# SURFDESIGN: EFFECTIVE PROTEIN DESIGN ON MOLECULAR SURFACES

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### ABSTRACT

Structure-based inverse folding has been extensively explored in recent years. In contrast, surface-conditioned protein generation is still an under-explored area. Molecular surfaces characterized by a compact and smooth composition of atoms at their boundary hold a more direct relevance to biomolecular interactions and function. In this work, we introduce a novel framework named SurfDesign with several key improvements. Firstly, considering the theoretical fact that the molecular surface is a continuous manifold with infinite resolution, we propose surface-based equivariant message passing (SEMP) to incorporate the normal vector and curvatures and get aware of the manifold's Euclidean locality. Besides, a hybrid parameter-efficient fine-tuning (PEFT) technique is employed to combine the knowledge of protein language models (PLMs) with the surface geometric encoder. We extensively evaluate SurfDesign on the CATH, TS50, TS500, and PDB datasets, achieving an average recovery of more than 70%. Our work opens another road to designing functional proteins, underscoring the importance of including surface attributes in protein discovery.

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#### 1 INTRODUCTION

028 Proteins, as intricately folded chains of 029 amino acids, are fundamental to biological processes such as transcription, trans-031 lation, signaling, and cell cycle control. The advent of generative deep learning 033 (DL) (Huang et al., 2016; Song et al., 034 2020; Rives et al., 2021) has revolutionized protein design, shifting the focus away from traditional physics-based methods (see Figure 1). One prevalent ap-037 proach is to first design a target backbone structure and then identify a sequence that folds into this backbone. Despite the sig-040 nificant progress (Ingraham et al., 2019; 041 Jing et al., 2020; Dauparas et al., 2022; 042 Hsu et al., 2022; Gao et al., 2022a; Mao 043 et al., 2023; Zheng et al., 2023; Wu & 044 Li, 2024a; Qiu et al., 2024; Wang et al., 2024), the goal of protein design goes be-

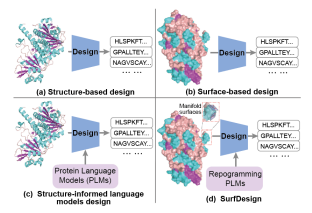


Figure 1: Prevailing setups of protein design, conditioned on backbone structures or molecular surfaces.

yond predicting a sequence that folds into a target backbone (Defresne et al., 2021). The ultimate
goal is to design proteins with desired functions, such as enzymes binding to specific substrates or
proteins inhibiting given targets. The inverse folding method has limitations, as it only specifies
geometric constraints through the backbone structure. To achieve desired functions, it is essential
to impose biochemical property constraints as well. For instance, two proteins with complementary
shapes may still not bind effectively due to poorly placed charges, polarity, or hydrophobicity at
their binding interface (Gainza et al., 2023).

Recent research (Song et al., 2024) has made strides in addressing this issue by designing functional proteins based on continuous surfaces augmented with biochemical properties. Albeit deep genera-

054 tive models show revolutionary capacity in this field, the current neural surface-conditioned protein 055 design still has undeniable flaws in devising more plausible proteins due to the limited expression 056 capabilities of their algorithms. First and foremost, in theory, molecular surfaces are continuous and 057 smooth 3D manifolds with infinite resolution (Lee et al., 2023; Sun et al., 2024). Existing point 058 cloud- (Song et al., 2024; Zhang et al., 2023) or mesh-based methods do not account for inherent connectivity and smoothness, treating surfaces as collections of discrete points. However, ideally, molecular surfaces are continuous, meaning no gaps or discrete points, and allow for differentiable 060 operations. Besides, the smoothness of manifolds indicates a well-defined tangent space at each 061 point and can be described using smooth functions. Secondly, the limited availability of experimen-062 tally determined protein surface data impedes progress in surface-conditioned design. For instance, 063 the known protein structures in the commonly-used CATH (Orengo et al., 1997) dataset are vastly 064 outnumbered by the sequence data in the UniRef (Suzek et al., 2015) sequence database. This dis-065 parity presents a challenge for data-hungry generative models, which struggle to comprehensively 066 explore the protein sequence space and often produce sub-optimal sequence predictions. Moreover, 067 from a biological perspective, molecular surfaces alone may not provide sufficient information, es-068 pecially in buried regions where sequential knowledge is more valuable yet largely neglected.

069 To address these challenges, we propose SurfDesign, a novel and effective algorithm for surfaceconditioned protein design (see Figure 2). SurfDesign captures the continuity and smoothness of 071 surface manifolds by analyzing the tangent space and curvatures near each point, where normal 072 vectors are used to approximate local geometry and curvatures are leveraged to measure deviations 073 from planarity. We then compute directional information between neighboring points and introduce 074 a surface-based equivariant message passing (SEMP) scheme to integrate manifold-specific geome-075 tries such as curvatures and directionality. Moreover, inspired by recent advances in employing pretrained protein language models (PLMs) for versatile protein design (Zheng et al., 2023; Gao et al., 076 2023; Qiu et al., 2024; Wang et al., 2024; Mao et al., 2023), we propose a hybrid parameter-efficient 077 fine-tuning (PEFT) technique to enhance our SEMP with the knowledge from PLMs. Comprehensive experiments have been conducted to evaluate our SurfDesign in the domain of inverse folding. 079 Our algorithm exhibits a substantial performance boost over current state-of-the-art methods, VFN-IF (Mao et al., 2023), KW-Design (Gao et al., 2023), and InstructPLM (Qiu et al., 2024), by a large 081 margin, achieving 74.13% and 72.14% sequence recovery on CATH 4.2 and 4.3 for single-chain 082 monomers. SurfDesign has also been trained on the entire PDB database with an impressive recov-083 ery of 81%. These results highlight SurfDesign's superior performance and potential in advancing 084 the field of protein design. Discussion on related works is put in Appendix B. 085

**Problem Statement.** Neural structure-conditioned protein design aims to find the amino acid sequence  $S = \{s_i \in Cat(20) : 1 \le i \le n\}$  folding into the desired structure  $\mathcal{X} = \{x_i \in \mathbb{R}^{4 \times 3} : 1 \le i \le n\}$ , where  $s_i$  belongs to one of the 20 residue types and  $\mathcal{X}$  denotes the spatial coordinates for 4 backbone atoms (*i.e.*,  $C_{\alpha}$ , C, N and O). It can be formulated as an end-to-end graph-to-sequence learning problem with a parameterized encoder-decoder neural network  $\mathcal{F}_{\vartheta}: \mathcal{X} \to S$ . Surfaceconditioned protein design is analogous to the structure-conditioned definition but generates functional proteins, which fold into expected surfaces  $\mathcal{Q}$  with associated biochemical properties (Song et al., 2024). Our objective therefore transfers to learn a function  $\mathcal{F}_{\vartheta}(\cdot)$ :

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$$\mathcal{F}_{\boldsymbol{\vartheta}}: \mathcal{Q} \to \mathcal{S} \tag{1}$$

Given sufficient surface-sequence paired data, the learning purpose is to maximize the conditional log-likelihood  $p(S|Q; \vartheta)$ . This approach allows for designing sequences that either have the highest likelihood or are generated using sampling algorithms to ensure diversity and novelty (Zheng et al., 2023). Remarkably, homologous proteins consistently share similar surfaces (Pearson & Sierk, 2005), so the surface-conditioned design is underdetermined. In other words, the valid amino acid sequence S may not be unique (Gao et al., 2022a).

- 102 2 METHOD
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104 2.1 PRELIMINARY AND BACKGROUND

**Surface Generation** The surface geometry of a protein is of crucial interest for protein-protein interaction analysis. We employ PyMol (DeLano et al., 2002) to obtain the raw molecular surface, where a probe of a certain radius ( $\sim 1$  Angstrom) is moved along the protein to calculate the Solvent

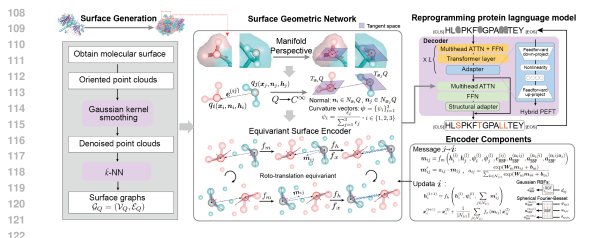


Figure 2: Illustration of SurfDesign. Smooth Surface graphs are acquired by PyMol or MSMS and processed via denoising. Then an equivariant surface encoder is appended to extract manifold representations. These features are further fed into the structural adapter of the protein language models for recovering masked amino acids.

128 Accessibility Surface (SAS) and Solvent Excluded Surface (SES). The consequent probe coordinates 129 are regarded as the molecular surface, defined by an oriented point cloud  $Q = \{q_i : 1 \le i \le m\}$ and m >> n. Each surface point  $q_i$  has a triplet of attributes  $(x_i, n_i, h_i)$ , where  $x_i \in \mathbb{R}^3$  and 130  $n_i \in \mathbb{R}^3$  is the 3D coordinates and unit normal vector, and  $h_i \in \mathbb{R}^{\phi_h}$  indicates the physicochemical 131 properties of  $q_i$  such as hydrophobicity, hbond, and charge. Then the surface graph is built via k-NN, 132 resulting in  $\mathcal{G}_Q = (\mathcal{V}_Q, \mathcal{E}_Q)$ . Notably, we also investigate the open source MSMS (Robinson et al., 133 2014) and BioPython (Cock et al., 2009) for surface generation and discover ignorable differences 134 in processing speeds among several toolkits. As raw point clouds generally carry noise and these 135 noisy points may limit the expressivity of molecular surfaces (Alexa et al., 2001), we borrow ideas 136 from Song et al. (2024) and apply the Gaussian kernel smoothing on raw point cloud data: 137

$$\boldsymbol{x}_{i} \leftarrow \sum_{j \in \mathcal{N}_{(i)}} \frac{\mathcal{K}(\boldsymbol{x}_{i}, \boldsymbol{x}_{j}) \cdot \boldsymbol{x}_{j}}{\sum_{t \in \mathcal{N}_{(i)}} \mathcal{K}(\boldsymbol{x}_{i}, \boldsymbol{x}_{t})}, \quad \mathcal{K}(\boldsymbol{x}, \boldsymbol{y}) = \exp^{-\frac{\|\boldsymbol{x}-\boldsymbol{y}\|^{2}}{\eta}},$$
(2)

where  $\mathcal{N}_{(i)}$  denotes the neighborhood of  $\boldsymbol{x}_i$  and  $\mathcal{K}(.,.)$  is the Gaussian kernel with  $\eta$  indicating distance scale in the point space. Here,  $\eta$  is set as  $\max\left(\left\{\|\boldsymbol{x}_i - \boldsymbol{x}_j\|^2\right\}_{j \in \mathcal{N}_{(i)}, i \in [m]}\right)$ .

#### 2.2 SURFACE GEOMETRIC NETWORK

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146 A Manifold Perspective for Molecular Surfaces. Theoretically, molecular surfaces are contin-147 uous manifolds with infinite resolution (Lee et al., 2023), which cannot be fully expressed by 148 existing mesh- (Gainza et al., 2020) or point-based (Sverrisson et al., 2021; Zhang et al., 2023; 149 Song et al., 2024) mechanisms. The key distinct property between manifold surfaces and con-150 ventional point clouds or meshes is that every point in the manifold is locally Euclidean. Mathe-151 matically, for  $\forall q_i \in Q$ , there exists a neighborhood  $U_{q_i}$  and a homomorphism  $f_{\text{homo}}(\cdot)$  such that  $f_{\text{homo}}: U_{q_i} \to V \subseteq \mathbb{R}^3$ , where V is an open ball in  $\mathbb{R}^3$ . In order to describe the local geometry of a manifold point  $q_i \in Q$ , we need to know at least (1) the linear approximation of the manifold in 152 153 its vicinity, which corresponds to the tangent space, and (2) how fast the surface bends or deviates 154 from being a plane near this point, which can be measured by *curvature*. 155

Towards this goal, we assume that the surface Q is a  $C^{\infty}$  differentiable manifold and  $T_{x_i}Q$  denotes the tangent space of any point  $x_i \in Q$ . Then we can acquire the unit normal vector  $n_i \in N_{x_i}Q$ perpendicular to  $T_{x_i}Q$ . If Q is implicitly described by a signed distance function (SDF) satisfying  $f_{\text{SDF}}(\cdot) = 0$ , then the normal at point  $x_i$  is equivalent to the gradient, *i.e.*,  $n_i = \nabla f_{\text{SDF}}(x_i)$ . Here, we draw the normal vector set  $\{n\}_{i=1}^m$  immediately from the software (*i.e.*, PyMol) and integrate this orientation knowledge into the geometric encoder to linearly approximate the manifold and achieve manifold-awareness. Prior studies (Zhang et al., 2023; Song et al., 2024) have seldom 162 163 considered this specialty of molecular surfaces and merely handle naive clouds. One exception, 163 dMaSIF (Strokach et al., 2020), notices this manifold uniqueness and computes the quasi-geodesic 164 distance as  $d_{ij} = ||\boldsymbol{x}_{ij}||^2 \cdot (2 - \boldsymbol{n}_i^\top \cdot \boldsymbol{n}_j)$  to naively resemble the geodesic coordinates in the tangent 165 space  $T_{\boldsymbol{x}_i}Q$ . However, its construction of tangent vectors destroys the equivariance.

Additionally, there are varying ways to define curvatures of 3D Riemannian manifolds intrinsically without reference to a larger space (Kobayashi & Nomizu, 1996), such as normal curvature  $k_n$ , geodesic curvature  $k_g$ , and geodesic torsion  $\tau_r$ . Those all relate the direction of curvatures to the unit normal vector  $n_i$ . Given a non-singular curve  $\gamma(q_i) \in Q$  parametrized by arc length, we can compute  $T_i = \gamma'(q_i)$  and  $t_i = n_i \times T_i$  to form the Darboux frame. The triple  $(T_i, t_i, n_i)$  defines a positively oriented orthonormal basis attached to each point of the curve  $\gamma(q_i)$ . Then the above  $T_i^{T} = (T_i) = (T_i) + (T_i)$ 

172 quantities are related by  $\begin{pmatrix} T' \\ t' \\ u' \end{pmatrix} = \begin{pmatrix} 0 & k_g & k_n \\ -k_g & 0 & \tau_r \\ -k_n & -\tau_r & 0 \end{pmatrix} \begin{pmatrix} T \\ t \\ u \end{pmatrix}$ . Inspired by progress in geometry 174

processing (Tian et al., 2023; Wu & Li, 2024b; Zhang et al., 2008), we estimate these quantities in a closed form from local points  $\mathcal{N}_{(i)}$ . Specifically, we first compute a covariance matrix for  $q_i$  and its neighborhood  $\mathcal{N}_{(i)}$ :

$$\boldsymbol{\Sigma} = \frac{1}{\|\mathcal{N}_{(i)}\|} \sum_{\mathbf{x}_j \in \mathcal{N}_{(i)}} \mathbf{x}_j \mathbf{x}_j^\top - \bar{\mathbf{x}} \bar{\mathbf{x}}^\top, \quad \boldsymbol{\Sigma} \in \mathbb{R}^{3 \times 3}.$$
 (3)

where  $\bar{\mathbf{x}}$  is the centroid of this point cluster. Then after the eigen-decomposition of  $\Sigma$  (*e.g.*, singular value decomposition or eigenvalue decomposition), eigenvalues can be attained as  $\epsilon_1, \epsilon_2$ , and  $\epsilon_3$  ( $\epsilon_1 \ge \epsilon_2 \ge \epsilon_3$ ). The three pseudo curvatures vectors  $\boldsymbol{\psi} = \{\psi_i\}_{i=1}^3$  can be therefore computed as:

$$\psi_i = \frac{\epsilon_i}{\sum_{j=1}^3 \epsilon_j}, \quad i \in \{1, 2, 3\}.$$
(4)

We employ  $\psi$  as a substitute and approximation of the Darboux frame  $(k_n, k_g, \tau_r)$ . It can be proved that this curvature feature  $\psi$  is roto-translation invariant (see Appendix E).

189 Directionality in Surface Point Clouds. The mani-190 fold characteristic of molecular surfaces introduces addi-191 tional directional information when considering pairwise 192 or ternary interactions among connected particles. To be 193 specific, for each neighboring point pair (i, j), two inter-194 secting planes (see Fig. 3) is formulated with respective 195 normals  $(n_i, n_j)$ . We denote the angles between normals 196 and the connecting directed line of two points  $(x_{ij}, x_{ji})$ by  $\varphi_{\mathbf{n}_i i j} = \angle n_i x_{i j}$  and  $\varphi_{\mathbf{n}_j j i} = \angle n_j x_{j i}$ . We denote 197 the dihedral angle between two half-phases as  $\theta_{\mathbf{n}_i i j \mathbf{n}_j} =$  $\angle \mathbf{n}_i \mathbf{n}_j \perp x_{ij}$ . In addition to the common distance 199  $\|\boldsymbol{x}_{ij}\|^2$ , these three angles provide a more comprehensive 200 view of understanding the relative position of  $(q_i, q_j)$  ly-201 ing in the surface manifold Q, which will also be incor-202

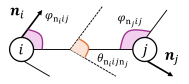


Figure 3: Angles hidden in the oriented surface point cloud, containing two intersection angles  $\varphi_{\mathbf{n}_i i j} = \angle \mathbf{n}_i \mathbf{x}_{i j}$  and  $\varphi_{\mathbf{n}_j j i} = \angle \mathbf{n}_j \mathbf{x}_{j i}$  as well as a dihedral angle  $\theta_{\mathbf{n}_i i j \mathbf{n}_i}$ .

porated into our surface modeling. For instance, for different values of  $(\varphi_{\mathbf{n}_i i j}, \varphi_{\mathbf{n}_j j i}, \theta_{\mathbf{n}_i i j \mathbf{n}_j})$ , a triplet of  $(\frac{\pi}{2}, \frac{\pi}{2}, 0)$  indicates a perfectly smooth region, while a triplet of  $(\pi, \pi, \pi)$  implies a severely sharp and steep curve.

**Equivariant Surface Encoder.** Finally, we draw inspiration from prevalent and modern equivariant algorithms (Satorras et al., 2021; Gasteiger et al., 2021; 2020b;a; Zhang et al., 2023; Song et al., 2024) and propose a surface-based equivariant message passing (SEMP) as the encoder of  $\mathcal{F}_{\vartheta}(\cdot)$ . Our SEMP architecture is roto-translation equivariant, leveraging both directional and curvature information. To begin with, by setting an interaction cutoff  $c_{int}$ , we calculate the 3D spherical Fourier-Bessel bases  $\left(\boldsymbol{a}_{\text{SBF}}^{(n_i ij)}, \boldsymbol{a}_{\text{SBF}}^{(n_j ji)}\right) \in 2 \times \mathbb{R}^{N_{\text{CBF}} \times N_{\text{SBF}} \times N_{\text{RBF}}}$  for two angles  $\varphi \in \left[\varphi_{\mathbf{n}_i ij}^{(l)}, \varphi_{\mathbf{n}_j ji}^{(l)}\right]$ to integrate orientation knowledge between each interactive particles in the surface:

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$$a_{\text{SBF},ovt}^{(l)}\left(\left\|\boldsymbol{x}_{ij}^{(l)}\right\|^{2},\varphi,\theta_{\mathbf{n}_{i}ij\mathbf{n}_{j}}^{(l)}\right) = \sqrt{\frac{2}{c_{\text{int}}^{3}j_{o+1}^{2}\left(z_{ov}\right)}}j_{o}\left(\frac{z_{ov}}{c_{\text{int}}}\left\|\boldsymbol{x}_{ij}^{(l)}\right\|^{2}\right)Y_{o}^{t}\left(\varphi_{\mathbf{n}_{i}ij}^{(l)},\theta_{\mathbf{n}_{i}ij\mathbf{n}_{j}}^{(l)}\right), \quad (5)$$

where  $o \in [N_{\text{CBF}}]$ ,  $v \in [N_{\text{SBF}}]$ , and  $t \in [N_{\text{RBF}}]$  control the degree, root, and order of the radial basis functions, respectively. Besides,  $j_o(\cdot)$  is the *o*-th degree spherical Bessel functions and  $z_{ov}$  is its corresponding *v*-th root.  $Y_o^t(\cdot)$  is the real *o*-th degree and *t*-th order spherical harmonics. Equ. 5 can be boiled down to a joint 2D basis if the order *t* is set to 0. By using  $Y_o^0(\cdot)$ , we obtain the 2D representation  $a_{\text{SBF}}^{(\mathbf{n}_i ij\mathbf{n}_j)} \in \mathbb{R}^{N_{\text{CBF}} \times N_{\text{SBF}}}$  based on  $\theta_{\mathbf{n}_i ij\mathbf{n}_j}^{(l)}$ .

Remarkably, those 2D/3D spherical Fourier-Bessel representations  $a_{\text{SBF}}^{(\mathbf{n}_i ij)}$ ,  $a_{\text{SBF}}^{(\mathbf{n}_j ji)}$ , and  $a_{\text{SBF}}^{(\mathbf{n}_i ij\mathbf{n}_j)}$ enjoy the roto-translation invariant property due to their exploitation of the relative distance as well as the invariant angles. Then those directional vectors along with pointwise curvatures are fed into SEMP to attain the initial messages  $m_{ij}$  as:

$$\boldsymbol{m}_{ij} = f_m \left( \boldsymbol{h}_i^{(l)}, \boldsymbol{h}_j^{(l)}, \boldsymbol{\psi}_i^{(l)}, \boldsymbol{\psi}_j^{(l)}, \boldsymbol{e}_{\mathsf{RBF}}^{(ij)}, \boldsymbol{a}_{\mathsf{SBF}}^{(\mathbf{n}_i ij)}, \boldsymbol{a}_{\mathsf{SBF}}^{(\mathbf{n}_i jj\mathbf{n}_j)}, \boldsymbol{a}_{\mathsf{SBF}}^{(\mathbf{n}_i ij\mathbf{n}_j)} \right), \tag{6}$$

where  $f_m$  is a multi-layer perception (MLP) appended with an activation function like SiLU (Nwankpa et al., 2018).  $e_{\text{RBF}}^{(ij)}$  is the radial basis function representation of the interatomic distance  $||\boldsymbol{x}_{ij}||^2$ . Then a softmax is employed to reweight the messages:

$$\boldsymbol{m}_{ij}' = a_{ij} \cdot \boldsymbol{m}_{ij}, \quad a_{ij} = \frac{\exp(\boldsymbol{W}_{\mathrm{m}}\boldsymbol{m}_{ij} + \boldsymbol{b}_{\mathrm{m}})}{\sum_{k \in \mathcal{N}_{(i)}} \exp(\boldsymbol{W}_{\mathrm{m}}\boldsymbol{m}_{ik} + \boldsymbol{b}_{\mathrm{m}})}$$
(7)

where the weight matrix  $W_m \in \mathbb{R}^{\phi_m \times 1}$  and vector  $b_m \in \mathbb{R}$  are learnable. After that, messages are propagated from the vicinity of each point  $q_i$  to update its node feature as well as coordinates:

$$\boldsymbol{h}_{i}^{(l+1)} = f_{h}\left(\boldsymbol{h}_{i}^{(l)}, \boldsymbol{\psi}_{i}^{(l)}, \sum_{j \in \mathcal{N}_{(i)}} \boldsymbol{m}_{ij}^{\prime}\right), \quad \boldsymbol{x}_{i}^{(l+1)} = \boldsymbol{x}_{i}^{(l)} + \frac{1}{\|\mathcal{N}_{(i)}\|} \sum_{j \in \mathcal{N}_{(i)}} f_{x}\left(\boldsymbol{m}_{ij}\right) \boldsymbol{x}_{ij}^{(l)}. \tag{8}$$

241 where  $f_h(\cdot)$  is another MLP and  $f_x: \mathbb{R}^{\phi_m} \to \mathbb{R}$  transforms  $m_{ij}$  into a scalar score to control 242 the impact of directional vector  $x_{ij}^{(l)}$ . Notably, as the position of each point  $x_i^{(l)}$  is moving as 243 the layer  $l \in [L]$  goes deeper with  $x_i^{(0)} = x_i$ , it is optional but recommended to adjust and re-244 calculate the curvature  $\psi_i$  and relevant angles  $(\varphi_{\mathbf{n}_i i j}, \varphi_{\mathbf{n}_j j i}, \theta_{\mathbf{n}_i i j \mathbf{n}_j})$  simultaneously. As angles 245  $(\varphi_{\mathbf{n}_i i j}, \varphi_{\mathbf{n}_j j i}, \theta_{\mathbf{n}_i i j \mathbf{n}_j})$  depend on each normal vector pair  $(\mathbf{n}_i \text{ and } \mathbf{n}_j)$ , we adopt the local least fit-246 ting method (Mitra & Nguyen, 2003) to estimate and renew  $\{n_i\}_{i=1}^m$ . In specific, for  $q_i$ 's updated 247 coordinates  $x_i^{(l)}$  at the *l*-th layer, we compute the covariance  $\Sigma^{(l)}$  according to Equ. 3 and decompose it to obtain three sorted eigenvalues as well as their corresponding eigenvectors  $(\nu_1, \nu_2, \nu_3)$ . 248 249 250 Then  $\nu_3$  with the least eigenvalue is selected as the normal vector  $n^{(l)}$  at the *l*-th layer.

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#### 2.3 REPROGRAMMING PROTEIN LANGUAGE MODELS

**PEFT for SurfDesign.** Recent works have explored the possibility of transforming PLMs (Rives 254 et al., 2021; Lin et al., 2022; Hu et al., 2022) into protein design models, and massive evidence 255 demonstrates that the emergent evolutionary knowledge hidden in those PLMs can vastly facili-256 tate the structure-conditioned protein design. Concretely, LM-Design (Zheng et al., 2023), In-257 structPLM (Qiu et al., 2024), KW-Design (Gao et al., 2023), and VFN-IF-ESM (Mao et al., 2023) 258 report improvements in CATH 4.2 of 10.8% (recovery  $50.22\% \rightarrow 55.65\%$ ), 73.9% (perplexity 259  $10.28 \rightarrow 2.68$ ), 14.4% (recovery 54.74%  $\rightarrow 62.67$ %), and 17.6% (recovery 51.66%  $\rightarrow 60.77$ %), 260 respectively. Motivated by this progress, we also leverage PLMs as the decoder of  $\mathcal{F}_{\vartheta}(\cdot)$  and stack 261 several parameter-efficient fine-tuning (PEFT) techniques to fully release the potential of PLMs and 262 significantly reduce the memory budget. Specifically, we utilize a hybrid PEFT method combined with a structural adapter (Zheng et al., 2023) and LoRA (Hu et al., 2021) with a rank of r = 4 and 264 a scaling constant of  $\alpha = 8$ . It is worth mentioning that there is still no consensus on which sort 265 of PEFT strategies are most suitable for PLMs (Sledzieski et al., 2024), and we practically find our hybrid mechanism more effective than a singular one for surface-conditioned protein design. 266

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Training. Following LM-Design (Zheng et al., 2023), we employ the conditional masked language modeling (CMLM) to better accommodate PLMs that are tasked with MLM (Devlin et al., 2018) as the training objective. Given the surface Q, CMLM decomposes the sequence into masked and

Table 1: Sequence design performance and ablation studies on CATH 4.2 held-out test split. The **best performance** is shown in bold, while the <u>best baseline</u> is indicated with an underline. ESM-IF is tested on CATH 4.2, although it was originally trained and evaluated on CATH 4.3.

Params. 1.4M / 1.4M 1.56M / 1.56M 2.1M / 2.1M 1.0M / 1.0M 3.6M / 3.6M 1.9M / 1.9M 142M / 142M	Short           8.29         8.39           7.09         7.23           7.32         6.21	Single-chain 8.74 8.83 7.49 7.84 7.63	All 6.40 6.63 6.05 5.36	Short 29.44 28.14 32.62 30.60	Single-chain 28.26 28.46 31.10	35.82
1.56M / 1.56M 2.1M / 2.1M 1.0M / 1.0M 3.6M / 3.6M 1.9M / 1.9M	8.39 7.09 7.23 7.32	8.83 7.49 7.84	6.63 6.05 5.36	28.14 32.62	28.46	35.91 35.82 37.64
2.1M / 2.1M 1.0M / 1.0M 3.6M / 3.6M 1.9M / 1.9M	7.09 7.23 7.32	7.49 7.84	6.05 5.36	32.62		35.82 37.64
1.0M / 1.0M 3.6M / 3.6M 1.9M / 1.9M	7.23 7.32	7.84	5.36		31.10	37.64
3.6M / 3.6M 1.9M / 1.9M	7.32			30.60		57.04
1.9M / 1.9M		7.63	6 20	2.000	28.95	39.47
	6.21		6.30	34.16	32.66	41.31
142M / 142M	0.21	6.68	4.61	36.35	34.43	45.96
	6.93	6.65	3.96	35.28	33.78	48.95
6.6M / 6.6M	6.04	6.31	4.55	39.84	38.53	51.66
5.0M / 659M	7.01	6.58	4.41	35.19	40.00	54.41
11.9M / 664M	6.77	6.46	4.52	37.88	42.47	55.65
5.0M / 659M	-	-	-	-	_	54.54
89.1M / 6.6B	<u>3.22</u>	3.17	2.68	61.59	59.29	57.51
6.4M / 798M	5.48	5.16	3.46	44.66	45.45	60.77
5.4M / 5.4M	5.70	5.86	4.17	41.34	40.98	54.74
5.4M / 15B	4.92	4.22	3.36	50.00	52.13	62.67
5.8M / 5.8M	-	-	3.13	-	-	57.78
5.3M / 5.3M	3.21	3.10	3.08	62.70	64.88	65.35
4.8M / 655M	3.08	2.93	2.76	65.43	67.06	66.27
5.3M / 656M	2.43	2.44	2.41	73.74	75.17	74.13
	11.9M / 664M 5.0M / 659M 89.1M / 6.6B 6.4M / 798M 5.4M / 5.4M 5.4M / 15B 5.8M / 5.8M 5.3M / 5.3M 4.8M / 655M	11.9M / 664M         6.77           5.0M / 659M         -           89.1M / 6.6B         3.22           6.4M / 798M         5.48           5.4M / 15B         4.92           5.8M / 5.8M         -           5.3M / 5.3M         3.21           4.8M / 655M         3.08	11.9M / 664M         6.77         6.46           5.0M / 659M         -         -           89.1M / 6.6B         3.22         3.17           6.4M / 798M         5.48         5.16           5.4M / 5.4M         5.70         5.86           5.4M / 15B         4.92         4.22           5.8M / 5.8M         -         -           5.3M / 5.3M         3.21         3.10           4.8M / 655M         3.08         2.93	11.9M / 664M         6.77         6.46         4.52           5.0M / 659M         -         -         -           89.1M / 6.6B         3.22         3.17         2.68           6.4M / 798M         5.48         5.16         3.46           5.4M / 5.4M         5.70         5.86         4.17           5.4M / 15B         4.92         4.22         3.36           5.8M / 5.8M         -         -         3.13           5.3M / 5.3M         3.21         3.10         3.08           4.8M / 655M         3.08         2.93         2.76	$\begin{array}{cccccccccccccccccccccccccccccccccccc$	$\begin{array}{c ccccccccccccccccccccccccccccccccccc$

observed ones as  $S = S_{\text{masked}} \cup S_{\text{obs}}$  and assumes a conditional independence over identities of target residues  $s_i \in S_{\text{masked}}$ . Then it requires the model to predict a set of target amino acids  $S_{\text{masked}}$  from the remaining observed residues  $S_{\text{obs}}$ :

$$p(\mathcal{S}_{\text{masked}}|\mathcal{S}_{\text{obs}}, \mathcal{Q}; \theta) = \prod_{s_i \in \mathcal{S}_{\text{masked}}} p(s_i|\mathcal{S}_{\text{obs}}, \mathcal{Q}; \theta)$$
(9)

where  $S_{\text{masked}}$  is randomly masked. Moreover, Zheng et al. (2023) presents a coarse-to-fine manner to reconstruct a protein native sequence from its corrupted version. We also explore this inference scheme with iterative refinement (Savinov et al., 2021) but discover no benefit.

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EXPERIMENTS

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We evaluate SurfDesign on various benchmarks for fixed backbone protein sequence design, including single-chain monomers (Sec. 3.1) and multi-chain protein complexes (Sec. 3.2). More experimental details, dataset statistics, and additional results are elaborated in the Appendix A.

306 **Baselines and Datasets.** A wide variety of baseline approaches are established for a fair com-307 parison and most of them are open source. Among them, StructGNN (Ingraham et al., 2019), 308 GraphTrans (Ingraham et al., 2019), GVP (Jing et al., 2020), ProteinMPNN (Dauparas et al., 2022), 309 AlphaDesign (Gao et al., 2022b), PiFold (Gao et al., 2022a), UniIF (Gao et al., 2024), and etc. are GNN-based algorithms. In contrast, DenseCPD (Qi & Zhang, 2020) is a CNN-based approach. 310 Besides, DPLM (Wang et al., 2024), InstructPLM (Qiu et al., 2024), LM-Design (Zheng et al., 311 2023), KW-Design (Gao et al., 2023) and VFN-IF-ESM (Mao et al., 2023) leverage and integrate 312 the knowledge of pretrained PLMs. SurfPro (Song et al., 2024) is a surface-based framework. Us-313 ing the same splitting strategy as the compared systems (Jing et al., 2020; Dauparas et al., 2022; 314 Gao et al., 2022a), proteins in CATH 4.2 were partitioned into 18,024/608/1,120 samples for train-315 ing, validation, and testing, respectively. To compare with ESM-IF (Hsu et al., 2022), structures 316 in CATH 4.3 were split into 16,153/1,457/1,797 samples for training, validation, and testing, sepa-317 rately. To provide a head-to-head comparison with ESM-IF, no extra data such as AF2DB (Varadi 318 et al., 2022) is utilized for training SurfDesign. To evaluate the generative quality thoroughly, we 319 report perplexity, and median recovery rate on short-chain, single-chain, and all-chain settings as 320 usual. The multi-chan protein design employs the dataset curated by Dauparas et al. (2022), which 321 was preprocessed by clustering sequences at 30% identity, resulting in 25,361 clusters. Following ProteinMPNN's setup, those clusters were divided randomly into 23,358/1,464/1,539 samples for 322 training, validation, and testing, respectively. This strategy ensures that none of the chains from the 323 target chain or biounits of the target chain were present in the other two sets.

Models	<b>Perplexity</b> $(\downarrow)$			Recovery Rate ( <sup>†</sup> )		
Models	Short	Single-chain	All	Short	Single-chain	All
GVP (Hsu et al., 2022)	7.68	<sup>†</sup> 6.12	6.17	32.60	39.40	39.20
ProteinMPNN (Dauparas et al., 2022)	6.31	6.32	4.85	40.30	39.02	48.25
ESM-IF (Hsu et al., 2022)	8.18	†6.33	6.44	31.30	38.50	38.30
+ 1.2M AF2 Data	6.05	†4.00	4.01	38.10	51.50	51.60
PiFold (Gao et al., 2022a)	5.88	5.55	4.47	42.86	43.69	50.68
VFN-IF (Mao et al., 2023)	-	-	-	45.34	53.70	52.18
UniIF (Gao et al., 2024)	-	-	-	45.41	54.46	53.05
LM-Design-MPNN (Zheng et al., 2023)	5.88	5.66	4.19	45.71	46.15	56.38
LM-Design-PiFold (Zheng et al., 2023)	5.66	5.52	4.01	46.84	48.63	56.63
KW-Design (Gao et al., 2023)	<u>5.47</u>	5.23	<u>3.49</u>	43.86	45.95	<u>60.38</u>
SurfDesign	5.08	4.97	3.12	66.74	71.30	72.14

#### Table 2: Sequence design on CATH 4.3. †: SINGLE-CHAIN in Hsu et al. (2022) is defined differently.

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#### 3.1 SINGLE-CHAIN PROTEIN DESIGN

340 **Results.** Table 1 and 2 document the results of SurfDesign in comparison to the comprehensive 341 strong baselines on the CATH (Orengo et al., 1997) benchmark. It can be concluded that SurfDe-342 sign consistently achieves state-of-the-art performance in distinct settings. In particular, we observe 343 that SurfDock is the foremost to exceed 70% recovery on not only CATH 4.2 but also CATH 4.3, 344 illustrating its superior capacity in restoring effective protein sequences. Besides, on the full CATH 4.2 benchmark, SurfDesign achieves a perplexity of 2.41 and a recovery of 74.13%, outpassing the 345 previous state-of-the-art VFN-IF-ESM (Mao et al., 2023) by 28.27% and 18.28%, separately. It also 346 induces recovery improvements of 19.72% and 26.78% on the short and single-chain subsets, respec-347 tively. Furthermore, SurfDesign surpasses SurfPro, another surface-based algorithm, by 23.00% and 348 28.29% in the overall metrics, respectively. The outstanding phenomenon exists for the CATH 4.3 349 benchmark as well, where SurfDesign outperforms the strongest competitor KW-Design (Gao et al., 350 2023) by 10.60% and 19.49% for perplexity and recovery, respectively. To summarize, SurfDesign 351 enhances surface-conditioned sequence generation with greater efficiency, thanks to the significant 352 advancements and open-source contributions from the entire community, building on the foundation 353 laid by previous pioneers.

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#### 3.2 MULTI-CHAIN PROTEIN COMPLEX DESIGN

357 Results. A protein only functions when it docks, combines, and interacts with other macro-358 molecules, forming multi-chain protein complexes. Therefore, studying protein sequence design 359 for multi-chain assembled structures is crucial for drug design. This motivates us to assess whether 360 SurfDesign can more effectively manage protein complex design. From Table 3, we conclude that the recovery is generally higher for longer proteins and all models achieve higher recovery rates on 361 PDB than CATH datasets. More importantly, SurfDesign attains the best performance with a recov-362 ery of more than 80%. This phenomenon indicates that SurfDesign can design both single-chain 363 proteins and multi-chain complexes. This makes SurfDesign more versatile regarding the categories 364 and scenarios where it can be deployed, creating opportunities to use it for designing specific protein 365 complexes.

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#### 3.3 ZERO-SHOT GENERALIZATION TO NEW PROTEIN FAMILIES

369 **Results.** TS50 and TS500 are commonly used independent test sets to assess model generalization 370 for unseen proteins introduced by Li et al. (2014). Towards this goal, we evaluate SurfDesign 371 trained on CATH 4.2 and 4.3 respectively and report the results in Table 4. It can be discovered that 372 SurfDesign outpasses prior studies by a large margin on all benchmarks. Specifically, it achieves a 373 perplexity of 2.05 and a recovery rate of 82.16 on TS50, which outperforms the previous state-of-374 the-art algorithm, VFN-IF-ESM, by 18.65% and 12.08%, respectively. Meanwhile, on the TS500 375 dataset, SurfDesign obtains a perplexity of 1.98 and a recovery rate of 84.70. These numbers are better than VFN-IF-ESM by 22.04% and 16.80%, separately. In addition, for those trained in CATH 376 4.3, SurfDesign consistently achieves the best. In a nutshell, SurfDesign is the pioneer to transcend 377 82% and 84% recovery on the TS50 and TS500.

380	Models		Recove	<b>rv</b> (†)	
381	length	L < 100	$100 \le L < 500$	$500 \le L < 1000$	Full
382	StructGNN (Ingraham et al., 2019)	0.41	0.41	0.42	0.41
383	GraphTrans (Ingraham et al., 2019)	0.40	0.39	0.40	0.40
384	GCA (Tan et al., 2023) GVP (Jing et al., 2020)	0.41 0.44	0.41 0.42	0.42 0.45	0.41 0.43
385	AlphaDesign (Gao et al., 2022b)	0.48	0.49	0.50	0.49
386	ProteinMPNN (Dauparas et al., 2022) PiFold (Gao et al., 2022a)	0.52 0.54	$0.53 \\ 0.58$	$0.55 \\ 0.60$	$0.53 \\ 0.58$
387	LM-Design-MPNN (Zheng et al., 2023)		_	_	0.61
388	LM-Design-GVP (Zheng et al., 2023)	-	-	-	0.62
389	KWDesign (Gao et al., 2023)	0.59	0.66	0.67	0.66
390	SurfDesign	0.74	0.79	0.82	0.81

Table 3: Performance on multi-chain protein complex dataset (*i.e.*, PDB).

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Table 4: Performance comparison on TS50 and TS500. Following prior literature, we mainly report the results using models trained on CATH 4.2. Numbers in the brackets are results from models trained on CATH 4.3.

Models		TS50		TS500			
Wodels	Perplexity $(\downarrow)$	Recovery (†)	Worst (†)	Perplexity $(\downarrow)$	Recovery (†)	Worst (†)	
DenseCPD (Qi & Zhang, 2020)	-	50.71	-	_	55.53	-	
StructGNN (Ingraham et al., 2019)	5.40	43.89	26.92	4.98	45.69	0.05	
GraphTrans (Ingraham et al., 2019)	5.60	42.20	29.22	5.16	44.66	0.03	
GVP (Jing et al., 2020)	4.71	44.14	33.73	4.20	49.14	0.09	
GCA (Tan et al., 2023)	5.09	47.02	28.87	4.72	47.74	0.03	
AlphaDesign (Gao et al., 2022b)	5.25	48.36	32.31	4.93	49.23	0.03	
KW-Design (Gao et al., 2023)	3.10	62.79	39.31	2.86	69.19	0.02	
VFN-IF (Mao et al., 2023)	3.58	59.54	-	3.19	63.65	-	
VFN-IF-ESM (Mao et al., 2023)	2.52	73.30	-	2.54	72.49	-	
InstructPLM (Qiu et al., 2024)	<u>2.29</u>	67.99	-	2.42	64.22	-	
ProteinMPNN (Dauparas et al., 2022)	3.93 (3.62)	54.43 (54.22)	37.24 (41.18)	3.53 (3.27)	58.08 (57.23)	0.03 (0.04	
PiFold (Gao et al., 2022a)	3.86 (3.70)	58.72 (59.68)	37.93 (38.14)	3.44 (3.70)	60.42 (59.95)	0.03 (0.05	
LM-Design-MPNN (Zheng et al., 2023)	3.82 (3.60)	56.92 (58.13)	35.17 (39.14)	2.13(2.15)	64.30 (63.76)	0.04 (0.04	
LM-Design-PiFold (Zheng et al., 2023)	3.50 ( <u>3.27</u> )	57.89 ( <u>61.38</u> )	39.74(46.75)	3.19 (3.09)	67.78 ( <u>66.56</u> )	0.02 (0.04	
SurfDesign	2.05(2.03)	82.16(83.44)	41.30(47.81)	1.98 (1.96)	84.70(85.12)	0.10 (0.0	

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#### 3.4 MORE RESULTS AND ANALYSIS

411 Ablation Studies. We conduct systematic experiments 412 to investigate the contributions of different components in SurfDesign, shown in Table 1. It can be observed that 413 the knowledge of PLMs provides a large improvement of 414 13.43% in recovery  $(65.35\% \rightarrow 74.13\%)$  and a decrease 415 of 24.29% in perplexity (3.21  $\rightarrow$  2.43). Moreover, the 416 incorporation of directionality and curvatures also con-417 tributes to the superiority of SurfDesign with an improve-418 ment of 11.86% in recovery and 12.68% in perplexity. 419

420 Structural Contexts. To further understand the ac-421 tion mechanism of SurfDesign, we dissect its perfor-422 mance according to different structural contexts in Fig-423 ure 4. Structure-based LM-Design shows high recov-424 ery on structurally constrained residues in the folding 425 core, while low recovery in structurally less constrained 426 residues on surface areas and loops. SurfDesign significantly enhances the recovery on structurally constrained 427 and less-constrained residues, particularly those on the 428 surface regions. 429

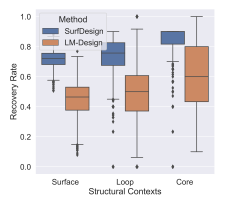


Figure 4: Comparison of sequence recovery w.r.t. structural contexts regarding SASA and interaction interface, on CATH 4.2 single-chain proteins.

Surface Recovery. Unlike the conventional structure-conditioned protein design, the ultimate goal 431 of our surface-based design is to generate proteins with higher surface similarity of key regions such

433	Table 5: Evaluation on the surface				Table 6: Structure recovery comparison based on the self-					
434	recovery on CATH 4.2.				consistent protocol from Yim et al. (2023). ‡: benchmarked results are quoted from Mao et al. (2023).					
435 436	Models	$\Big  \ IoU \ (\uparrow)$	$CD\left( \downarrow\right)$	$NC(\uparrow)$						
430	IM Declar	0.00	5 072	0 4026	Motrice DiFold <sup>‡</sup> I M Design <sup>‡</sup> VEN IF FSM <sup>‡</sup> SurfDesign					

	Models	100(1)	CD (↓)	$\mathbf{NC}(1)$					
436	LM-Design	0.90	5.972	0.4236	Metrics	PiFold <sup>‡</sup>	LM-Design <sup>‡</sup>	VFN-IF-ESM <sup>‡</sup>	SurfDesign
437	VFN-IF-ESM	0.92	4.688	0.4859	scTM > 0.5	90.98%	89.42%	93.29 %	96.17%
438	SurfDesign	0.98	2.873	0.6241	scRMSD < 2.0	60.35 %	58.41%	64.16%	72.83%
					-				

as the binding or interaction site (Lai et al., 2024). In order to evaluate the similarity between two 3D molecular shapes, we follow ideas from (Sun et al., 2024) and use three evaluation metrics commonly used in 3D modeling from three aspects: volume, distance, and normal vectors. They are Volumetric Intersection over Union (IoU), Chamfer distance (CD), and Normal Consistency (NC) (computational details are in Appendix A.2). As shown in Table 5, SurfDesign can reconstruct the molecular surfaces well, which accords with the motivation of our surface-conditioned design. Visualization of generated and ground truth surfaces are available in Appendix A.3.

Structure Recovery. We compare SurfDesign with strong baselines in terms of protein struc-448 ture recovery on CATH 4.2, reported in Table 6. Following standard evaluation procedures (Yim 449 et al., 2023; Mao et al., 2023), ESMFold was used to predict structures of designed sequences. A 450 case study of visualization comparison using Alphafold-3 is displayed in Appendix C. Two selfconsistent metrics, scTM ( $\uparrow$ ) and scRMSD ( $\downarrow$ ) are leveraged to assess the similarity between desired 452 and designed protein structures. It can be found that SurfDesign is more likely to generate protein sequences with expected structures. 454

455 Scalability of PLMs. The scaling law w.r.t model sizes 456 of PLMs has recently been studied (Zheng et al., 2023; 457 Qiu et al., 2024). To understand the influence of PLM 458 model sizes over SurfDesign's capacity, we increase the 459 parameters of ESM-2 from 8M to 3B. As indicated in Fig-460 ure 5, a similar phenomenon has been discovered where 461 the performance of SurfDesign improves as PLMs scale. When integrating knowledge from the largest PLM (3B), 462 SurfDesign achieves a 76.01% recovery rate on CATH 463 4.2. This coincidence highlights the great potential of em-464 powering surface-conditioned design with cutting-edge 465 PLMs (Kaplan et al., 2020). 466

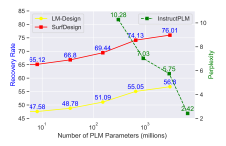


Figure 5: Performance in terms of model scales of PLMs using ESM2.

#### 4 CONCLUSION

We propose SurfDesign, a novel method that integrates the geometric and biochemical information from molecular surfaces to design proteins with the knowledge of protein language models. SurfDesign is the foremost model that achieves 70% recovery on CATH 4.2, CATH 4.3, TS50, TS500, and PDB, demonstrating its generalizability and effectiveness. We look forward to future efforts in extending its application to real-world problems such as antibody and enzyme discovery.

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740 741 742	A EXPERIMENTAL DETAILS
743 744 745 746	<b>Training and metrics.</b> The models were trained up to 50 epochs by default using the Adam optimizer on 4 A100 GPUs. We used the same training settings as ProteinMPNN (Dauparas et al., 2022) and LM-Design (Zheng et al., 2023), where the batch size was set to approximate 6000 residues, and the Adam optimizer was aligned with a NOAM learning rate scheduler. Following previous works,

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Implementation. PyMol is adopted for surface generation in our implementation. We have tried
 the fast sampling algorithm introduced by dMaSIF (Sverrisson et al., 2021) and used by later studies (Wu & Li, 2024b), which approximates the protein surface as the level set of a smooth distance
 function. However, this sampling mechanism has unacceptable randomness and is abandoned for
 SurfDesign. As for the biochemical feature computation, we follow MaSIF (Gainza et al., 2020)
 and calculate three key invariant point inputs, including the Poisson Boltzmann electrostatics using

CHAIN set contains proteins recorded as a single chain in PDB (Berman et al., 2002).

perplexity and *median* recovery scores are reported. In Table 1 and 2, two subsets of the entire test

set are also reported. Particularly, the SHORT set contains proteins up to length 100, and the SINGLE

APBS <sup>1</sup>, the hydrophobicity <sup>2</sup>, and the free electrons/protons <sup>3</sup>. After a further ablation study, we discover that the hydrophobicity and the charge are pivot to the performance improvement while the electrostatics is not necessary.

For Figure 4, we employ RSA to determine the surface and core. To be specific, residues with RSA greater than 0.25 are considered on the surface, while residues with RSA less than 0.1 are regarded as core residues. We use the DSSP algorithm to decide the loop regions.

#### 764 A.1 DATASET INFORMATION

Table 7 documents the vertex count statistics for the CATH datasets. We observe an equal distribution over vertex in different splits. Besides, comparing our surface with SurfPro (Song et al., 2024), it can be found that our surface is more sparse with nearly half of the average vertex per residue. This difference is due to the different computation techniques adopted by various software for surface generation (*e.g.*, PyMol and MSMS).

Table 7: Vertex counts statistics for surfaces from the CATH 4.2 and CATH 4.3 datasets.

Vertex Count		CATH 4.2		CATH 4.3			
vertex Count	Train	Validation	Test	Train	Validation	Test	
Average Vertex Count Per Residue	53.47	53.56	53.31	53.36	55.27	53.11	
Maximum Vertex Count	27,817	25,614	25,433	27,110	27,817	25,968	
Minimum Vertex Count	1,923	2,315	2,022	1,923	2,011	2,000	
Preprocess Time Per Protein		0.38s			0.36s		

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#### A.2 SURFACE COMPARISON

#### 783 A.2.1 EVALUATION METRICS 784

Motivated by DSR (Sun et al., 2024), we employ IoU, CD, and NC to assess the similarity between the molecular surfaces of designed proteins and target proteins. For simplicity, these three metrics are all normalized to a range of 0 - 1. They provide a comprehensive evaluation of the model's performance from different perspectives and are defined as follows.

**IoU.** IoU compares the reconstructed volume with the ground truth shape (higher is better). For two arbitrary shapes  $A, B \subseteq \mathbb{S} \in \mathbb{R}^n$  is attained by  $IoU = \frac{|A \cap B|}{|A \cup B|}$ .

**CD.** CD is a standard metric to evaluate the distance between two point sets  $\mathcal{X}_1, \mathcal{X}_2 \subset \mathbb{R}^n$ (lower is better) as  $d_C(\mathcal{X}_1, \mathcal{X}_2) = \frac{1}{2} (d_{\overrightarrow{C}}(\mathcal{X}_1, \mathcal{X}_2) + d_C(\mathcal{X}_2, \mathcal{X}_1))$ , where  $d_{\overrightarrow{C}}(\mathcal{X}_1, \mathcal{X}_2) = \frac{1}{|\mathcal{X}_1|} \sum_{\boldsymbol{x}_1 \in \mathcal{X}_1} \min_{\boldsymbol{x}_2 \in \mathcal{X}_2} ||\boldsymbol{x}_1 - \boldsymbol{x}_2||$ .

NC. NC evaluates estimated surface normals (higher is better). Normal consistency between two normalized unit vectors  $n_i$  and  $n_j$  is defined as the dot product between the two vectors. For evaluating the surface normals, given the object surface points and normal vectors:  $X_{\text{pred}} = \{(\boldsymbol{x}_i, \overrightarrow{n_i})\}$ , and the ground truth surface points and normal vectors:  $X_{gt} = \{(\boldsymbol{y}_j, \overrightarrow{m_j})\}$ , the surface normal consistency between  $X_{\text{pred}}$  and  $X_{gt}$ , denoted as  $\Gamma$ , is defined as:  $\Gamma(X_{gt}, X_{\text{pred}}) = \frac{1}{|X_{gt}|} \sum_{j \in |X_{gt}|} |\overrightarrow{n_j} \cdot \overrightarrow{m}_{\theta}(\boldsymbol{y}_j, X_{\text{pred}})|$ , where  $\theta(\boldsymbol{y}_j, X_{\text{pred}} := \{(\boldsymbol{x}_i, \overrightarrow{n_i})\}) = \underset{i \in |X_{\text{pred}}|}{\text{argmin}} \|\boldsymbol{y}_j - \boldsymbol{x}_i\|_2^2$ .

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&</sup>lt;sup>1</sup>https://github.com/LPDI-EPFL/masif/blob/master/source/triangulation/
computeAPBS.py

<sup>&</sup>lt;sup>2</sup>https://github.com/LPDI-EPFL/masif/blob/master/source/triangulation/ computeHydrophobicity.py

<sup>&</sup>lt;sup>3</sup>https://github.com/LPDI-EPFL/masif/blob/master/source/triangulation/ compuSurfDesignteCharges.py

# 810 A.3 SURFACE VISUALIZATION

In addition to protein structure restoration, we show the surface similarity between designed and
ground truth proteins. We envision the surface of designed proteins and target proteins in Figure 6.
A heavy overlap can be found between the point clouds of the designed protein surface and the
ground truth protein surface with a pretty low CD and significantly high IoU. These all indicate that
SurfDesign produces proteins with expected surface shapes.

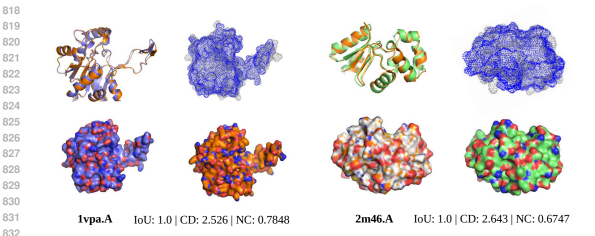


Figure 6: Comparision between original and designed surfaces, where molecular surfaces are visualized from two perspectives: the point cloud view and the manifold view.

#### B RELATED WORK

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Structure-based Protein Design. The protein folding problem, a longstanding challenge in 840 biology, has seen significant advancements through the application of AI techniques like Al-841 phaFold (Jumper et al., 2021) and RoseTTAFold (Baek et al., 2021). The complementary problem, 842 known as *inverse folding*, has also garnered increasing attention. The most primitive group relies on 843 multi-layer perceptrons (MLPs) to predict the probability of 20 amino acids for each residue based 844 on handcrafted features. SPIN (Li & Koehl, 2014) achieved a 30% recovery rate on the TS50 dataset 845 by incorporating torsion angles, sequence profiles, and energy profiles. This was further improved by 846 SPIN2 (O'Connell et al., 2018), which added features such as backbone angles, local contact num-847 ber, and neighborhood distance, reaching a 34% recovery rate. Concurrently, other methods, like 848 the one proposed by Wang et al. (2018), employed features including backbone dihedrals, solvent-849 accessible surface area (SASA) of backbone atoms, secondary structure types, and unit direction vectors, achieving a 33% recovery rate. Another category uses 2D or 3D CNN to extract protein fea-850 tures. For instance, SPROF (Chen et al., 2019) adopts 2D CNN to learn residue representations from 851 the distance matrix and achieves a 40.25% recovery on TS500. ProDCoNN (Zhang et al., 2022a) 852 designs a nine-layer 3D CNN with multi-scale convolution kernels and achieves 42.2% recovery on 853 TS500. DenseCPD (Qi & Zhang, 2020) further enhanced recovery to 55.53% using the DenseNet 854 architecture (Huang et al., 2017). As proteins can be natured represented as graphs, GNNs are exten-855 sively employed to consider structural constraints, with nodes and edges representing residue infor-856 mation and pairwise interactions, respectively. Notable works include GraphTrans (Ingraham et al., 857 2019), which introduced a graph attention encoder and autoregressive decoder, and GVP (Jing et al., 858 2020), which incorporated geometric vector perceptrons for learning from scalar and vector features. 859 Subsequent developments enclose GCA (Tan et al., 2023), which introduces global graph attention 860 for learning contextual features, AlphaDesign (Gao et al., 2022b), which presents a simplified graph 861 encoder and a constraint-aware decoder based on GVP, ProteinMPNN (Dauparas et al., 2022), which capitalizes on the benefits of an auto-regressive encoding-decoding scheme and message-passing up-862 dating techniques, and PiFold (Gao et al., 2022a), which improves the traditional encoding-decoding 863 framework by introducing virtual atoms and backbone dihedrals. Lately, VFN (Mao et al., 2023)

proffers learnable vector computations between coordinates of frame-anchored virtual atoms and
 exhibits an impressive 62.67% recovery.

Despite the enormous advancements, the diversity of generated sequences is limited by the small 867 scale of training data. ESM-IF (Hsu et al., 2022) addressed this by leveraging the accurate pro-868 tein folding predictions of AlphaFold2 to train a large-scale inverse folding framework using GVP. LM-Design (Zheng et al., 2023) tackles the data limitation by fine-tuning ESM models and em-870 ploying embeddings from pre-trained structural encoders to recover design sequences through con-871 ditional mask prediction. Subsequently, InstructPLM (Qiu et al., 2024) utilizes the cross-modality 872 alignment in LLMs and introduces structure prompts to fine-tune ProGen2 (Nijkamp et al., 2023). 873 KW-Design (Gao et al., 2023) proposes a knowledge-aware module that refines low-quality residues 874 with knowledge from ESM and GearNet (Zhang et al., 2022b). Another interesting line (Wang et al., 2024) demonstrates their self-supervised discrete diffusion probabilistic framework is versatile pro-875 tein learners for tasks like structure-conditioned sequence generation. 876

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878 **Protein Surface Modeling.** The characteristics of the molecular surface dictate the type and 879 strength of the interactions that a protein can have with other molecules. It is defined by van der 880 Waals (vdW) radii(Connolly, 1983) and is commonly represented as meshes derived from signed 881 distance functions. MaSIF (Gainza et al., 2020) pioneered the use of mesh-based geometric DL 882 to abstract the internal parts of the protein fold and explore protein interactions. A subsequent study (Sverrisson et al., 2021) reduced pre-computation costs by modeling molecular surfaces as 883 point clouds with atom categories assigned to each point. Other seminal works have linked protein 884 surfaces with structural information in a multimodal manner (Somnath et al., 2021) incorporat-885 ing comprehensive pretraining strategies (Wu & Li, 2024b) using implicit neural representations 886 (INRs) (Park et al., 2019) for self-supervised learning Lee et al. (2023) and dynamic structure mod-887 eling Sun et al. (2024). Despite these efforts, protein design based on surface features remains underexplored. Recent advancements, such as the work by Gainza et al. (2023) on expanding MaSIF 889 for *de novo* binder design, and SurfPro (Song et al., 2024), which eliminates the need for handcrafted 890 feature calculations, have started to address this gap by generating functional proteins directly from 891 surface data.

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Parameter-efficient Fine-tuning for Language Models. The development of protein language 894 models (PLMs) has been accelerated by the availability of vast datasets of amino acid se-895 quences (Rives et al., 2021; Lin et al., 2022; Rao et al., 2019; Elnaggar et al., 2020; Madani et al., 896 2020; Nijkamp et al., 2023). However, training and storing full copies of large PLMs for various 897 downstream tasks are increasingly impractical, necessitating parameter-efficient fine-tuning (PEFT) 898 methods. Recent works (Sledzieski et al., 2024; Zeng et al., 2023) have shown that PEFT techniques, 899 such as LoRA (Hu et al., 2021) and prompt tuning (Lester et al., 2021), achieve competitive or superior performance compared to full fine-tuning, with significantly reduced memory requirements 900 901 for tasks like protein-protein interaction prediction, signal peptide prediction, and homo-oligomer symmetry prediction. In addition, biologists attempt to incorporate structural information into PLMs 902 using advanced PEFT tools. For example, LM-Design (Zheng et al., 2023) introduces a lightweight 903 adapter to realize structural awareness, referred to as structural surgery on PLMs. SES-Adapter (Tan 904 et al., 2024) integrates structural data by converting it into sequential vectors through tools like Fold-905 Seek (Van Kempen et al., 2024) and DSSP (Kabsch & Sander, 1983), enabling cross-modal atten-906 tion calculations. It defeats structure-aware PLMs such as SaProt (Su et al., 2023) across standard 907 datasets, including those for thermostability, metal ion binding, gene ontology (GO) annotations, 908 and subcellular localization predictions.

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## C VISUALIZATION RESULTS

In this section, we visualize several protein structure restoration results of SurfDesign, as shown in Figrue 7 and Figure 8. The designed structures were obtained using the latest AlphaFold 3 (Abramson et al., 2024)<sup>4</sup>

<sup>&</sup>lt;sup>4</sup>We employed the Alphafold Server for inference at https://alphafoldserver.com/.

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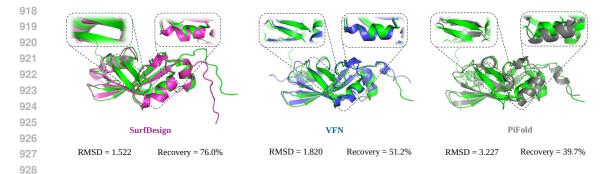


Figure 7: Visualization results of a challenging sample (PDB 2KRT). We use AlphaFold3 to recover the structure based on the predicted sequence and compare it against the experimentally determined ground-truth structure.

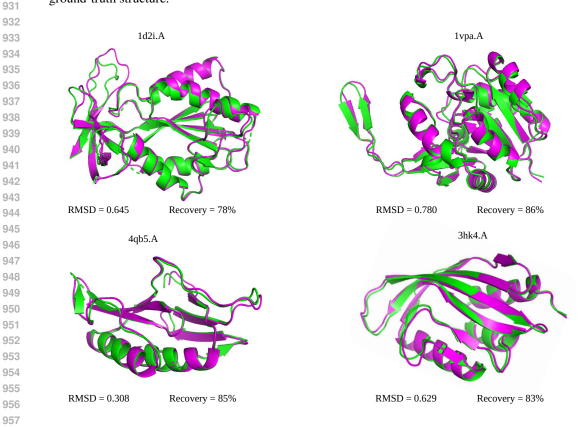


Figure 8: Visalization of SurfDesign, where the green and pink ones are ground truth and designed structures, respectively.

### D REFOLDABILITY ANALYSIS

964 Following Wang et al. (2023), firstly, to assess whether the generated sequences can respect the 965 structure condition, we evaluate the agreement of the ground truth structure with the predicted struc-966 tures using the TM-score (Zhang & Skolnick, 2004). We refer to this metric as Ref-TM. Further-967 more, to evaluate the folding stability of the generated sequences, we compute the mean value of 968 the per-residue confidence estimate pLDDT predicted by the structure prediction models, referred 969 to as Ref-pLDDT. As pLDDT is a reliable predictor of disorder (Tunyasuvunakool et al., 2021), AlphaFold2 (Jumper et al., 2021), OmegaFold (Wu et al., 2022), and ESMFold (Lin et al., 2022) 970 are leveraged as a structure prediction model, which helps minimize deviations due to the choice of 971 model.

We examine SurfDesign on the same 82 test samples of the CATH dataset and results are reported in Table 8. We observe that SurfDesign stands out as the leading design method across the refoldability metrics, competitive with ProteinMPNN. It achieves 0.89 Ref-TM and 89.42 Ref-pLDDT with Al-phaFold2 prediction. ProteinMPNN is slightly behind with a 0.87 Ref-TM and 87.89 Ref-pLDDT, followed by LM-Design.

Design method	ES TM	<b>MFold</b> pLDDT	Om TM	e <b>gaFold</b> pLDDT	Alpl TM	h <b>aFold2</b> pLDDT	Recovery%
Wildtype	0.80	74.91	0.75	78.39	0.90	89.87	100
Uniform Natural frequencies	0.05	27.68 30.53	0.05 0.07	31.53 35.59	0.06	33.68 35.02	5.00 5.84
AF-Design ESM-Design	0.53	61.37 59.65	0.53 0.38	72.04 62.66	0.52	75.29 60.02	15.95 17.33
StructTrans GVP ProteinMPNN PiFold	$ \begin{array}{c c} 0.72 \\ 0.73 \\ \underline{0.80} \\ 0.71 \\ 0.73 \end{array} $	68.85 69.67 <u>76.53</u> 67.55 72.12	$ \begin{array}{c c} 0.64 \\ 0.67 \\ \underline{0.76} \\ 0.64 \\ 0.70 \\ \end{array} $	70.35 74.33 <b>80.75</b> 70.21 77.58	$ \begin{array}{c c} 0.79 \\ 0.83 \\ \underline{0.87} \\ 0.82 \\ 0.85 \end{array} $	80.66 84.29 <u>87.89</u> 82.54 87.26	35.89 39.46 41.44 44.86
LM-Design SurfDesign	0.73 0.81	72.12 79.35	0.70 <b>0.76</b>	<u>80.11</u>	0.85 0.89	87.20 <b>89.42</b>	<u>51.23</u> <b>70.19</b>

Table 8: Refoldability metric and recovery metric on the CATH dataset. We employ **bold** and <u>underline</u> to highlight the best and suboptimal results on each metric. We use TM and pLDDT to represent Ref-TM and Ref-pLDDT.

In addition to the amino acid recovery rate, we have incorporated Foldable Diversity and sc-TM as recommended in to further verify the diversity and self-consistency of the generated sequences. Foldable Diversity evaluates only those sequence pairs that are structurally consistent with the in-put protein backbone, providing a more targeted diversity metric that avoids penalizing high-quality, diverse designs. Self-consistency TM score (sc-TM), following [D], gauges the consistency of struc-tural predictions for generated sequences, leveraging a fixed threshold of  $TM_{\min} = 0.7$  as imple-mented by [B]. We refer to https://github.com/flagshippioneering/pi-rldif for computation, and the results are shown below. The analysis shows that SurfDesign maintains high structural consistency with competitive diversity, outperforming other methods on foldable diversity metrics and provid-ing substantial evidence of the model's capability to generate high-quality, diverse sequences that remain faithful to the structural constraints of input proteins. 

1008	Dataset Model	Foldable Diversity $\uparrow$	sc-TM ↑
1009 1010	ProteinMPNN (T=0, RD)	20%	0.80
1011	ProteinMPNN (T=0.1)	23%	0.67
1012	ProteinMPNN (T=0.2)	3%	0.30
1012	ProteinMPNN (T=0.3)	0.1%	0.14
1014	PiFold (T=0.1) PiFold (T=0.2)	$23\% \\ 8\%$	0.72 0.38
1015	KWDesign (T=0.1)	18%	0.38
1016	KWDesign (T=0.2)	23%	0.58
1017	SurfDesign	23%	0.84
1018	Suitesign	<b>23</b> 70	0.01

Table 9: Foldable diversity on CATH-all.

#### E MATHEMATICAL ANALYSIS

Here we demonstrate that the curvature feature  $\psi$  is roto-translation invariant. Firstly, suppose we translate the entire neighborhood  $\mathcal{N}_{(i)}$  by a vector  $\mathbf{t} \in \mathbb{R}^3$ , so each point  $\mathbf{x}_j \in \mathcal{N}_{(i)}$  is transformed

to  $\mathbf{x}'_j = \mathbf{x}_j + \mathbf{t}$ . - When computing the covariance matrix  $\Sigma$ , the centroid  $\overline{\mathbf{x}}$  is subtracted from each point in  $\mathcal{N}_{(i)}$ . The centroid after translation becomes  $\overline{\mathbf{x}}' = \overline{\mathbf{x}} + \mathbf{t}$ , so the translated covariance matrix becomes:

$$\boldsymbol{\Sigma}' = \frac{1}{\left\| \mathcal{N}_{(i)} \right\|} \sum_{\mathbf{x}_j \in \mathcal{N}_{(i)}} \left( \mathbf{x}_j + \mathbf{t} \right) \left( \mathbf{x}_j + \mathbf{t} \right)^\top - \overline{\mathbf{x}}' \overline{\mathbf{x}}^\top.$$
(10)

1032 Expanding this, we get:

 $\boldsymbol{\Sigma}' = \frac{1}{\|\mathcal{N}_{(i)}\|} \sum_{\mathbf{x}_j \in \mathcal{N}_{(i)}} \mathbf{x}_j \mathbf{x}_j^\top + \mathbf{t} \mathbf{t}^\top + 2\mathbf{t} \cdot \sum_{\mathbf{x}_j \in \mathcal{N}_{(i)}} \mathbf{x}_j^\top / \|\mathcal{N}_{(i)}\| - (\overline{\mathbf{x}} + \mathbf{t})(\overline{\mathbf{x}} + \mathbf{t})^\top.$ (11)

1036 This simplifies back to the original  $\Sigma$  since t terms cancel out in the computation of  $\Sigma$  after trans-1037 lating by t. Therefore, the covariance matrix  $\Sigma$  is invariant under translations.

Suppose we apply a rotation  $\mathbf{R} \in SO(3)$  to all points in  $\mathcal{N}_{(i)}$ , where  $\mathbf{R}$  is an orthogonal matrix with determinant 1. Then each point  $\mathbf{x}_j \in \mathcal{N}_{(i)}$  is transformed to  $\mathbf{x}'_j = \mathbf{R}\mathbf{x}_j$ . The centroid  $\overline{\mathbf{x}}$  also transforms under the rotation, so the new centroid is  $\overline{\mathbf{x}}' = \mathbf{R}\overline{\mathbf{x}}$ . The covariance matrix  $\Sigma'$  after rotation becomes:

$$\boldsymbol{\Sigma}' = \frac{1}{\left\| \mathcal{N}_{(i)} \right\|} \sum_{\mathbf{x}_j \in \mathcal{N}_{(i)}} \mathbf{R} \mathbf{x}_j \left( \mathbf{R} \mathbf{x}_j \right)^\top - \overline{\mathbf{x}}' \overline{\mathbf{x}}'^\top.$$
(12)

1045 Expanding the terms, we obtain:

$$\boldsymbol{\Sigma}' = \mathbf{R} \left( \frac{1}{\left\| \mathcal{N}_{(i)} \right\|} \sum_{\mathbf{x}_j \in \mathcal{N}_{(i)}} \mathbf{x}_j \mathbf{x}_j^\top - \overline{\mathbf{x}} \overline{\mathbf{x}}^\top \right) \mathbf{R}^\top = \mathbf{R} \boldsymbol{\Sigma} \mathbf{R}^\top.$$
(13)

Since a rotation is a similarity transformation, the eigenvalues of  $\Sigma'$  are the same as those of  $\Sigma$ . Therefore, the eigenvalues  $\epsilon_1, \epsilon_2$ , and  $\epsilon_3$ , which are used to compute  $\psi$ , remain unchanged under rotations.