

# Optimized atomic statistical potentials: assessment of protein interfaces and loops

Guang Qiang Dong, Hao Fan, Dina Schneidman-Duhovny, Ben Webb and Andrej Sali\*

Department of Bioengineering and Therapeutic Sciences, Department of Pharmaceutical Chemistry and California Institute for Quantitative Biosciences (QB3), University of California, San Francisco, CA 94158, USA

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## ABSTRACT

**Motivation:** Statistical potentials have been widely used for modeling whole proteins and their parts (e.g. sidechains and loops) as well as interactions between proteins, nucleic acids and small molecules. Here, we formulate the statistical potentials entirely within a statistical framework, avoiding questionable statistical mechanical assumptions and approximations, including a definition of the reference state.

**Results:** We derive a general Bayesian framework for inferring *statistically optimized atomic potentials* (SOAP) in which the reference state is replaced with data-driven 'recovery' functions. Moreover, we restrain the relative orientation between two covalent bonds instead of a simple distance between two atoms, in an effort to capture orientation-dependent interactions such as hydrogen bonds. To demonstrate this general approach, we computed statistical potentials for protein-protein docking (SOAP-PP) and loop modeling (SOAP-Loop). For docking, a near-native model is within the top 10 scoring models in 40% of the PatchDock benchmark cases, compared with 23 and 27% for the state-of-the-art ZDOCK and FireDock scoring functions, respectively. Similarly, for modeling 12-residue loops in the PLOP benchmark, the average main-chain root mean square deviation of the best scored conformations by SOAP-Loop is 1.5 Å, close to the average root mean square deviation of the best sampled conformations (1.2 Å) and significantly better than that selected by Rosetta (2.1 Å), DFIRE (2.3 Å), DOPE (2.5 Å) and PLOP scoring functions (3.0 Å). Our Bayesian framework may also result in more accurate statistical potentials for additional modeling applications, thus affording better leverage of the experimentally determined protein structures.

**Availability and implementation:** SOAP-PP and SOAP-Loop are available as part of MODELLER (<http://salilab.org/modeller>).

**Contact:** sali@salilab.org

**Supplementary information:** Supplementary data are available at *Bioinformatics* online.

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## 1 INTRODUCTION

Computational modeling can be used to predict the structures of whole proteins or their parts (e.g. loops and sidechains) as well as complexes involving proteins, peptides, nucleic acids and small molecules (Audie and Swanson, 2012; Baker and Sali, 2001; Dill and MacCallum, 2012; Ding *et al.*, 2010; Skolnick

*et al.*, 2013; Wass *et al.*, 2011). A modeling method requires a conformational sampling scheme for proposing alternative structures and a scoring function for ranking them. Significant progress has been made on both fronts (Fernández-Recio and Sternberg, 2010; Moult *et al.*, 2011). In particular, many physics-based energy functions and statistical potentials computed from known protein structures have been described (Andrusier *et al.*, 2007; Benkert *et al.*, 2008; Betancourt and Skolnick, 2004; Betancourt and Thirumalai, 1999; Brenke *et al.*, 2012; Chuang *et al.*, 2008; Colovos and Yeates, 1993; Cossio *et al.*, 2012; Dehouck *et al.*, 2006; Fan *et al.*, 2011; Ferrada *et al.*, 2007; Gao and Skolnick, 2008; Gatchell *et al.*, 2000; Hendlich *et al.*, 1990; Huang and Zou, 2010; Jones, 1999; Keasar and Levitt, 2003; Kocher *et al.*, 1994; Li *et al.*, 2013; Liu and Gong, 2012; Liu and Vakser, 2011; Lu and Skolnick, 2001; Lu *et al.*, 2008; McConkey *et al.*, 2003; Melo and Feytmans, 1997; Melo *et al.*, 2002; Miyazawa and Jernigan, 1996; Park and Levitt, 1996; Pierce and Weng, 2007; Qiu and Elber, 2005; Rajgaria *et al.*, 2008; Rata *et al.*, 2010; Reva *et al.*, 1997; Rojnuckarin and Subramaniam, 1999; Rykunov and Fiser, 2010; Samudrala and Moult, 1998; Shapovalov and Dunbrack, 2011; Shen and Sali, 2006; Simons *et al.*, 1997; Sippl, 1993; Summa *et al.*, 2005; Tanaka and Scheraga, 1975; Wang *et al.*, 2004; Xu *et al.*, 2009; Zhang and Zhang, 2010; Zhao and Xu, 2012; Zhou and Skolnick, 2011; Zhou and Zhou, 2002; Zhu *et al.*, 2008).

Derivation of a statistical potential has often been guided by an analogy between a sample of known native structures and the canonical ensemble in statistical mechanics, suggesting that the distributions of spatial features in the sample of native structures follow the Boltzmann distribution (Sippl, 1990). Thus, statistical potentials are generally calculated in two steps: (i) extracting a probability distribution of a spatial feature (e.g. a distance spanned by a specific pair of atom types) from a sample of known protein structures and (ii) normalizing this distribution by a reference distribution (e.g. the distribution of all distances, regardless of the atom types). Statistical potentials can differ in a number of aspects, including the sample of known protein structures, the protein representation (e.g. centroids of amino acid residues,  $C_{\alpha}$  atoms and all atoms), the restrained spatial feature (e.g. solvent accessibility, distance, angles and orientation between two sets of atoms), the sequence features (e.g. amino acid residue types, atom types, residue separation in sequence and chain separation), the treatment of sparse samples and the definition of the reference state. Here, we optimize the accuracy of a statistical potential over most of these aspects. This optimization challenge is addressed by formulating a statistical

\*To whom correspondence should be addressed.

potential independently from any assumptions grounded in statistical mechanics; instead, we rely on a Bayesian approach based on data alone. Although the proposed theory applies to any kind of a statistical potential, we illustrate it by deriving specific statistical potentials for protein–protein docking and loop modeling.

## 2 METHOD

We begin by defining statistical potentials in terms of distributions extracted from known protein structures (Section 2.1), followed by a description of a protocol to actually compute a statistical potential (Sections 2.2–2.7, Fig. 1).

### 2.1 Theory

For structure characterization of a given protein sequence by either experiment or theory, we ideally need a joint probability density function (pdf) for the structure, given everything we know about it (Shen and Sali, 2006). In general, our knowledge can come from different kinds of experiments with the protein (e.g. X-ray crystallography), physical theories (e.g. a molecular mechanics force field) and/or statistical inference (e.g. all known structures or only homologous known structures). Here, we focus on a joint pdf for a given sequence based on the knowledge of all known protein structures deposited in the Protein Data Bank (PDB) (Kouranov *et al.*, 2006); thus, our joint pdf is a statistical potential.

To derive the joint pdf for a structure of a sequence, we need to approximate it by using terms that can actually be computed from the PDB. The structure  $X$  of an amino acid sequence is defined by the set of its features  $\{f^{(m)}\}$ ,  $m = 1 \dots n$ , such as a distance between two specific

atoms. Thus, we can approximate the joint pdf by the product of pdfs (restraints) for individual features:

$$p(X) \approx \prod_{1 \leq m \leq n} p(f^{(m)}), \quad (1)$$

Without any loss of accuracy, we define the restraint  $p(f^{(m)})$  as the ratio between the feature distribution  $p(f^c | Q_K)$  from a sample of informative features in a set of proteins  $Q_K$  with known structures (e.g. for a distance, all distances spanned by the same atom types in  $Q_K$ ) and an unknown recovery function  $g(f^c | Q_K)$ :

$$p(f^{(m)}) = p(f^c | Q_K) / g(f^c | Q_K), \quad (2)$$

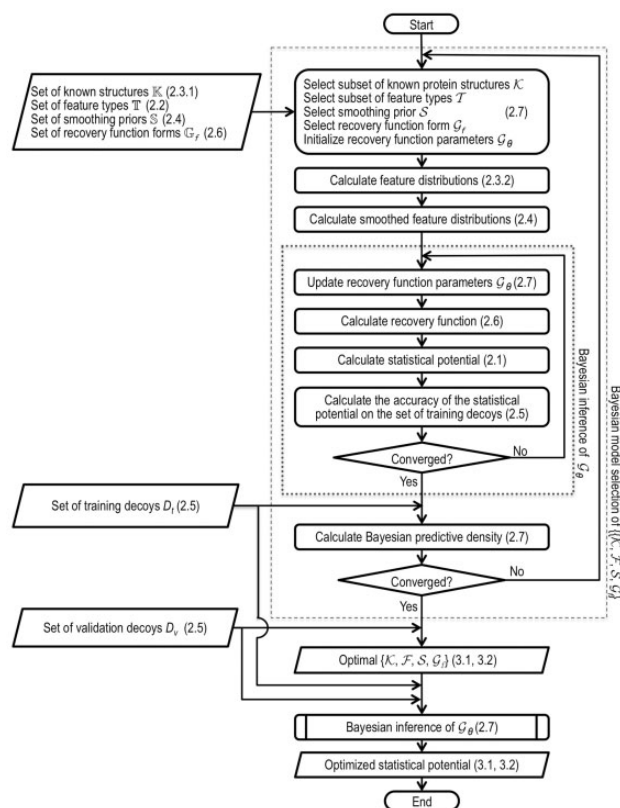
In other words, the recovery function is defined such that the product of restraints approximates the joint pdf as well as possible (*c.f.*, Equation 1), while minimizing the number of parameters that need to be fit to the data. Construction of the sample of informative features involves a compromise between including only features of known structures that are most likely to resemble the predicted feature  $f^{(m)}$  (which minimizes sample size) and minimizing the statistical noise (which maximizes sample size). The features used in the sample are termed to be of the same type  $c$  as the inferred feature (Section 2.2). The restraints on all features of  $X$  of type  $c$  are calculated from the same set of informative features and thus are the same. Here, the sample of informative features includes all features of the same type from representative known protein structures (Section 2.3).

### 2.2 Feature types

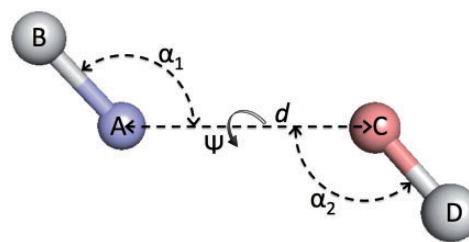
To illustrate the general theory mentioned earlier in the text, we derive optimized statistical potentials for assessing protein–protein interfaces (SOAP-PP) and loop conformations (SOAP-Loop). We restrain the following feature types:

**2.2.1 Atomic distance** Distance  $d|a_1, a_2, b_s$  is considered to depend on atom types  $a_1$  and  $a_2$  as well as the ‘covalent separation’ between the two atoms ( $b_s$ ). The atom type depends on the residue type, resulting in the total of 158 atom types for the 20 standard residue types (Shen and Sali, 2006). Covalent separation is measured in three ways. First, by the minimum number of covalent bonds between the two atoms (bond separation). Second, by the number of residues separating the two atoms in the polypeptide chain (residue separation). Third, by chain separation, which is 0 if the atoms are in the same chain and 1 otherwise. The distance is mapped in the range from 0 to a parameterized distance cutoff, such as 15 Å.

**2.2.2 Orientation between a pair of covalent bonds** Orientation  $d, \alpha_1, \alpha_2, \psi|t_1, t_2, b_s$  is defined by a distance  $d$ , two angles  $\alpha_1, \alpha_2$  and a dihedral angle  $\psi$  (Fig. 2). It is considered to depend on covalent bond types ( $t_1, t_2$ ) defined in turn by their atom types and covalent separation ( $b_s$ ); there are 316 covalent bond types for the 20 standard residue types.



**Fig. 1.** Flowchart for optimizing statistical potentials. The corresponding sections in the text are indicated



**Fig. 2.** Distance and angles between two covalent bonds, A–B and C–D.  $d$ , distance between atoms A and C.  $\alpha_1$ , angle between atoms B, A and C.  $\alpha_2$ , angle between atoms A, C and D.  $\psi$ , dihedral angle between atoms B, A, C and D.  $b_s$  is defined using atoms A and C

**2.2.3 Relative atomic surface accessibility** Accessibility  $s|a$  is considered to depend on the atom type ( $a$ ) (Sali and Blundell, 1993).

## 2.3 Feature distributions

**2.3.1 Known protein structures** A small fraction of the known protein structures from the PDB (and their decoy structures) is used only for assessing the accuracy of statistical potentials (Section 2.5). The remaining structures from the PDB are filtered to construct the known protein structure set  $\mathbb{K}$ , including only structures determined by X-ray crystallography at the resolution better than 2.2 Å and  $R_{\text{free}}$  better than 25%. Three additional subsets of representative structures were obtained by requiring at most 30, 60 and 95% sequence identity to any other representative structure, respectively, with preference for structures determined at higher resolutions and with lower  $R_{\text{free}}$  values. A statistical potential is optimized by choosing among the entire set  $\mathbb{K}$  or its three subsets to estimate the feature distributions  $p(f^c|Q_K)$ .

**2.3.2 Calculation of feature distributions** The sample for computing this distribution is the set of the individual features of type  $c$  in protein set  $Q_K$ , where each feature is represented by the distribution of this feature  $-p(f^{c(m)}|Q_K)$ . The feature distribution  $p(f^c|Q_K)$  is the average of these sample distributions. For a distance and an angle,  $p(f^{c(m)}|Q_K)$  is approximated by a Gaussian distribution  $p'(f^{c(m)}|Q_K)$  with the mean equal to the observed value and the standard deviation computed by the propagation (Neuhauser, 2010) of the uncertainties of individual atomic positions, which in turn are estimated from the atomic isotropic temperature factors (Carugo and Argos, 1999; Cruickshank, 1999; Schneider, 2000). For relative atomic surface accessibility,  $p(f^{c(m)}|Q_K)$  is approximated using a delta function  $p'(f^{c(m)}|Q_K)$  centered at feature  $f^{c(m)}$  in  $K$ . The approximated feature distribution  $p'(f^c|Q_K)$  is then computed from the approximated sample distributions  $p'(f^{c(m)}|Q_K)$ .

## 2.4 Bayesian smoothing and smoothing priors

The feature distributions  $p'(f^c|Q_K)$  can be noisy when the sample  $K$  is relatively small, as is often the case for the orientation between a pair of covalent bonds (Fig. 3A). Thus, we use Bayesian inference to calculate a smooth feature distribution:

$$p(p(f^c|Q_K)|p'(f^c|Q_K)) \propto p(p'(f^c|Q_K)|p(f^c|Q_K)) \cdot p(p(f^c|Q_K)) \quad (3)$$

where  $p(f^c|Q_K)$  is the ideal distribution without noise from an infinitely large set of known structures. Both the likelihood  $p(p'(f^c|Q_K)|p(f^c|Q_K))$  and the prior  $S \equiv p(p(f^c|Q_K))$  are multivariate Gaussian distributions (Rasmussen and Williams, 2005). The smoothness of  $p(f^c|Q_K)$  is specified by the prior  $S$ ; here, the prior is a multivariate Gaussian distribution with a zero mean and a squared exponential covariance function (Mackay, 2003). The characteristic length scale of the covariance function defines the range over which the two points are still correlated (the

smoothness of the curve). We set the characteristic length equal to a scale parameter  $L$  multiplied by 0.2 Å for distance, 10° for angles and 0.1% for atomic surface accessibility. A set of smoothing priors  $S$  is obtained by varying  $L$ . Using a scale of 2.0 as an example, the inferred  $p(f^c|Q_K)$  is significantly smoother than  $p'(f^c|Q_K)$  (Fig. 3B).

## 2.5 Decoys and assessment criteria

**2.5.1 Learning set for SOAP-PP** This set consists of 176 native complex structures in the pairwise protein docking benchmark 4.0 (Hwang et al., 2010) and ~4500 decoys for each of the complexes generated using PatchDock (Duhovny et al., 2002).

**2.5.2 Testing set for SOAP-PP** This set consists of 176 native complex structures in the pairwise protein docking benchmark 4.0 (Hwang et al., 2010) as well as ~212 000 decoys for each of the complexes generated using PatchDock (Duhovny et al., 2002) and ~54 000 decoys for each of the complexes generated using ZDOCK (Pierce et al., 2011).

**2.5.3 Assessment criteria for SOAP-PP** Each model is assessed for accuracy based on root mean square deviation (RMSD) from the native structure, as used at CAPRI (Lensink et al., 2007). A docking model is considered *acceptable* if the ligand  $C_\alpha$  RMSD after superposition of the receptors is <10 Å or the interface  $C_\alpha$  RMSD is <4 Å. A docking model is of *medium* accuracy if ligand  $C_\alpha$  RMSD is <5 Å or interface  $C_\alpha$  RMSD is <2 Å. The success rate for SOAP-PP is the percentage of benchmark cases with at least one medium or acceptable accuracy model in the top  $N$  predictions.

**2.5.4 Learning set for SOAP-Loop** This set consists of 3838 native loop conformations of 4–20 residues and ~500 decoys for each loop generated using MODELLER (Fiser and Sali, 2003; Sali and Blundell, 1993). Loops were extracted from X-ray crystallography structures in the PDB using DSSP (Kabsch and Sander, 1983; Joosten et al., 2011). We only considered protein structures determined at a resolution better than 2 Å,  $R_{\text{free}}$  better than 0.25 and crystallized between pHs 6.5 and 7.5; no pair of source structures had sequence identity higher than 30%. Each loop has only standard residues, no missing non-hydrogen atoms, average atomic surface accessibility between 5 and 60%, no crystal contacts, no clashes with nearby atoms, no contacts with metal ligands and does not occur in the PLOP loop modeling decoy set (Jacobson et al., 2004).

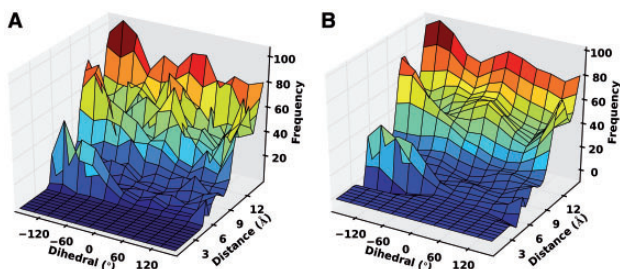
**2.5.5 Testing set for SOAP-Loop** This set consists of 833 native loop conformations of 4–12 residues and ~450 decoys for each loop generated using PLOP (Jacobson et al., 2004).

**2.5.6 Assessment criteria for SOAP-Loop** Each model is assessed for accuracy based on its main-chain RMSD to the native conformation, after superposition of all non-loop atoms ( $\text{RMSD}_{\text{global}}$ ) (Fiser et al., 2000); main-chain atoms include amide nitrogen,  $C_\alpha$ , as well as carbonyl carbon and oxygen. SOAP-Loop is assessed by the average  $\text{RMSD}_{\text{global}}$  of the top ranked model for each loop.

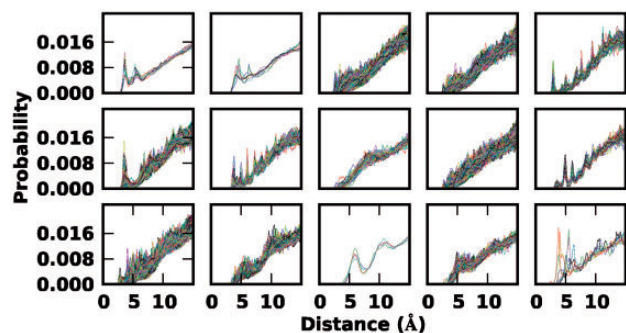
## 2.6 Recovery functions and functional forms

We estimate the recovery function  $g(f^c|Q_K)$  by optimizing the accuracy of the corresponding statistical potential on a benchmark of interest. To avoid overfitting, we assume either a single recovery function for all feature types or the same recovery function for a subset of similarly distributed feature types.

The set of recovery function forms  $G_f$  is different for distances, angles and accessibility: The recovery function for the atomic distance is modeled using one of three functional forms: (i)  $d^q$ , where  $d$  is distance and  $q$  is a constant (Zhou and Zhou, 2002); (ii) the ideal gas distribution in spheres with varying radii (Shen and Sali, 2006); and (iii) spliced cubic splines. For orientation, the recovery function is defined as the product of



**Fig. 3.** Distance and dihedral angle joint distribution between alanine  $N-C_\alpha$  and alanine  $O-C$ , when  $\alpha_1 \in [60^\circ, 90^\circ]$  and  $\alpha_2 \in [60^\circ, 90^\circ]$ . (A) Original distribution. (B) Smoothed distribution



**Fig. 4.** Distance distributions  $p(f^c|Q_k)$  for different atom pairs are clustered into 15 different groups. Each line represents a distance distribution from a pair of atoms of certain types. Each group has 6–8401 distributions. During k-mean clustering, the number of clusters was set to 20, resulting in 14 clusters with  $>5$  distributions and 6 clusters with  $<5$  distributions; the latter 6 clusters are grouped together (bottom right panel)

a recovery function for  $d$ ,  $\alpha_1$ ,  $\alpha_2$  and  $\psi$ , respectively. The recovery functions for angles  $\alpha_1$ ,  $\alpha_2$  and dihedral angle  $\psi$  are modeled using two different functional forms: (i) the feature distribution calculated using the ideal gas assumption and (ii) spliced cubic splines. For the relative atomic surface accessibility, the recovery function form is spliced cubic splines. Control points of cubic splines are defined by their  $x$  and  $y$  values. When searching for the best cubic spline recovery function, the  $x$  values of the control points are either fixed at discrete sampling values or inferred together with the  $y$  values.

To optimize the recovery functions, we need to balance minimizing noise and maximizing precision. Thus, for atomic distances, we clustered the distance distributions  $p(f^c|Q_k)$  for different atom type pairs using k-mean clustering and assumed that the pairs of atom types with similar distance distributions have a similar recovery function (Fig. 4).

## 2.7 Bayesian inference and model selection

A statistical potential is defined by four discrete input variables (the known protein structure subset  $\mathcal{K}$ , the feature type subset  $\mathcal{F}$ , the smoothing prior  $\mathcal{S}$  and the recovery function form  $\mathcal{G}_f$ ) and a vector of continuous input variables (the recovery function parameters  $\mathcal{G}_\theta$ ). We elected to define the best values for the four discrete variables are those that result in the most generalizable statistical potential, as judged by the Bayesian predictive densities (Vehtari and Lampinen, 2002), whereas the best values for the recovery function parameters are those that result in the most accurate statistical potential, as judged by a given benchmark. Because each of the five variables can be sampled at many values, enumeration of all combinations is not computationally feasible. Thus, the search for the best values is carried out in four stages, as follows.

First, irrespective of the final restrained feature  $\mathcal{F}$ , we begin with the atomic distance and a single recovery function for all atom type pairs. The optimal values of the discrete variables ( $\mathcal{F}, \mathcal{K}, \mathcal{S}, \mathcal{G}_f$ ) are found by an iterative discrete search:

- (1) Choose an arbitrary starting value for each variable out of their possible value sets  $\{\mathbb{F}, \mathbb{K}, \mathbb{S}, \mathbb{G}_f\}$  (Supplementary Table S1 and S2).
- (2) For each variable, choose the best value and eliminate the worst value in the value set using Bayesian model selection based on Bayesian predictive densities (Vehtari and Lampinen, 2002). The Bayesian predictive density for each value is calculated with other variables fixed at their best previous values:

$$\prod_{\{t, v\}} \int p(D_v|\mathcal{F}, \mathcal{K}, \mathcal{S}, \mathcal{G}_f, \mathcal{G}_\theta) \cdot p(\mathcal{G}_\theta|\mathcal{F}, \mathcal{K}, \mathcal{S}, \mathcal{G}_f, D_t) d\mathcal{G}_\theta \quad (4)$$

where the learning decoys  $D$  are randomly separated multiple times into a training set  $D_t$  and a validation set  $D_v$ , from which the integrals are estimated using Monte Carlo sampling (Evans and Swartz, 2000).  $p(\mathcal{G}_\theta|\mathcal{F}, \mathcal{K}, \mathcal{S}, \mathcal{G}_f, D_t)$  is calculated following the Bayes rule:

$$p(\mathcal{G}_\theta|\mathcal{F}, \mathcal{K}, \mathcal{S}, \mathcal{G}_f, D_t) \propto p(D_t|\mathcal{F}, \mathcal{K}, \mathcal{S}, \mathcal{G}_f, \mathcal{G}_\theta) \cdot p(\mathcal{G}_\theta|\mathcal{G}_f) \quad (5)$$

here the likelihood  $p(D_t|\mathcal{F}, \mathcal{K}, \mathcal{S}, \mathcal{G}_f, \mathcal{G}_\theta)$  is a half-normal distribution whose corresponding normal distribution has the mean equal to the accuracy of an imaginary statistical potential generating scores that correlate perfectly with the decoy-native RMSD and the standard deviation computed by dividing the mean by the number of the cases in the training set  $D_t$ ; the prior  $p(\mathcal{G}_\theta|\mathcal{G}_f)$  is an informative prior defining a reasonable range for  $\mathcal{G}_\theta$ .

- (3) Repeat step 2 until the best values do not change.
- (4) Repeat five times steps 1–3 for different random initial values.
- (5) Keep the best performing variable values.

Second, keeping the optimal values from the previous step fixed, we find the optimal values for the feature type, smoothing length scale and the number of spline anchor points using the same 5-step iterative discrete search outlined earlier in the text.

Third, if the optimal spatial feature selected in the previous step is not orientation, we vary the number of recovery functions and the number of anchor points to optimize their values, again using the 5-step iterative discrete search.

Fourth, using the selected  $\{\mathcal{F}, \mathcal{K}, \mathcal{S}, \mathcal{G}_f\}$ , we infer the best recovery function parameter values  $\mathcal{G}_\theta$  by maximizing  $p(\mathcal{G}_\theta|\mathcal{F}, \mathcal{K}, \mathcal{S}, \mathcal{G}_f, D)$  (Equation 5). The optimized statistical potential is then calculated (Equation 2) and assessed on testing decoy sets.

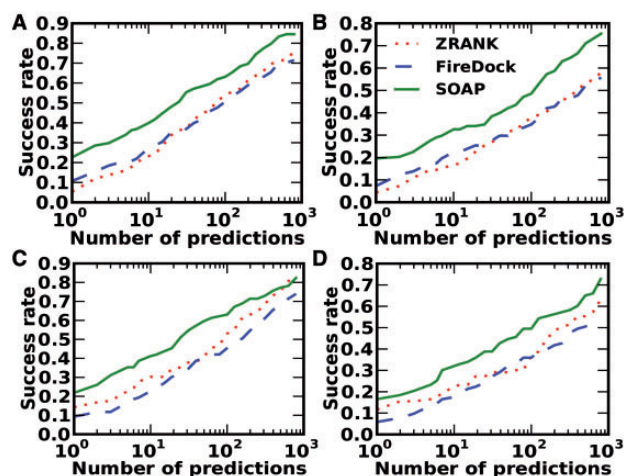
SOAP-PP and SOAP-Loop are available as part of MODELLER (<http://salilab.org/modeller>). All the training, learning, testing, decoys, benchmark sets and scripts are available at <http://salilab.org/soap>.

## 3 RESULTS

### 3.1 Scoring protein–protein interfaces

SOAP-PP is an atomic statistical potential for assessing a binary protein interface, computed with our Bayesian framework by optimizing its accuracy on the learning set for SOAP-PP (Supplementary Table S1).

Using the recovery function parameters optimized for 15 sets of training decoys (each set is randomly selected 50% of the learning set), the average top10 success rate (Section 2.5.3) is  $44.7 \pm 1.2\%$  on the sets of training decoys and  $38.4 \pm 1.7\%$  on the sets of validation decoys. The relatively small difference between the two success rates likely results from overfitting. To investigate overfitting, we increased the size of the training decoy set from 50 to 67% of the entire learning set of 176 proteins. As a result, the average top10 success rate on the training decoys decreased from 44.7 to 44.2%, but the average success rate on the validation decoys (the remaining 33% of the learning set) increased from 38.4 to 39.8%. This observation suggested that increasing the size of the training set may be an effective way of reducing overfitting (Murphy, 2012). Thus, we optimized SOAP-PP using the entire learning set of 176 proteins as the training set, even though this forces subsequent testing on the training protein sequences. To estimate the resulting overfitting, we calculated six optimized statistical potentials, each one of which was based on a training set that included a random subset of  $\sim 67\%$  of the learning set. Next, we tested these potentials on two testing sets: the

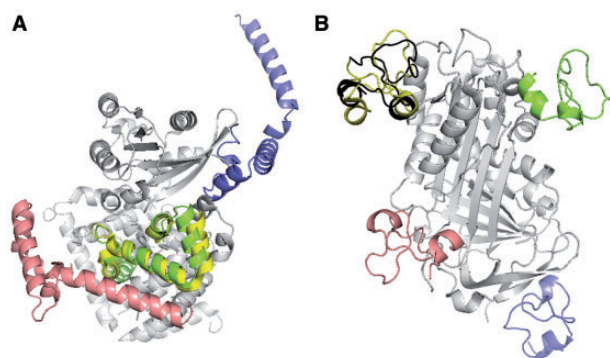


**Fig. 5.** Success rates of SOAP-PP, ZRANK and FireDock on the PatchDock and ZDOCK decoy sets. (A) Success rates on the PatchDock decoy set, where a success is defined as having an acceptable accuracy structure in the top  $N$  predictions ( $x$ -axis). (B) Success rates on the PatchDock decoy set for picking structures with medium accuracy. (C) Success rates on the ZDOCK decoy set for picking structures with acceptable accuracy. (D) Success rates on the ZDOCK decoy set for picking structures with medium accuracy

first set consisted only of the training proteins; the second set consisted of the remaining learning proteins. The average top10 success rate for the PatchDock decoys is 41.1% and 38.6% for the first and second test set, respectively; for the ZDOCK decoys, the average top10 success rate is 40.0 and 38.9% for the first and second test set, respectively. Therefore, given that increasing the training set size reduces overfitting as shown previously, the accuracy of SOAP-PP estimated based on a completely different testing set is expected to be within 2.5% of the current estimate (later in the text).

SOAP-PP was assessed on the PatchDock (Schneidman-Duhovny *et al.*, 2012) and ZDOCK decoy sets (Pierce *et al.*, 2011) (Fig. 5). For PatchDock decoys, the top10 success rate of SOAP-PP is 40% (Fig. 5A) compared with 23% for ZRANK and 27% for FireDock. If only models of medium or better accuracy are considered, the top10 success rate is 33% for SOAP, 17% for ZRANK and 23% for FireDock (Fig. 5B). For ZDOCK decoys, the top10 success rate of SOAP-PP is 41% (Fig. 5C) compared with 30% for ZRANK and 22% for FireDock. If only models of medium or better accuracy are considered, the success rate is 32% for SOAP-PP, 22% for ZRANK and 17% for FireDock (Fig. 5D).

High accuracy of SOAP-PP can sometimes be attributed to the weaker short-distance repulsion (Fig. 6A) compared with ZRANK (Pierce and Weng, 2007) and FireDock (Andrusier *et al.*, 2007), both of which use a modified van der Waals repulsion term; thus, the clashes of the best sampled structure with a receptor are likely less penalized by SOAP than by ZRANK and FireDock. Although SOAP-PP is more successful than ZRANK and FireDock overall, picking near-native protein–protein complex models out of decoys remains a hard problem



**Fig. 6.** Comparison of the top ranked, best sampled and native configurations. (A) 2G77. (B) 1OC0. The receptor is shown in gray. The ligand is shown in the native configuration (yellow), the best sampled configuration (green for 2G77 and black for 1OC0) and the top ranked configuration by SOAP (green), FireDock (blue) and ZRANK (red)

(Fig. 5). For some cases, all three scoring functions perform badly, especially when the protein–protein interfaces are small and have poor shape complementarity (Fig. 6B).

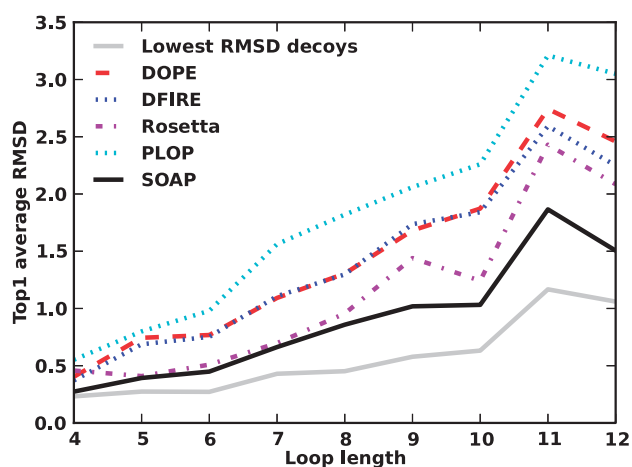
### 3.2 Scoring loops

SOAP-Loop is an atomic statistical potential for assessing protein loop conformations, computed with our Bayesian framework by optimizing its accuracy on the learning set for SOAP-Loop (Supplementary Table S2).

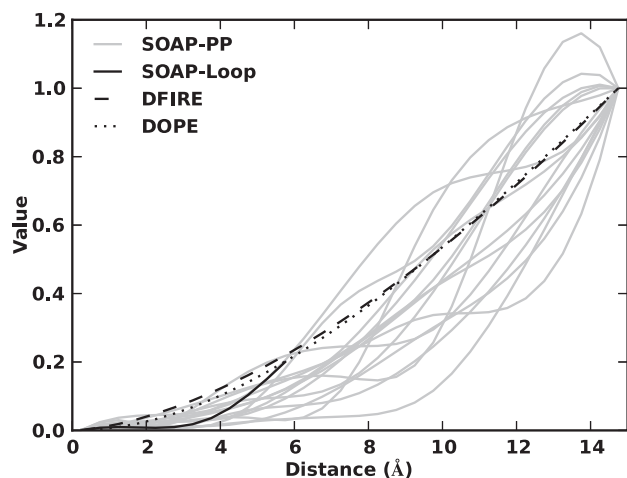
SOAP-Loop was assessed on the PLOP loop modeling decoy set (Jacobson *et al.*, 2004). We compare SOAP-Loop with DOPE (Shen and Sali, 2006), DFIRE (Zhang *et al.*, 2004), Rosetta 3.3 (Simons *et al.*, 1999) and PLOP 25.6 scoring functions (Jacobson *et al.*, 2004) (Fig. 7A). For short loops, SOAP-Loop and Rosetta perform similarly and better than the other tested scoring functions: the main-chain RMSD of SOAP-Loop's top ranked structure is close to that of the best decoy structure. For longer loops, the accuracy differences become larger. SOAP-Loop is still able to pick structures close to the best decoy structures: for 12-residue loops, the average main-chain RMSD of the best scored conformations by SOAP-Loop is 1.5 Å, close to the average RMSD of the best decoy conformations (1.2 Å) and significantly better than that by DOPE (2.5 Å), DFIRE (2.3 Å), Rosetta (2.1 Å) and PLOP scoring functions (3.0 Å). We note that this assessment should not be used to rank the PLOP scoring function because the decoy set used here was generated with PLOP. Thus, we further compare different scoring functions by their average all-atom RMSD values of the best scored conformations using our learning set for SOAP-Loop (Section 2.5.4 and Supplementary Table S3).

Although no testing protein occurs in the learning set, 11 pairs of testing-learning loops have the same sequence. Excluding these 11 loops from the testing set, the average RMSD of the top ranked loop by SOAP-Loop increases insignificantly from 0.895 Å to 0.897 Å; the average RMSD of the best decoy conformations also increases insignificantly from 0.566 Å to 0.567 Å.

The relative success of SOAP is attributed to the scoring of the orientation instead of distance and the use of the recovery functions instead of a reference state (Fig. 8). However, SOAP-Loop



**Fig. 7.** Accuracy of SOAP-Loop. The average main-chain RMSD of top ranked structures by DOPE, DFIRE, Rosetta, PLOP and SOAP-Loop on PLOP loop modeling decoys. The average RMSD of the most accurate conformations sampled by PLOP is plotted by a dash-dotted line

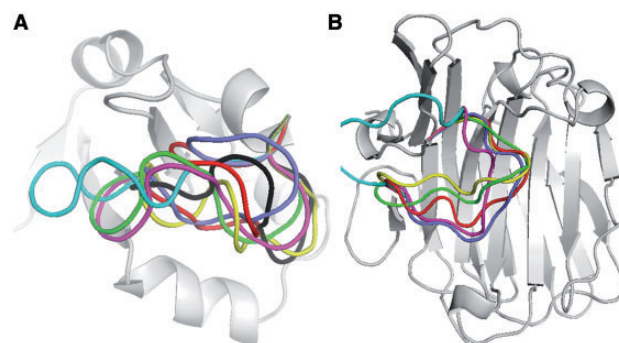


**Fig. 8.** Recovery functions for SOAP-PP and SOAP-Loop are compared with DOPE and DFIRE's reference states

still fails to identify the best-sampled conformation in some cases. For a loop in 1CYO, for example, the failure can be attributed to the lack of a sufficiently native conformation among the tested conformations and the absence of significant interactions between the loop and the rest of the protein (Fig. 9A). It is also possible that some interactions, such as long-range interactions, are not treated accurately by any scoring function, indicating the need for further development of the theory of statistical potentials.

## 4 DISCUSSION

We developed a Bayesian approach to optimizing statistical potentials based on probability theory and without recourse to questionable statistical mechanical assumptions and approximations. We also applied this approach to calculate optimized



**Fig. 9.** Comparison of the top ranked, best sampled and native configurations. (A) 1CYO. (B) 2AYH. The native structure is shown in light gray. The loop is shown in the native configuration (yellow), the best sampled configuration (black for 1CYO and green for 2AYH) and the top ranked configuration by SOAP (green), DOPE (blue), DFIRE (red), Rosetta (magenta) and PLOP (light blue)

statistical potentials for assessing protein interactions (SOAP-PP) and loops (SOAP-Loop). These two statistical potentials perform better than others in their class. For PatchDock and ZDOCK decoys, the top10 success rate of SOAP-PP is >10% higher than that of FireDock and ZRANK (Fig. 5). For 12-residue loops in the PLOP benchmark, the average main-chain RMSD of the best scored conformations by SOAP-Loop is 1.5 Å, close to the average RMSD of the best sampled conformations (1.2 Å) and significantly better than that from DOPE (2.5 Å), DFIRE (2.3 Å), Rosetta (2.1 Å) and PLOP scoring functions (3.0 Å) (Fig. 7). The relative accuracy of SOAP-PP and SOAP-Loop results primarily from normalizing the raw distributions by the recovery functions instead of a reference state, restraining of orientation instead of only distance and thoroughly optimizing parameter values while avoiding overfitting.

Next, we discuss three points in turn. First, we describe our recovery functions and compare them with the reference states used for other statistical potentials. Second, we discuss the importance of restraining orientation and using covalent separation as an independent variable. Finally, we conclude by commenting on future improvements of our Bayesian approach and its applications.

### 4.1 Cubic splines as a recovery function form

A key difference between statistical potentials is the definition of their reference states, which are often derived by assuming that the PDB provides a Boltzmann ensemble of structural features (Sippl, 1990). Here, we replace the reference state by data-driven recovery functions, defined self-consistently without recourse to these questionable statistical mechanical assumptions (Finkelstein *et al.*, 1995; Shen and Sali, 2006). In an extreme case, we use cubic splines to compute optimal recovery functions, relying on Bayesian inference to obtain parameter values that result in the most accurate statistical potential given a benchmark.

The use of splines as recovery functions is motivated by a qualitative analysis of the recovery function (Supplementary Equation S2). The distribution  $p(f^{(m)}|Q_K)$  of a single feature

$f^{c(m)}$  is the product of the restraint on  $f^{c(m)}$  and an integral involving the restraints on  $Q_K$ 's other features (i.e. the environment restraint). Then, the recovery function  $g(f^c|Q_K)$  is the distribution of feature type  $c$  in structure set  $\mathcal{K}$  resulting from the environmental restraints alone (Supplementary Equation S2). We now discuss three implications of this perspective.

First, if we assume that atoms are placed randomly within the protein shell, a recovery function will be similar to the DFIRE and DOPE reference states based on the ideal gas assumption (Shen and Sali, 2006; Zhou and Zhou, 2002).

Second, using the distance  $d$  between atoms A and C in Figure 2 as an example, the environment restraint on  $d$  is a consequence of the restraints on distances between A–D, C–B and B–D as well as the bonds between A–B and C–D. The restraints on A–D, C–B and B–D distances have short-range repulsion components. Thus, the environment restraint on the distance A–C will include an effective short-range repulsion. This qualitative analysis is consistent with the observed recovery functions for SOAP-PP and SOAP-Loop, which all have lower values at short distances than the DOPE reference state based on the ideal gas assumption (Fig. 8).

Finally, the recovery functions for different feature types can vary, because of their different environments, as observed for the recovery functions for 15 clusters of atom type pairs used in SOAP-PP (Fig. 8).

Although splines can mimic almost any smooth function given a sufficient number of anchor points, its flexibility could also lead to overfitting; moreover, a large number of anchor points could lead to oscillations (Fig. 8). Although our Bayesian model selection method helps with the generalizability of the optimized cubic spline (Vehtari and Lampinen, 2002), it is conceivable that applying Bayesian model selection to a less flexible but appropriate functional form will result in a more accurate and general statistical potential than that based on splines.

## 4.2 Spatial and sequence features

Our orientation restraints score a spatial relationship between two sets of atoms in more detail than distance restraints alone, and should be particularly useful for scoring spatial relationships between polar atoms, especially for hydrogen bond donors and acceptors. In fact, the relative accuracy of SOAP-Loop can be attributed to the use of orientation and recovery functions instead of distance and reference state, respectively (Supplementary Table S1). However, using orientation did not result in a better statistical potential for ranking protein interfaces (Supplementary Table S2). Although we may not have found the globally optimal statistical potential for orientation, a more likely reason is insufficient accuracy of the tested conformations produced by rigid docking.

Covalent separation is another important factor affecting the accuracy of the derived statistical potentials. Surprisingly, for ranking protein interfaces, statistical potentials derived from intra-chain non-local atom pairs (bond separation  $>9$ ) work better than statistical potentials derived from inter-chain atom pairs (chain separation = 1) (Supplementary Table S1). A likely reason is that many protein interfaces in the PDB result from crystal contacts that do not reflect interfaces between proteins in solution (Carugo and Argos, 1997; Krissinel, 2010). In the

future, a better statistical potential for ranking protein interfaces might be obtained if only true biological interfaces from PDB are used.

## 4.3 Bayesian inference

Statistical potentials can be derived for many different values of the input variables, with little or no *a priori* reasons to choose one set of values over the others. The Bayesian model selection based on Bayesian predictive densities provides a statistically rigorous way of choosing the values that result in most generalizable statistical potentials (Vehtari and Lampinen, 2002). However, one limitation of this method is that the calculation of predictive densities is computational intensive, often requiring more than tens of thousands of evaluations of the statistical potential on the benchmark. Thus, such calculations are not always practical. Fortunately, increases in the available computer power will enable us to find more accurate statistical potentials in an increasingly larger parameter space in the future. Another approach to improving the search for optimal parameter values is to use physically motivated feature types, functional forms and allowed value ranges.

In principle, normalizing the feature distributions by recovery functions to obtain a statistical potential (Equation 2) is not necessary. Instead, we could use parametric (e.g. the mathematical functional forms used in molecular mechanics force fields) or non-parametric functions to represent the statistical potential and directly infer the optimal statistical potential by its accuracy on a benchmark of interest. However, this approach might not provide an accurate statistical potential in practice because of the large number of parameters whose values would need to be optimized.

Our method for smoothing feature distributions is a generalization of the two related methods used in calculating statistical potentials (Sippl, 1990) and homology restraints (Sali and Blundell, 1993). Both methods are equivalent to our Bayesian smoothing method with a diagonal covariance matrix as the smoothing prior. Their prior distribution is equivalent to the mean of our prior  $\mathcal{S}$ , whereas the weights on their prior distributions are defined by the standard deviation in our covariance matrix.

In conclusion, our Bayesian framework can be applied to derive an optimized statistical potential for many other kinds of modeling problems for which sample structures are available, thus affording better leverage of the experimentally determined protein structures. Examples include membrane protein topology and complexes of proteins with small molecules or peptides.

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