autocorrelation curves were acquired for a total acquisition time of 600 s at each temperature.

Flow cytometric analysis

A431-NS cells (ATCC #CRL-2592) were grown to sub-confluence and harvested in Ca2+/Mg2+ free PBS containing 1 mM EDTA. Cells were washed, resuspended in FACS buffer (1% BSA, 0.1% sodium azide, PBS), and plated at 2 x 105 cells/well in 96-well round-bottom tissue culture plates. To evaluate scFv binding to the cells under conditions of equilibrium, the reaction took place overnight at 4°C. The next day cells were washed and fixed with 1% paraformaldehyde, and cell-bound scFv was detected with 1 mg/ml FITC-labeled Penta-His antibody (Qiagen) for 1 hour on ice. Cells were washed, fixed again, and analyzed. To determine whether P2224 or P2224KV3_LV1DE blocked cetuximab binding, cells were pre-bound with 3 µM of the indicated scFv overnight at 4°C. The next day, cetuximab was pre-labeled with equimolar concentrations of FITC-conjugated donkey anti-human IgG (Jackson ImmunoResearch) for 30 min on ice. The cetuximab:FITC conjugate was applied to the scFv pre-blocked cells for 1 hour on ice. Cells were washed again and resuspended in 1% paraformaldehyde in PBS. All fluorescently labeled cells were acquired using a FACSScan instrument (Becton Dickinson) and the data were analyzed by FlowJo software (FlowJo, LLC).

Disclosure of Potential Conflicts of Interest

No potential conflicts of interest were disclosed.

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Supplemental Material

Supplemental data for this article can be accessed on the publisher’s website.

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