Improving computational protein design by using structure-derived sequence profile

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Abstract
Designing a protein sequence that will fold into a predefined structure is of both practical and fundamental interest. Many successful, computational designs in the last decade resulted from improved understanding of hydrophobic and polar interactions between side chains of amino acid residues in stabilizing protein tertiary structures. However, the coupling between main-chain backbone structure and local sequence has yet to be fully addressed. Here, we attempt to account for such coupling by using a sequence profile derived from the sequences of five residue fragments in a fragment library that are structurally matched to the five-residue segments contained in a target structure. We further introduced a term to reduce low complexity regions of designed sequences. These two terms together with optimized reference states for amino-acid residues were implemented in the RosettaDesign program. The new method, called RosettaDesign-SR, makes a 12% increase (from 34 to 46%) in fraction of proteins whose designed sequences are more than 35% identical to wild-type sequences. Meanwhile, it reduces 8% (from 22% to 14%) to the number of designed sequences that are not homologous to any known protein sequences according to psi-blast. More importantly, the sequences designed by RosettaDesign-SR have 2–3% more polar residues at the surface and core regions of proteins and these surface and core polar residues have about 4% higher sequence identity to wild-type sequences than by RosettaDesign. Thus, the proteins designed by RosettaDesign-SR should be less likely to aggregate and more likely to have unique structures due to more specific polar interactions.

Keywords
protein design; structure-derived sequence profile; RosettaDesign; local sequence-structure coupling

INTRODUCTION
Computational protein design attempts to identify the amino acid sequences that will fold into a given target structure with sufficient stability and solubility and, ultimately, perform a desired function by using computational methods. The research in this area is motivated by the practical need for improving, modifying, and/or expanding the function of natural
proteins as well as the fundamental interest in understanding the relationship between protein sequences, structures, and functions. Significant progress has been made in the last two decades in de novo design of proteins. In 1997, Dahiya and Mayo demonstrated through NMR structure determination that a computationally designed sequence can fold into a unique structure. Since then, designed proteins have been experimentally validated to fold into all-alpha, all-beta, and mixed α-beta proteins. A protein with a new protein fold was also designed successfully. The advance in computational design leads to a wide variety of applications ranging from increasing protein stability and folding rate, improving fold recognition, identifying functional sites, enhancing or creating binding specificity, and catalytic activity to designing protein conformational switches.

Despite of significant progresses, we have not yet reached the point that successful design is guaranteed and protein design can be routinely applied to any given (known or novel) structure folds. For example, in a large-scale design of nine proteins by RosettaDesign, only “half of the folded designs have NMR spectra and temperature melts typical of tightly packed proteins.” The actual success rate of a fully automatic method is likely even lower because manual inspection and human experience are often involved in selecting designed sequences for final experimental validations. More recently, an experimental testing of five computationally designed, high-affinity ligand-binding proteins revealed that none of the designed proteins performs as expected due to instability, aggregation, or lack of detectable, designed ligand binding.

What causes the low success rate described above? A commonly used criterion for judging the potential success of a design is sequence identity between the designed sequence and the original wild-type sequence for the target structure. The existing protein-design techniques have been reported to reach an average sequence identity of 30–37%. However, most studies are limited to a small number of test proteins while fixing certain residue types (e.g., Gly, Cys, Pro in EGAD, and Cys in RosettaDesign). In fact, as we shall see in this article, RosettaDesign, EGAD, and Liang-Grishin all achieved an average sequence identity of 30–32% with a set of 33 globular proteins. It is known that 30–32% sequence identity is in a so-called twilight zone where two proteins may or may not have the same structure. This twilight zone is confirmed in Figure 1 by making 22,207,780 pairwise structural comparisons by TM-align between 6665 proteins (2Å resolution or better, pairwise sequence identity of 90% or less, chain length between 60 and 300 residues, from the pisces server). There is a transition from structurally dissimilar pairs (TM-score around 0.3) to structurally similar pairs (TM-score >0.5) at sequence identity around 30%. Here, TM-score >0.5 denotes structurally similar pairs because at TM-score >0.5 the difference between the two structures is often limited to a small portion of the structures, for proteins with high sequence identity, in particular. That is, locating in the twilight zone is likely one of the reasons for low success rate. On the other hand, Figure 1 shows that structurally dissimilar pairs essentially disappear when sequence identity is above 35%. Thus, even small improvement from 30 to 32% sequence identity achieved by existing methods could potentially lead to a large improvement in success rate. Indeed, a recent study showed that optimizing sequence identity (2% improvement) and specific reference states for all-beta proteins lead to the first successful design of a beta sandwich protein called Tenascin without the need of negative design.

Improving sequence identity to wild type sequences requires a more effective scoring function because sequences are predicted by placing them onto a target structure and minimizing the scoring function. Some scoring functions are made of various physical energy terms, including van der Waals interaction, solvation, electrostatics, and hydrogen bonding. Others, on the other hand, took the advantage of the knowledge-based
information derived from known protein structures. These energy functions are usually built to capture tight and specific tertiary packing interactions. The relation between local sequence and backbone (or secondary) structure, however, is only described by a single-residue property of amino acid propensity for a given backbone torsion angle (as used in RosettaDesign) or for a given secondary structure (as used in ORBIT). However, it is known from secondary structure or backbone torsion-angle prediction that a local sequence segment of about 20 residues long is needed to determine about 80% three-state secondary structure or 83% of both backbone $\phi$ and $\psi$ torsion angles within 60 degree from their native values. Thus, it is necessary to go beyond single-residue properties to account for the coupling between local structure and sequence for protein design.

One way to take into account the coupling between local structure and sequence is to search structurally similar fragments for a given target fragment structure and produce structure-derived sequence profiles from the sequences of structurally similar fragments. We have successfully applied this structure-derived sequence profile for improving fold recognition and protein structure prediction. In this work, we explore the usefulness of this structure-derived sequence profile as an energetic term in protein design. We implemented this energy term into two separate protein-design programs, RosettaDesign and EGAD. We found that implementing this energetic term in RosettaDesign leads to about 3% improvement from 31 to 34% for the sequence identity between the top-1 energy-ranked sequence and the wild-type sequence (based on design of 944 proteins). Larger improvement (8%) is also observed when the sequence profile is implemented in EGAD. We further show that structure-derived sequence profiles together with energy terms for reducing repetitive regions and optimized reference states lead to a method (called RosettaDesign-SR) that increases 12% (12%) for the fraction of designed proteins with more than 35% (30%) sequence identity to wild-type sequences and makes a 8% reduction to the fraction of designed proteins that do not have homologous hits to any existing sequences according to psi-blast. Moreover, sequences designed by RosettaDesign-SR contain more polar residues than those by RosettaDesign at both surface and core regions of proteins and, thus, are potentially more soluble and likely fold into unique conformations.

**THEORY**

**Datasets for protein design**

We downloaded a nonredundant protein set with resolution better than 2.0Å, pair-wise sequence identity of less than 30% from the pisces server (October 17, 2008). This set contains 4803 protein chains that are further reduced to 2528 chains after removing the chains with missing residues or backbone atoms. We randomly selected 40 globular proteins between 60 and 150 residues as a set for parameter training. Because RosettaDesign failed to design sequences for 7 of the 40 proteins, we finally have 33 proteins as the training set of protein design programs. The number of residues of these proteins ranges from 66 to 132, with an average of 104. We use a small set of 33 proteins to reduce the computational time for training.

We further selected the first 1000 smallest proteins from the above 2528 proteins. Because RosettaDesign program failed to design 56 of the 1000 proteins, the total number of proteins for testing design programs is 944. The number of residues of these proteins range from 60 to 160, with an average of 112.4.

Finally, we used the fragments contained in the structures of 2495 (2528–33) protein chains to build the fragment template library. All datasets used in this study are available at http://sparks.informatics.iupui.edu.
Structure-derived sequence profile

The method for generating structural-based sequence profile is essentially the same as that developed by Zhou and Zhou. Given the structure of a protein to be designed (target structure), we first divided it into a number of structural fragments with a sliding widow of length $L_\text{frag}$ along the sequence. For a protein with $N_{\text{res}}$ residues, there are $N_{\text{res}} - L_\text{frag} + 1$ structural fragments. Each target fragment was then structurally aligned to the same-length fragments contained in a fragment library. The sequences of the top $N_{\text{frag}}$ library fragments with the lowest root-mean-squared distance (RMSD) to the target fragment are used to calculate sequence profile, or, the probability of a residue type in a given target sequence position. RMSD values between fragments are calculated based on four backbone heavy atoms (N, C$_\alpha$, C, and O) plus pseudo C$_\beta$ atom because the locations and types of native residues are supposed to be unknown. Pseudo C$_\beta$ atoms are generated based on standard 1.54Å for the C$_\alpha$-C$_\beta$ bond length, 109.53° for the bond angle N-C$_\alpha$-C$_\beta$ and 109.46° for the dihedral angle C-N-C$_\alpha$-C$_\beta$. The profile $[P_{\text{profile}}(i,I) | i, I = 1..20]$ is obtained by averaging over the sliding windows. We optimized $L_{\text{frag}}$ and $N_{\text{frag}}$ to maximize the sequence identity between the wild-type sequence and the consensus sequence that corresponds to the residue type with the highest probability at each sequence position in the profile. The optimal values for $L_{\text{frag}}$ and $N_{\text{frag}}$ are 5 and 25, respectively, by using the above mentioned fragment library of 2495 proteins. During optimizing $L_{\text{frag}}$ and $N_{\text{frag}}$, the sequence profile of a given protein in the 2495 protein set is generated from the fragment library of the rest 2494 proteins. The average sequence identity between the consensus sequences and wild-type sequences is 24.9% averaged over 2495 proteins. This is 3% higher than ~22% in our previous work by residue-depth dependent fragment comparison with $L_{\text{frag}} = 9$ and $N_{\text{frag}} = 25$. Here, residue depth refers to the depth of a residue from the surface of a protein and was used to compare fragments in similar level of solvent exposure. We did not use residue-depth here because using residue depth did not lead to further improvement in sequence identity. A shorter five-residue fragment used here is closely related to the jump between helical and strand structures.

Finally, structure-derived sequence profiles for the 33 protein set (training set) are generated from the fragments of 2495 proteins.

RosettaDesign, EGAD, and Liang-Grishin programs for protein design

RosettaDesign (versions 2.3 and 3.0) and EGAD were download from http://www.rosettacommons.org/ and http://egad.ucsd.edu/EGAD_manual/index.html, respectively. We also obtained a copy of protein design program by Liang and Grishin. In RosettaDesign 2.3, the command “Rosetta -design -fixbb -ex1” was executed, while in RosettaDesign 3.0, the command “fixbb.linuxgccrelease -ex1” was executed. In EGAD, the JOB-TYPE is set to “MC” and the VARIABLE_POSITIONS is set to “total_sequence_design.” In Liang’s program, its default setting was used. We have modified RosettaDesign 2.3, rather than RosettaDesign 3.0 because the latter was not yet available at the time we started this study. After RosettaDesign 3.0 was released, we found that the difference between sequences designed by RosettaDesign 3.0 and RosettaDesign 2.3 is small and thus we continued our focus on modification of RosettaDesign 2.3.

Modification of RosettaDesign 2.3 and EGAD energy functions

We have modified the energy functions in RosettaDesign as follows:

$$\Delta E = E - E_{\text{original}} = - w_{\text{profile}} \Sigma \ln P_{\text{profile}}(i, I) + w_{\text{rep}} \Sigma \ln N_{\text{rep}} + \Sigma r E_{\text{ref}}(s, I)$$

(1)
where $E_{\text{original}}$ is the original energy function of a given program, $P_{\text{profile}}(i, I_i)$ is the structure-derived sequence profile at a given sequence position $i$ and residue type $I_i$ at the position ($1/25$ is added to each value of $P_{\text{profile}}(i, I_i)$ to prevent ln0), $N_{\text{rep}}^i$ is the number of nearest and second nearest neighboring residues that repeat the residue type at the sequence position $i$ (ranging from $i - 2$ to $i + 2$ including itself), $E_{\text{ref}}(s_i, I_i)$ is the reference-state energy term that depends on secondary structure ($s_i$) and residue type ($I_i$), and $w_{\text{profile}}$ and $w_{\text{rep}}$ are weight parameters for the profile and residue-repeat terms, respectively. Here, secondary structure assignment was made by STRIDE. The purpose of the first term is to incorporate restraints on sequence from local structure. We added the second repetitive energy term because adding the sequence profile term leads to a significant increase in number of repeats of same residue types such as LLL and VVV. The second term is a simplified definition of sequence complexity that measures the extent of randomness in a sequence by Shannon's entropy. It leads to a higher (unfavorable) energy if the designed sequence repeats a certain residue type within a local region. We found that this term is effective for removing low-complexity regions of designed sequence defined by the SEG program. We added this term because low sequence complexity is often associated with disordered regions of proteins. The last term in Eq. (1) is for optimizing the reference states of residue types at different secondary structure environments so that the occurrence of each residue type is close to that of wild-type sequences. Secondary structure-dependent reference states were introduced previously.

Equation (1) is also implemented in EGAD. However, only the first term is used because the purpose is to confirm the effect of structure-derived sequence profile in protein design.

RESULTS

Sequence identity to wild type sequences

To obtain consistent, comparative information on the sequence identity of designed sequences to wild type sequences, we have applied EGAD, RosettaDesign 2.3, RosettaDesign 3.0, and Liang-Grishin to 33 training proteins. We found that the average sequence identity of 100 designed sequences per protein (a total of 3300 sequences) are 30.3% by EGAD (fixing Cys, Gly and Pro), 31.9% by Rosetta 2.3, and 29.0% by Rosetta 3.0, and 30.0% by Liang-Grishin, respectively. Thus, the average sequence identities given by different programs are between 29 and 32% for this set of training proteins.

Modification of RosettaDesign 2.3

The weight parameters and reference values are trained with the 33-protein set. Side chains of each protein structure are stripped off and pseudo C$\beta$ atoms are built before the structure is used as the target structure. That is, all amino acid residues are designed including Cys which is typically fixed by RosettaDesign. Results shown in Table I are averaged over 100 designed sequences per protein. Compared with RosettaDesign, it is clear that structure-derived sequence profiles alone has substantially lower sequence identity (8% lower) with significantly higher low complexity regions (9%). We first implemented only the structure-derived sequence profile [the first term in Eq. (1)] into RosettaDesign (called RosettaDesign-S for convenience) and optimized $w_{\text{profile}}$ based on the average sequence identity of designed sequences to the wild type sequences. Figure 2 shows that sequence identity changes from 31.9% at $w_{\text{profile}} = 0$ to 35.2% at $w_{\text{profile}} = 1$ with a peak at 35.6% at $w_{\text{profile}} = 0.75$. The change of sequence identity is small around $w_{\text{profile}} = 0.75$, suggesting the stability of improvement. Without fixing Cys, the RosettaDesign modified by incorporating structure-derived sequence profiles does not produce any sequences containing Cys residues as the original RosettaDesign does.
Sequences designed by RosettaDesign-S, however, have frequent appearance of multiple repeats of same residue types such as LLL (three sequentially connected Leu residues) and VVV (three Val residues). This significant increase in LLL and VVV appearance occurs along with significant increase in low-complexity regions. As shown in Table I, low complexity regions defined by program SEG\textsuperscript{72} increase from 6.4\% by RosettaDesign to 21.3\% by RosettaDesign-S, compared to only 2.1\% for wild type sequences. These repetitive regions, signaled low complexity of designed sequences, are often associated with disordered regions.\textsuperscript{73} The occurrence of low complexity regions indicates the hidden bias of local-fragment-derived sequence profiles toward small hydrophobic residues such as L and V. Indeed, we observed a large drop in fraction of polar residues from 44\% in RosettaDesign to 37\% by RosettaDesign-S. The reduction is the most significant in helical regions (decreases 11\% from 48\% to 37\%). Thus, we introduced two additional terms in Eq. (1) to increase sequence complexity and improve the balance of hydrophobic and hydrophilic residues. More specifically, weight parameter $w_{\text{rep}}$ and reference state energy $E_{\text{ref}}$ are optimized for reducing repetitive sequences and increasing the fraction of hydrophilic residues. These parameters are optimized by a trial-and-error approach. We only modified $E_{\text{ref}}$ for eight residue types that have a significant deviation from their occurrence frequencies of native proteins. The final optimized value for $w_{\text{rep}}$ is 0.4 and the values for $E_{\text{ref}}$ are listed in Table II with $w_{\text{profile}}$ unchanged at 0.75. The final version of the method is labeled as RosettaDesign-SR with R for repetitive and reference energy terms.

The effect of repetitive and reference-state energy terms is illustrated in Table I. Although sequence identity has decreased by about 1\% from RosettaDesign-S, the fraction of low complexity regions and the occurrence of LLL and VVV as well as the frequency of polar residues are now more in line with wild-type sequences. More importantly, the amount of polar residues in designed sequences given by RosettaDesign-SR is significantly closer to that of wild type sequences than RosettaDesign. For example, the fraction of polar residues in strand regions are 34\% by RosettaDesign, 28\% by RosettaDesign-S, and 40\% by RosettaDesign-SR, compared with 41\% for wild-type. The fraction of polar residues is important for maintaining the solubility of proteins, preventing aggregation, and enhancing specific interactions.

**RosettaDesign-SR**

RosettaDesign-SR is tested with the dataset of 944 proteins. We designed 100 sequences for each protein. These sequences are ranked by the modified energy in Eq (1). The average sequence identities to wild type sequences for the final version of the modified RosettaDesign are 33.9\% for top 1 ranked by energy, 36.8\% for the highest sequence ID among top 10, 39.0\% for the highest sequence ID in 100 designed sequences and 33.2\% for the average over 100 designed sequences. They are 2.5\%, 2.4\%, 2.3\%, and 2.5\% improvement over RosettaDesign, respectively. Because we found that averaging top 1, or the best in top 10, or 100 designed sequences over 944 proteins yield similar trends in comparison with Rosetta-Design, we will report the results based on statistics of top 1, here and hereafter.

Table III compares the results given by wild-type sequences, sequences designed by RosettaDesign 2.3, and by RosettaDesign-SR. Similar to the results for the training set, RosettaDesign-SR provides a consistent improvement in sequence identity to wild-type sequences (2.5\%), in amount of low complexity regions (5.1\%), and in fraction of polar residues (2.6\%). The sequence identity to wild-type sequences increases in a similar magnitude across different secondary structure. RosettaDesign-SR makes a 2.7\% (from 26.2\% to 28.9\%), 1.6\% (from 35.6\% to 37.2\%), and 2.7\% (from 35.0\% to 37.7\%) improvement over RosettaDesign for helical, strand, and coil regions, respectively (not shown in Table III for clarity). As in RosettaDesign, helical residues from RosettaDesign-
SR have the lowest sequence identities to the wild type sequences. The improvement in fraction of polar residues, on the other hand, is the most significant for strand regions. The sequences from RosettaDesign-SR contain 5% more polar residues in strands than the sequences from RosettaDesign although still 3% less than wild-type sequences.

Figure 3 compares the fraction of all 20 amino acid types in designed sequences to those in wild type sequences. The frequencies of most residue types given by RosettaDesign-SR are slightly closer than those given by RosettaDesign to the corresponding frequencies in the wild type sequences. The average absolute deviation between each amino acid frequency in designed sequences and wild-type sequences decreases from 1.1% (RosettaDesign) to 0.9% (RosettaDesign-SR).

It is of interest to know how the matching residues between designed sequences and wild type sequences are distributed on the surface or inside proteins and what fraction of polar residues are on the surface of proteins. We measure solvent exposure by residue contact number, the number of residues (C$_\alpha$ atoms) within 10 Å from a specific residue. We exclude six nearest sequentially linked residues ($i \pm 1$, $i \pm 2$, and $i \pm 3$) because they are more likely within 10Å distance cutoff. Contact numbers are divided into four bins (0–5, 6–10, 11–15 and >16 contacts). We use residue contact numbers because they are found evolutionarily more conserved than solvent accessibility.$^{75,76}$ More importantly, residue contact number based on C$_\alpha$ atoms can be evaluated in the absence of sidechains.

Table IV compares sequence identity at different contact numbers. RosettaDesign-SR improves sequence identity over RosettaDesign by 3% at the surface (residue contact number <5) and 0.5% at the deep core (residue contact number >16). This is because high sequence identity inside the core (50%) is more difficult to improve further. Identical residues between wild type sequences and designed sequences are further analyzed according to the locations of polar and nonpolar residues in wild-type sequences. As Table IV indicates, the most significant improvement is in sequence identity of polar residues across different levels of solvent exposure. There are 2.9%–4.5% more polar residues in RosettaDesign-SR sequences that are identical to those in wild type sequences than in RosettaDesign from deep core to surface. The smaller but consistent improvement is also observed in sequence identity between nonpolar residues of designed and those of wild-type sequences (between 0.2 and 1.6% at different contact numbers).

Improvement in the polar content of designed sequences is also reflected from the fraction of polar residues at different contact numbers. As shown in Table IV, sequences designed by RosettaDesign-SR have more polar residues than those by RosettaDesign and closer in the fraction of polar residues to wild type sequences at all regions of proteins except that our new method yields 1% more polar residues than wild type sequences on the surface of proteins (residue contact numbers <5).

Because the sequence identity to wild type sequences is the lowest for surface-exposed residues that have few contacts with other residues, we suspect that higher fraction of surface residues in a protein (less globular-shaped) would lead to smaller sequence identity between designed sequences and wild type sequences. Figure 4 shows that such a trend indeed exists. When the fraction of surface residues (contact number <9) is greater than 80%, all designed proteins have less than 30% sequence identity to their respective wild type sequences. Meanwhile, when the fraction of surface residues is less than 30%, all designed proteins have more than 20% sequence identity to their respective wild type sequences.

We further examine how sequence identity and fraction of polar residues change for different structural folds. Results are shown in Table V. As in RosettaDesign, sequence identity to wild type sequences given by RosettaDesign-SR is the highest for all-β proteins.
(37%). On the other hand, the fraction of polar residues given by RosettaDesign is 7.4% below wild type sequences for all-β proteins. RosettaDesign-SR cuts this gap more than half to 3%. Improvement on other β-strand containing proteins (others) is also observed. This suggests that the sequences designed by RosettaDesign-SR for β-strand containing proteins are more likely soluble than the sequences designed by RosettaDesign.

The above results appear to suggest a small incremental change from RosettaDesign to RosettaDesign-SR. However, as Figure 1 suggests, what is more important is the fraction of proteins whose designed sequences are more than 35% sequence identity to wild type sequences. We found that this fraction increases by 12% (33.9% by RosettaDesign and 45.9% by RosettaDesign-SR, respectively). If a 30% sequence identity cutoff is used, there is also a 12% increase from 60.1% by RosettaDesign to 72.1% by RosettaDesign-SR. Underlying significant improvement is also revealed by the fact the average sequence identity between sequences designed by Rosetta-Design and those by RosettaDesign-SR is only 51%.

Another way to verify the designed sequence is to perform a psi-blast search and to locate if there is any sequence homologous to designed ones. Because psi-blast search performs a local alignment, we define that a homologous sequence is found only if its length is at least 2/3 that of the designed sequence with an e-value of 0.01 or less. Figure 5 shows the highest sequence identity found in psi-blast search as a function of the sequence identity to wild type sequences. For the majority of proteins, the designed sequences have slightly higher sequence identity to the homologous sequences found in sequence library than to the wild type sequences. The average sequence identity between designed sequences and the closest homologous sequences is 29.9% by RosettaDesign and 34.3% by RosettaDesign-SR. (34.3% is only 0.4% higher than 33.9%, the average sequence identity to wild type sequences. This small change is due to many sequences having no homologous hits. See below.) This is consistent with about 3% improvement for sequence identity to wild-type sequences. This small improvement in average sequence identity, however, leads a significantly larger 15% increase in the fraction of designed proteins that share more than 35% identity with nearest homologous sequences in sequence library. The fractions of proteins are 54.6% by RosettaDesign and 70.0% by RosettaDesign-SR, respectively. For a 30% identity cutoff, the fractions of proteins are 74.3% by RosettaDesign and 84.5% by RosettaDesign-SR.

We further found that there is significant improvement in number of designed sequences without homologous hits in sequence library. There are 22% (or 209 proteins) without hits for 944 proteins designed by RosettaDesign. This number is reduced to only 14% (or 134 proteins) for proteins designed by RosettaDesign-SR. Most of these “orphan” sequences are lower than 30% sequence identity to their corresponding wild-type sequences and connected to the structures with high percentage surface residues as shown in Figure 4. That is, sequences designed for proteins with 60% or more surface residues (defined as contact number <9) are often unreliable.

**EGAD**

To further demonstrate the usefulness of structure-derived sequence profile for protein design, we also implemented the profile in EGAD as in Eq. (1) but without repetitive and reference energy terms because optimizing the last two terms makes minor changes to sequence identity. We found that there is an 8.5% improvement in sequence identity from 30.3% to 37.8% for 33 training proteins after introduction of the sequence profile term with \( w_{\text{profile}} = 7.0 \). The higher sequence identity is due to fixing of Gly, Pro, and Cys in EGAD. Because EGAD is too slow for testing on all 944 proteins, we randomly selected 55 proteins from the set. Application of the modified EGAD to the set of 55 proteins continues to make an 8.0% improvement in sequence identity over EGAD. Here, EGAD was run without
including native rotamers. This result confirms that the structure-derived sequence profile for protein design is effective for improving protein design.

**DISCUSSION**

In this article, we have tested whether a structure-derived sequence profile is useful for protein design. We found that its implementation in RosettaDesign and EGAD (RosettaDesign-S and EGAD-S) leads to about 4 and 8% improvement in sequence identity to wild type sequences, respectively. Further refining RosettaDesign-S with repetitive and reference-state energy terms leads to RosettaDesign-SR. On the basis of analysis of designed sequences for 944 proteins, we found that the most significant improvement of RosettaDesign-SR over RosettaDesign is in amount and location of polar residues in designed sequences. The fraction of polar residues increases by 3% (5% for strand residues) while sequence identity between polar residues of designed and that of wild-type sequences improves by 4% crossing all levels of surface exposure. More polar residues on the surface of the proteins for the sequences designed by RosettaDesign-SR will make them less likely to aggregate, a significant problem in designed sequences. Meanwhile, more polar residues inside the proteins for the sequences designed by RosettaDesign-SR will make them more likely to fold into a unique structure as hydrophilic residues form specific hydrogen bonding and salt bridges. This small improvement in sequence identity leads to a large change in overall designed sequences (the average sequence identity between sequences designed by RosettaDesign and those by RosettaDesign-SR is only 51%) and 12% increase in the fraction of proteins with more than 35% sequence identity to wild-type sequences.

Our structure-derived sequence profiles are obtained with a library of proteins with 30% or less sequence identity between each other. It is of interest to know if this sequence identity cutoff would have strong influence in the quality of sequence profiles. To answer this question, we obtained an updated dataset of 2592 proteins with a cutoff of 25% sequence identity (high-resolution X-ray structures, 2.0Å or better) by PISCES server. This dataset is larger than the set of 2495 proteins at 30% cutoff because this new set was downloaded 9 months later. We found that the sequence identity between the most popular residue types and wild-type sequences (averaged over proteins) is 25.3% for the new fragment library, compared to 24.9% for the library of 2495 proteins. Thus, homologous sequences if existing in proteins based on the 30% sequence identity cutoff have minor effect on structure-derived sequence profiles.

Implementing structure-derived sequence profiles only leads to about 3% improvement in sequence identity to wild type sequences. What has stopped us to achieve a higher sequence identity? Certainly, a protein can mutate frequently without a significant change in its three-dimensional structure. In fact, the average sequence identity is 56% (averaged over 944 proteins) between the wild type sequences and the consensus sequences from the sequence profile derived from homologous sequence alignments by psi-blast. That is, there is room for further improvement. A close examination of 100 sequences designed by either RosettaDesign or RosettaDesign-SR indicates that they are highly homologous among each other: the average sequence identity within themselves is 67.8% by RosettaDesign and 78.3% by RosettaDesign-SR. That is, the designed sequences converge to a single family of sequences (single solution) with different initial guesses. This suggests that it is not insufficient sampling of sequence space but rather the lack of a good energy function causing wild-type sequences often not within the family of designed sequences. Indeed, if we include a biased energy term to sample sequences with high sequence identity (SeqID) to wild type sequences \(E = -w_{\text{seq}}|\text{SeqID} - \text{SeqID}_0|\), we found that the energy of a designed sequence increases significantly in a parabolic curve as SeqID either increases toward to...
100% or decreases toward 0%. Moreover, the energy of a wild type sequence is close to the energies of random sequences with about 0% sequence identity to wild-type sequence. That is, the energy function does not favor sequences that are close to 100% wild-type sequence as one of the solutions. Although native sequences are not necessarily optimized for their structures, the energy difference between wild-type sequences and optimized designed sequences is too large to be realistic. In other words, the quality of an energy function remains as the bottleneck for successful, computational design.

One important observation is higher recovery of native polar residue types in protein cores by RosettaDesign-SR. All polar residues inside core (defined as having more than 10 contacting residues) are highly conserved with an average conservation index of 0.80 from PSI-BLAST (1 for 100% conservation). RosettaDesign-SR recovered a subset (23%) of core polar residues with an average conservation index of 0.91, compared to 20% by RosettaDesign with an average conservation index of 0.93. Analysis of correctly predicted pairs of polar residues indicates that RosettaDesign-SR consistently has about 40% more pairs than RosettaDesign, regardless of the distance between the pair of polar residues. These highly conserved and connected polar residues likely play an important role in structure stabilization.

This study uses sequence identity to wild type sequence as one of the criteria for judging the potential success of the designed sequence. Completely relying on this criterion is certainly a questionable practice because most sequences constrained to have high sequence identity to wild type sequence will not fold into the same structure of the wild-type sequence. For example, an average of 80% sequence identity will be resulted from replacing all surface polar residues (contact number <6) by nonpolar residues (Table IV). This sequence, however, will more likely aggregate rather than fold. Moreover, 35% sequence-identity threshold observed in Figure 1 may not be applicable to designed sequences because the structures can be disrupted by a few bad predictions of amino acid residues. Thus, sequence identity metric has to be used in combination of other native properties such as sequence complexity, the fraction of polar residues, and the frequency of 20 amino acid residues. We are currently searching for more robust theoretical criteria for judging the potential success of a designed sequence before expensive experiments are carried out as the final proof.

This article is limited to design with a fixed backbone to reduce the conformational space of sampling. A fixed backbone may have prohibited the sampling of more native-like sequences. Indeed, Saunders and Baker showed that the sequence identity between designed sequences and wild-type sequences can be improved after allowing flexibility. RosettaDesign-SR is not limited to a fixed backbone because newly implemented terms are sequence-dependent only. We will defer design with backbone flexibility to our future studies.

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Figure 1.
Population density of protein pairs as a function of the sequence identity between a protein pair calculated by FASTA \(^46\) (X axis) and the TM-score describing the structural similarity between a pair of proteins (Y axis). The population density is normalized for each sequence identity. The population density is described in the unit of the population density at a uniform distribution. The initial data is a 18 × 20 grid with sequence identity stepsize of 0.05 and TMscore stepsize of 0.05 based on pairwise comparison of 6665 proteins. The data is further smoothed by interpolating to a grid of 900×1000.
Figure 2.
Average sequence identity between designed sequences and wild type sequences for 33 training proteins as a function of the weight $w_{\text{profile}}$ for the sequence profile term.
Figure 3.
The frequency of 20 amino acid residues in sequences of 944 test proteins. Black bar corresponds to wild-type sequences. Gray bar corresponds to designed sequences by the original RosettaDesign. Open bar corresponds to designed sequence by RosettaDesign-SR.
Figure 4.
The sequence identity between top-1 ranked, designed sequences for 944 proteins and wild-type sequences as a function of fraction of surface residues (residue contact number <9). Open circles are those designed sequences that do not have any hit in a psi-blast search.
**Figure 5.**
The sequence identity between designed and wild-type sequences versus the highest sequence identity between the designed and the sequence obtained by performing a psi-blast search for designed sequences. Circle denotes the sequences designed by RosettaDesign-SR, and plus denotes the sequences designed by original RosettaDesign. Points with zero sequence identity between the designed and the sequence obtained by performing a psi-blast search for designed sequences refer to those designed sequences without any hits from a psi-blast search.
Table I

Comparison Between the Results Given by Wild-Type Sequences, Sequences Designed by RosettaDesign 2.3, RosettaDesign and Structure-Derived Sequence Profile (RosettaDesign-S), RosettaDesign-S Plus Repetitive, and Reference-State Energy Terms (RosettaDesign-SR)

<table>
<thead>
<tr>
<th>% Polar residues$^a$</th>
<th>Seq. ID (%$^b$</th>
<th>% LC$^c$</th>
<th>% LLL$^d$</th>
<th>% VVV$^e$</th>
<th>All</th>
<th>α</th>
<th>β</th>
<th>Coil</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wild-type</td>
<td>100</td>
<td>2.1</td>
<td>0.030</td>
<td>0.060</td>
<td>49.1</td>
<td>51.2</td>
<td>41.4</td>
<td>52.6</td>
</tr>
<tr>
<td>RosettaDesign</td>
<td>31.9</td>
<td>6.4</td>
<td>0.113</td>
<td>0.033</td>
<td>44.0</td>
<td>48.4</td>
<td>33.8</td>
<td>46.9</td>
</tr>
<tr>
<td>Sequence profile$^f$</td>
<td>24.2</td>
<td>15.1</td>
<td>0.030</td>
<td>0.714</td>
<td>28.2</td>
<td>24.2</td>
<td>21.0</td>
<td>38.3</td>
</tr>
<tr>
<td>RosettaDesign-S</td>
<td>35.6</td>
<td>21.3</td>
<td>0.783</td>
<td>0.597</td>
<td>37.4</td>
<td>36.9</td>
<td>28.2</td>
<td>45.1</td>
</tr>
<tr>
<td>RosettaDesign-SR</td>
<td>34.4</td>
<td>1.4</td>
<td>0.029</td>
<td>0.030</td>
<td>48.2</td>
<td>51.3</td>
<td>40.0</td>
<td>50.9</td>
</tr>
</tbody>
</table>

The results are obtained for the training set of 33 proteins averaged over 100 sequences designed for each protein.

$^a$Frequency of polar residues for helical, strand, and coil positions, respectively. Polar residues include D, E, H, K, N, Q, R, S, T, and Y.

$^b$The average sequence identity to wild type sequences of target proteins.

$^c$Percent of low-complexity regions defined by program SEG$^{72}$.

$^d$Percent of occurrence for three sequentially linked Leu residues.

$^e$Percent of occurrence for three sequentially linked Val residues.

$^f$Results based on the consensus sequence from structure derived sequence profile.
Table II

Optimized Reference State Energy $E_{ref}(s,I)$ Values for 8 Amino Acid Residue Types

<table>
<thead>
<tr>
<th>Type</th>
<th>A</th>
<th>E</th>
<th>G</th>
<th>I</th>
<th>K</th>
<th>L</th>
<th>P</th>
<th>V</th>
</tr>
</thead>
<tbody>
<tr>
<td>Helix</td>
<td>0.8</td>
<td>0.7</td>
<td>0</td>
<td>0</td>
<td>0.2</td>
<td>0.9</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Strand</td>
<td>0.3</td>
<td>0</td>
<td>0</td>
<td>0.9</td>
<td>0</td>
<td>0.4</td>
<td>0.1</td>
<td>1.2</td>
</tr>
<tr>
<td>Coil</td>
<td>0</td>
<td>0</td>
<td>0.2</td>
<td>1</td>
<td>0</td>
<td>0</td>
<td>1.6</td>
<td>0</td>
</tr>
</tbody>
</table>

Others are set to 0.
Table III

Comparison Between the Results Given by Wild-Type Sequences, Sequences Designed by RosettaDesign 2.3 and by RosettaDesign-SR

<table>
<thead>
<tr>
<th>Seq. ID (%a)</th>
<th>% LCb</th>
<th>% LLLd</th>
<th>% VVVe</th>
<th>All</th>
<th>α</th>
<th>β</th>
<th>Coil</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wild-type</td>
<td>100</td>
<td>3.1%</td>
<td>0.063</td>
<td>0.045</td>
<td>50.1</td>
<td>51.8</td>
<td>44.7</td>
</tr>
<tr>
<td>RosettaDesign</td>
<td>31.4</td>
<td>9.1%</td>
<td>0.244</td>
<td>0.093</td>
<td>45.6</td>
<td>50.1</td>
<td>36.1</td>
</tr>
<tr>
<td>RosettaDesign-SR</td>
<td>33.9</td>
<td>4.0%</td>
<td>0.039</td>
<td>0.038</td>
<td>48.2</td>
<td>51.1</td>
<td>41.5</td>
</tr>
</tbody>
</table>

The results are obtained for the test set of 944 proteins with the top 1 ranked sequence in 100 sequences designed for each protein.

*a Frequency of polar residues for helical, strand, and coil positions, respectively.

b The average sequence identity to wild type sequences of target proteins.

c Percent of low-complexity regions defined by program SEG72

d Percent of occurrence for three sequentially linked Leu residues.

e Percent of occurrence for three sequentially linked Val residues.
Table IV
Comparison Between the Results at Different Solvent Exposures Given by Wild-Type Sequences, Sequences Designed by RosettaDesign 2.3 and by RosettaDesign-SR

<table>
<thead>
<tr>
<th>% Pol. ID</th>
<th>% Seq. ID</th>
<th>% Polar residues</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Wild-type</td>
<td>RosettaDesign</td>
</tr>
<tr>
<td></td>
<td>100</td>
<td>22.4 (14.0/35.8)</td>
</tr>
<tr>
<td>0-5</td>
<td>100</td>
<td>25.7 (15.7/39.9)</td>
</tr>
<tr>
<td>6-10</td>
<td>100</td>
<td>36.3 (18.3/50.5)</td>
</tr>
<tr>
<td>11-15</td>
<td>100</td>
<td>49.6 (23.4/60.1)</td>
</tr>
<tr>
<td>≥16</td>
<td>100</td>
<td>60.5</td>
</tr>
</tbody>
</table>

The results are obtained for the test set of 944 proteins with the top 1 ranked sequence in 100 sequences designed for each protein.

- **% Pol. ID**: Fraction of polar residues in protein sequences.
- **% Seq. ID**: Average sequence identity between designed sequences and wild-type sequences.
- **% Polar residues**: Population of residues in four residue-contact bins.
- **Solvent exposure** is measured by residue contact number, the number of residues (Cα atoms) within 10 Å from a specific residue (three nearest sequentially linked residues are excluded from counting). They are divided into 4 bins.
- **Population of residues in four residue-contact bins.**
- **Sequence identity to polar and nonpolar residues in wild-type sequences.**
Table V

Sequence Identity Between Designed Sequences and Wild Type Sequences and Fraction of Polar Residues for All-α, All-β, and Other Proteins

<table>
<thead>
<tr>
<th></th>
<th>% Seq. ID&lt;sup&gt;a&lt;/sup&gt;</th>
<th>% Polar residues&lt;sup&gt;b&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>All-α&lt;sup&gt;c&lt;/sup&gt; (255)</td>
<td>All-β&lt;sup&gt;c&lt;/sup&gt; (166)</td>
</tr>
<tr>
<td>Wild-type</td>
<td>100</td>
<td>100</td>
</tr>
<tr>
<td>RosettaDesign</td>
<td>26.8</td>
<td>35.3</td>
</tr>
<tr>
<td>RosettaDesign-SR</td>
<td>30.2</td>
<td>36.9</td>
</tr>
</tbody>
</table>

<sup>a</sup> Average sequence identity between designed sequences and wild-type sequences.

<sup>b</sup> Fraction of polar residues in protein sequences.

<sup>c</sup> All-α proteins refer to the proteins with >10% helical residues and <10% strand residues. All-β proteins refer to the proteins with <10% helical residues and >10% strand residues.77

<sup>d</sup> The number of proteins in a given structural fold.