Comparing pairwise-additive and many-body Generalized Born models for acid/base calculations and protein design

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Abstract

Generalized Born (GB) solvent models are common in acid/base calculations and protein design. With GB, the interaction between a pair of solute atoms depends on the shape of the protein/solvent boundary and therefore the positions of all solute atoms, so that GB is a many-body potential. For compute-intensive applications, the model is often simplified further, by introducing a mean, native-like protein/solvent boundary, which removes the many-body property. We propose a method for both acid/base calculations and protein design that uses Monte Carlo simulations in which side chains can explore rotamers, bind/release protons, or mutate. The solvent is treated with GB. The fluctuating protein/solvent dielectric boundary is treated in a way that is numerically exact (within the GB framework), in contrast to a mean boundary. Its originality is that it captures the many-body character while retaining the residue-pairwise complexity given by a fixed boundary. The new "Fluctuating Dielectric Boundary" method (FDB) is implemented in the Proteus protein design software. It yields a slight but systematic improvement for acid/base constants in nine proteins and a significant improvement for the computational design of three PDZ domains. It eliminates a significant source of model uncertainty which will facilitate the analysis of other model limitations.

Key words: protein electrostatics, generalized Born model, Proteus program, molecular mechanics, implicit solvent

1 Introduction

Continuum electrostatic treatments of aqueous solvent [1] are an important ingredient of many biophysical models [2–5]. They describe a protein solute as a low dielectric medium, embedded in a high dielectric solvent. Atomic charges within the protein are shielded by the solvent, and the extent of shielding depends on the shape of the protein-solvent boundary [6, 7]. As a result, the interaction between two protein atoms depends on the position of the other protein atoms. Thus, to compute the forces and energy one must solve a many-body problem [8–10].

Generalized Born (GB) models are simplified treatments that preserve much of the physics of continuum electrostatics [11–13]. In particular, the GB interaction between two protein atoms depends on their "solvation radii", which approximate their distance from solvent and depend on the positions of the other protein atoms [14–17]. In simulations, the many-body nature of GB affects computational efficiency. Therefore, in some applications, an additional approximation is introduced that makes the GB model "pairwise additive". The GB energy then takes the form of a sum over pairs of solute atoms, similar to the Coulomb and van der Waals energies. One way to do this is to assume the atomic solvation radii are constant, and can be computed ahead of time using a consensus structure, such as a crystal structure. With some GB variants, if the solvation radii are held constant, the computational cost of a molecular dynamics simulation is divided by two.

Two applications that require very efficient energy calculations are acid/base calculations and computational protein design (CPD). CPD starts from a 3D structural model and explores a vast space of sequences and conformations to identify protein variants that have certain predefined properties, such as stability or ligand binding. Acid/base properties of proteins are important for their structure, stability, and function [18, 19]. Acid/base calculations are related to CPD, since proton binding/unbinding to a side chain can be treated formally as a kind of mutation. Indeed, pK_a calculations have been used recently to test CPD model and software packages [20–22].

Several recent models for CPD [23–31] and acid/base problems [32–44] combine atomistic simulations of the protein motions, a Poisson-Boltzmann (PB) or GB solvent, and a Monte Carlo (MC) exploration of side chain mutations and/or protonation. In these models, the protein backbone is held fixed, and its motions described implicitly, through the protein dielectric constant [45]. Side chains explore a discrete set of preferred rotamers, so that conformational space is finite. The energy is approximated as a sum of pairwise

terms that involve just one or two amino acids; i.e., it is pairwise additive [31, 46–50]. With a discrete conformational space and a pairwise additive energy, interaction energies can be computed ahead of time and stored in a lookup table, called the *energy matrix*. Conformational space is then explored through a Monte Carlo simulation, which includes side chain protonation changes and/or mutations.

The pairwise additivity approximation affects surface area calculations, but its main effect is on the GB energy term [22, 50]. In previous work, to make the GB term pairwise additive, we and others have assumed that for any given protein conformation, each amino acid pair experiences a mean, effective dielectric environment that is native-like and remains constant over time. We refer to this as the "Native Environment Approximation", or NEA. Such pairwise additivity approximations for the solvent are routine in CPD, and are common for acid/base calculations [34, 51] and problems like protein:ligand docking [52, 53]. We recently reported an extensive pK_a benchmark test that used the NEA [22].

Here, we go beyond the NEA level of theory and use a method that takes into account rigorously the dielectric environment of each amino acid and its fluctuations over time [54]. In other words, the method preserves the many-body property of GB and continuum electrostatics. The originality of the method is that it also preserves the pairwise complexity of the energy matrix and the MC calculation. We refer to it as the "Fluctuating Dielectric Boundary" method, or FDB. It exploits the fact that with GB, the dielectric environment of a residue pair is completely characterized by a small set of "atomic solvation radii". These radii are themselves pairwise sums over the protein atoms [14, 16]. Our method involves two steps. First, we compute an average solvation radius B_I for each residue I, that will be used for the calculation of residue–residue interaction energies. This averaging leads to a GB variant we call "Residue-GB" [54]. Second, we express the GB interaction energy of each residue pair I, J as a power series relative to the B_I , B_J . Since the residue solvation radii are themselves sums over residue pairs, we thus reduce the GB energy calculation to a pairwise complexity, as was the case with NEA. The new method requires additional bookkeeping and overhead during both the pre-calculation of the energy matrix and the MC simulation, leading to an increased cost. For the energy matrix, the cost increase of FDB compared to the NEA is only a few percent. For the MC simulation, the computational cost is increased by a factor of about 5-10, depending on the protein. Including the cost of the energy matrix, the overall cost increase with FDB, compared to the NEA is about a factor of four. The method is implemented in our CPD package Proteus [31, 47, 55].

Below, we describe briefly the implementation of the new FDB method. To test it, we first performed an extensive pK_a benchmark, including 9 proteins and 149 titratable groups, for which the NEA and FDB methods were compared. One of the proteins was hemoglobin, for which we also computed the Bohr effect: the number of protons that bind when the protein switches from its deoxy to its oxy form, which is of physiological interest [56, 57]. Overall, the FDB performance was slightly but systematically better than NEA, with an rms deviation of 1.1 between the computed and experimental pKa's (vs. 1.2 with NEA), a higher correlation with experiment, and fewer large errors. The alkaline Bohr effect in hemoglobin is fairly well reproduced with FDB, and the overall Bohr effect is superior to the NEA result. Although the FDB performance is not as good as the empirical PropKa tool [58], it is comparable for groups with significantly shifted pK_a 's and it provides a more physically-transparent interpretation than an empirical model.

As a second test, we did protein design calculations with Proteus for three PDZ proteins, and compared the NEA with FDB. PDZ domains are small, globular domains of around 90 amino acids that establish protein-protein interaction networks in the cell [59–64]. They were redesigned through MC simulations where all the amino acids (except Gly, Pro) were allowed to mutate freely. The simulations used a low and physically-realistic protein dielectric constant of four. The lowest-energy Proteus sequences were compared to natural sequences from the Pfam database and to sequences generated by the widely-used Rosetta package [23, 65–67]. The similarity between the Proteus and Pfam sequences was comparable to the similarity between the Pfam sequences themselves. It was also comparable to the Pfam similarity of sequences produced by Rosetta. Compared to the NEA method, there was a significant improvement. Overall, the rigorous treatment of GB and its many body character leads to improved quality for both sequence design and acid/base constants; it eliminates a significant source of model uncertainty, and thus facilitates the interpretation of the model predictions.

2 Computational methods

2.1 Fluctuating Dielectric Boundary method

In GB models, the electrostatic energy includes both a direct, Coulomb term and a contribution from the solvent, polarized by the solute charges. Treating the protein and the solvent as two distinct, homogeneous, dielectric media, the total electrostatic energy has

the form:

$$E^{\text{elec}} = E^{\text{Coul}} + \Delta G^{\text{solv}}$$

$$= \frac{1}{2} \sum_{i \neq j} \frac{q_i q_j}{r_{ij}} + \frac{1}{2} \sum_{ij} g_{ij}, \qquad (1)$$

where the sums are over all pairs of protein charges and the second sum includes diagonal terms, i = j. The term g_{ij} represents a GB interaction or screening energy. In the most common GB model [11], this term is approximated by

$$g_{ij} = g(r_i, r_j) = \tau q_i q_j \left(r_{ij}^2 + b_i b_j \exp[-r_{ij}^2/4b_i b_j] \right)^{-1/2}, \tag{2}$$

where $\tau = \frac{1}{\epsilon_w} - \frac{1}{\epsilon_p}$, ϵ_w is the solvent dielectric constant, ϵ_P is the protein dielectric constant and b_i , b_j are the solvation radii of the atoms i and j. The solvation radii are approximated by a simple, analytical function of the positions of all the solute atoms: $b_i = b_i(r_1, r_2, ..., r_N)$. Different GB variants use different functional forms [11, 14–17, 68, 69]. The essential point is that in most variants, including the one considered here, the self-energy takes the form of a sum over pairs of atoms.

With the new, FDB method, we modify the GB formulation to employ "residue" solvation radii, leading to a "Residue" GB model. We define a self-energy contribution corresponding to a particular residue pair I, J by the expression

$$E_{IJ}^{\text{self}} = \sum_{i \in I} E_{ij}^{\text{self}}, \tag{3}$$

where the sum extends over atom pairs where i belongs to residue I and j to residue J. The self-energy of residue I can be written

$$E_I^{\text{self}} = \sum_I E_{IJ}^{\text{self}} \tag{4}$$

and the total self-energy can be written

$$E^{\text{self}} = \sum_{I} E_{I}^{\text{self}}.$$
 (5)

This residue-decomposition of the self-energy is exact within the GB model. We then define the residue solvation radius B_I by the relation

$$E_I^{\text{self}} \stackrel{\text{def}}{=} \tau \sum_{i \in I} \frac{q_i^2}{2B_I}.$$
 (6)

We also have

$$\left(\sum_{i\in I} q_i^2\right) \frac{1}{B_I} = \sum_{i\in I} \frac{q_i^2}{b_i}.$$
 (7)

Thus, B_I is a harmonic average over the b_i , $i \in I$, weighted by the squared charges.

We now define the contribution g_{IJ} of residues I and J to the total screening energy ΔG^{solv} :

$$g_{IJ} = \sum_{i \in I, j \in J} \tau q_i q_j \left(r_{ij}^2 + B_I B_J \exp[-r_{ij}^2 / 4B_I B_J] \right)^{-1/2}$$
 (8)

For I = J, the double summation in Eq. (8) is actually restricted to pairs of distinct atoms, $i \neq j$. We note that, for fixed interatomic distances r_{ij} , $g_{IJ}(B_IB_J)$ is a slowly varying function of $B \equiv B_IB_J$. This dependency can be approximated by a low-order power expansion [54]:

$$g_{IJ}(B) \approx c_1^{IJ} + c_2^{IJ}B + c_3^{IJ}B^2 + c_4^{IJ}B^{-1/2} + c_5^{IJ}B^{-3/2}$$
 (9)

The coefficients c_n^{IJ} can be pre-computed and stored in the energy matrix. To keep the notations simple, their dependency on the particular rotamer combination r_I , r_J is not indicated explicitly. The quantities B_I and B_J can be obtained from residue pair contributions stored in the energy matrix. Thus, with (9), the Fluctuating Dielectric Boundary method only involves quantities that depend on residue pairs.

2.2 FDB implementation details

For all the proteins except hemoglobin (Hb), all the residue pairs were treated using the new, FDB method. For Hb, due to its size, the FDB method was applied to all pairs belonging to one of the Hb subunits, say α_1 , and to interactions between α_1 residues and other residues less than 6 Å away. For the other pairs, we used the NEA. For each FDB residue pair and all possible rotamer combinations, the GB interaction energy was fitted to the power expansion (9) in the range B=1–150 Å² using our locally-modified version of the Xplor program, which is itself part of the Proteus package [31]. The code was based on the general linear fit subroutine LFIT from Numerical Recipes [54, 70]. The fit is controlled at the level of the Xplor script language [71] by a statement of the form: pick gbfit <selection1> <selection2>, which computes the GB interaction energy between two groups of atoms R1, R2 defined by the two selections, which occupy a specific conformation. The individual solvation radii of R1 and R2 are not computed from the protein structure but are defined by the relation $B_{R1}B_{R2} = B$. Xplor performs the fit

and stores the fitting coefficients in the script variables coef1, ..., coef5, which can be printed out by a script command, e.g., display coef1. In the energy matrix, this information is stored along with the other interaction energy terms. The contribution of each residue pair to the GB self energy is also stored as a separate item in the energy matrix. The matrix entry for a pair I, J and a particular rotamer combination r_I , r_J is shown in Fig. 1.

With the NEA method, at each step t of the Monte Carlo simulation, if a residue I is displaced, the resulting energy change $\Delta E(t)$ can be computed from energy matrix elements of the form E_{IJ} . With FDB, the solvation radii change over time and so additional operations are needed:

- 1. Throughout the trajectory, we maintain an up-to-date list of all the residue solvation radii B_I , whose values fluctuate over the trajectory. This is possible since the GB self energy information is available in the matrix. At each MC step, B_I is only updated if a residue close to residue I (based on a neighbor list built ahead of time) is displaced or mutated.
- 2. At each MC trial move, if a solvation radius B_I changes, then residue I will contribute to the energy change $\Delta E(t)$, since its GB interaction energies g_{IJ} with all other residues J are affected. In fact, the contributions to $\Delta E(t)$ that result from the change in B_I are only computed for residues J within a certain cutoff distance of I. These J values are read out of a second neighbor list, built ahead of time, based on the size of the fitting coefficients c_n^{IJ} (Eq. 9): small coefficients indicate distant neighbors. For each neighbor J, the appropriate (rotamer-dependent) g_{IJ} value is obtained from the product $B_I B_J$ and the fitting coefficients c_n^{IJ} , n = 1, ..., 5, via Eq. 9.
- 3. At regular intervals (about every 1000 MC steps), the entire energy and all the solvation radii are recomputed from scratch, to avoid propagation of numerical errors.

With these implementation choices, the FDB method is efficient. For the proteins BPTI, barnase, and lysozyme, using FDB for the entire protein, the MC simulation times for a single pH value were 25, 86, and 107 minutes, respectively. This represents a cost increase, relative to NEA, by a factor of 8, 17, and 17, respectively. For Hb (125721 residue pairs), we applied FDB to all the pairs where I and J both belong to one subunit or its neighborhood (groups from other subunits less than 6 Å away); we applied NEA to all

other pairs. The simulation time per pH value was 30 minutes, which is four times longer than a calculation using NEA for all pairs. The MC simulations at different pH values can be done in parallel on multiple compute nodes. These simulation times can be compared to the time for energy matrix calculation. For BPTI, where we compute interactions for 861 pairs of residues (with multiple rotamers each), the whole matrix calculation on a 20-core desktop machine (Intel Xeon, 2.10 Ghz) required around 37 minutes with NEA and 39 minutes with FDB. For hemoglobin, where there are 125721 residue pairs, the matrix calculation took about 6 hours on a cluster of 100 cores (Intel Xeon, 2.67 Ghz). For BPTI, barnase, and lysozyme, the matrix compute time on 100 cores is about 0.5 hours. Overall, for the three small proteins, the total cost of a full pH scan increases from about 0.5 hours to about two hours. For Hb, the cost increase is much smaller because the matrix part dominates.

2.3 Monte Carlo simulations at constant pH

The framework for constant-pH Monte Carlo has been described several times [22, 34, 37–40]. We consider a dilute solution of the protein of interest, with a constant volume V and temperature T, closed with respect to protein and water, but open with respect to protons. If we compare two states that differ by the addition of a proton to a specific titratable side chain, with the protein in a given conformational state J, the ratio of Boltzmann probabilities has the form:

$$\frac{P_J(N+1)}{P_J(N)} = e^{-\beta(E_J(N+1) - E_J(N))} e^{\beta\mu} = e^{-\beta(E_J(N+1) - E_J(N)) - 2.303 \,\text{pH} + \beta\mu^0}$$
(10)

The index J identifies protein the conformation (rotamers) but also the specific location of all the protons bound to the protein; N is the number of bound protons, and μ is the chemical potential of the proton, which depends on the pH. The last equality uses the relation between μ and the pH: $\beta\mu = \beta\mu^0 - 2.303$ pH, where μ^0 is the proton chemical potential in aqueous solution in the standard state at pH = 0. For simplicity, in (10), we have kept the same index J for the two states, even though the N+1 state has an extra proton; this notation emphasizes that the protein conformation is otherwise unchanged, only a proton has been added to a particular location. The Boltzmann probability distribution in Eq. (10) is readily sampled using the standard, Metropolis, Monte Carlo algorithm [72].

Here, proton binding is described within a classical mechanical, molecular mechanics framework [2, 73–78]. The titrating protons, like the other atoms in the system, are

treated as classical mechanical particles, bearing a partial charge, and interacting with the other atoms through force field terms [79]. The proton's unit charge is distributed among the atoms of the titrating side chain. The classical mechanical formalism cannot describe bond formation, so we use a thermodynamic cycle [73], where the two horizontal legs represent the protonation of a titratable residue in the protein environment and in solution, respectively. In effect, when we remove a proton from the protein, we add it to a corresponding model compound in solution. The protonation free energy difference between the two reactions is computed with the force field, assuming that quantum mechanical effects associated with bond breaking cancel out. Thus, we actually obtain the shift $\Delta p K_a$ of each residue's pK_a, compared to the model compound in solution (pK^{model}_a):

$$pK_a^{\text{protein}} = pK_a^{\text{model}} + \Delta pK_a$$
 (11)

$$\Delta \Delta G^{\circ} = \Delta G^{\circ}_{\text{protein}} - \Delta G^{\circ}_{\text{model}}$$
$$= -2.303 kT \Delta p K_a \tag{12}$$

We can express the fraction x of the protonated species as a function of the standard protonation free energy and the pH, through the well-known titration curve:

$$x = \frac{1}{1 + 10^{(pH - pK_a)}}\tag{13}$$

The pK_a represents the pH at the inflection point of this curve, where the populations of the protonated and deprotonated states are equal.

2.4 Energy function and energy matrix

To describe each system, we used the Amber ff99SB force field [79, 80], combined with a GB variant whose parameters were optimized earlier [46], that we call GB/HCT [14, 81]. The GB energy term is computed with either the NEA or the FDB method. For the protein dielectric constant, we compared two values: 4 or 8. The solvent dielectric constant was set to 80. The energy also included a solvent accessible surface area term [46, 48]. The contact areas between each side chain pair, and between each side chain and the backbone were computed by the Lee and Richards method [82], using a probe sphere of radius 1.5 Å. The atomic surfaces were multiplied by empirical atomic solvation parameters, which describe the tendencies of various atom types to be buried or exposed, and were optimized to give good results for the stability changes due to a large collection of point mutants in various proteins: alkane = -5, polar = -8, ionic = -9, aromatic = -12, hydrogen = 0

(in cal/mol/Å²). Negative values indicate a preference to be solvent-exposed. To avoid overcounting of surface burial, for residue pairs involving at least one buried side chain, the surface energy was scaled by 0.65 [22, 46, 47, 50].

Interaction energies between all residue pairs in each protein were computed and stored in an energy matrix, allowing for all their possible rotamers and protonation states. The backbone was kept fixed as well as prolines and cysteines in disulfide bonds. In addition, to alleviate bad steric contacts due to the rotamer approximation, each pair was energy-minimized through 30 conjugate-gradient steps, in the presence of the fixed backbone and in the absence of the other side chains. During the minimization, each side chain torsional angle was subjected to a flat-bottomed harmonic restraint with a width of $\pm 5^{\circ}$ and a force constant of 200 kcal/mol/rad². Calculations were done with a locally-modified version of Xplor [71], which is part of our in-house Proteus package, developed for computational protein design [31, 47]. In addition to the modified Xplor, Proteus is made up of a large and sophisticated set of scripts in the Xplor command language, C code for MC exploration, and a set of perl and shell scripts.

2.5 Acid/base model compounds

Each titratable side chain has an associated model compound, which depends on the side chain type X. Here, we use the molecule 2N-acetyl-1N-methyl-X-1-amide, which consists of the backbone and side chain of X, capped with two blocking methyl groups [77]. X can designate either the protonated or deprotonated side chain form. We consider the titratable side chains Asp, non-disulfide Cys, Glu, His, Lys, and Tyr. The only titratable Cys residues in our test proteins are in hemoglobin. Each titratable side chain is assigned a "reference" free energy, $G_X^{\rm ref}$, which represents the free energy of the model compound in solution, estimated with the force field and including a pH-dependent term. The important quantities are the differences between the two or three protonation states of each model compound. These free energy differences can be expressed in the form:

$$\Delta G_X^{\text{ref}} = \Delta E_X^{\text{ref}} - kT \ln \frac{n_{XH}}{n_X} + \delta g_X^{\text{conf}} - 2.303 \, kT (pH - pK_a^X)$$
 (14)

Here, $\Delta E_X^{\rm ref}$ is the mean energy difference between the protonated and deprotonated states of X, averaged over a Monte Carlo simulation; n_{XH} and n_X are the rotamer numbers of the two states; pK_a^X is the acid/base constant of the model compound, taken from experiment [37, 83]; the rightmost term is the pH-dependent term, and $\delta g_X^{\rm conf}$ is the remaining free

energy. This term would be zero if all the rotamers of each protonation state had equal energies and were equally populated. The pK_a^X values were as follows: Asp, 3.9; Glu, 4.3; Cys, 8.6; His, 6.5 (N_{\delta} form) or 7.0 (N_{\epsilon} form); Lys, 10.4; Tyr, 9.8. By construction, titrating each model compound with Proteus using the reference free energies given by (14) should give back the experimental pK_a . This condition determines the value of $\delta g_X^{\rm conf}$. Numerical values were estimated by running a Monte Carlo simulation of each model compound with Proteus, at a pH equal to its experimental pK_a and adjusting $\delta g_X^{\rm conf}$ so that the two protonation states were equally populated. The values are reported in Table 2.

2.6 Monte Carlo protocol for acid/base calculations

The MC simulation starts from a random choice of protonation states and rotamers, using the mt19937 random number generator from the GNU Software Library. At each step, a move is chosen randomly and subjected to the Metropolis test [22, 72, 84]. Possible moves include changes of one or two rotamers, with or without an associated change of the protonation state. Move selection probabilities were 0.42 (single rotamer change), 0.08 (single protonation change), 0.25 (two rotamer changes), and 0.25 (two protonation changes). For two-position moves, the first position was chosen randomly and the second was chosen among positions that had at least one conformation where the unsigned interaction with the first side chain was 3 kcal/mol or more. To determine the pK_a's of all titrating residues, we scanned a pH range from 0 to 15, in steps of 0.5 units. The MC run at each pH value consisted of 10 million steps at room temperature (except myoglobin, which used 20 million steps, properly sampled the behavior of a coupled side chain pair). The quality of the MC sampling was tested for the deoxy form of HbA by performing scans with 5, 10 and 20 million steps per pH value. These runs yielded pK_a values within 0.2 units and Hill coefficients (see below) within 0.1.

2.7 Titration curves: fitting to the modified Hill equation

From each MC simulation, we computed the fractional occupancy x_i of the protonated state for each titrating site i at the given pH. After a full pH scan (300 million MC steps), we obtained the titration curve $x_i(pH)$ of each group. Each curve was fitted to a sigmoidal function, described by the modified Hill equation:

$$x = \frac{[AH]}{[A^-] + [AH]} = \frac{1}{1 + 10^{n(pH - pK_a)}}$$
 (15)

The exponent n is called the Hill coefficient; it is proportional to the maximum slope of the curve, which occurs at the inflection point. It is a measure of cooperativity, due to multiple, interacting, titrating sites in the protein [85]. Values close to 1 indicate non-cooperative proton binding. To determine the inflection point and the pK_a , we used a simple, iterative, grid search with focusing, implemented in a perl script. Both n and pK_a were chosen to minimize the mean square deviation between the fitted curve and the MC populations x(pH). This led to unbiased estimates of the pK_a 's when fitting to data generated by adding gaussian noise to the theoretical populations (15), using Mathematica [86]. The standard deviations of the fitted pK_a 's (over 10000 noise samples) were within 20% of that of the noise itself (0.1 or 0.05 pH units).

2.8 Protein setup for acid/base calculations

The test proteins are listed in Table 1. Staphylococcal nuclease (SNase) refers to a hyperstable mutant known as " Δ +PHS" [87]. In most cases, the precise form of the protein was the same in the pK_a measurements and the X-ray structure, with all protein atoms well-defined in the PDB coordinate file. N- and C-terminal groups were added in all cases but three. For OMTKY3 and barnase, several atoms at the N-terminus are disordered in the crystal structure and were omitted from our computational model, so no N-terminal ammonium was added. For SNase, several residues at each terminus were disordered, and so no N-terminal or C-terminal groups were added. Hydrogen atoms were positioned with ideal stereochemistry using the Xplor hbuild commmand and subjected to 1000 steps of conjugate gradient minimization, with everything else kept fixed. Myglobin has a heme cofactor, modeled using published force field parameters [88]. The proximal His93, which coordinates the heme iron, was maintained in its deprotonated form and kept in its crystal conformation.

Human adult Haemoglobin (Hb) is an $(\alpha/\beta)_2$ tetramer formed of two α and two β subunits (141 and 146 residues, respectively). Each subunit binds a heme cofactor, to which it is linked by a proximal histidine (α His87; β His92). We considered two forms of Hb; one has a CO molecule bound to the heme iron; this form is representative of oxy Hb. The other, deoxy form has no bound oxygen or CO. Following Zheng et al [57], the atomic coordinates were taken from the high resolution (1.25 Å) X-ray structures of the deoxy (PDB 2DN2) and oxy forms (PDB 2DN3) [89]. The deoxy form was crystallized at T = 298 K, pH = 6.5 with ammonium sulfate and ammonium phosphate, and is thought [57] to provide a good representation of the solution structure. The oxy form was crystallized at

T=100~K, pH=6.7, with sodium phosphate, potassium phosphate and glycerol. The rms deviation between the oxy/deoxy crystal structures is 2.3 Å (nonhydrogen atoms). Crystal waters were removed and hydrogens added. For each protein subunit, we established covalent linkages between the N_{ϵ} atom of the proximal histidine and the iron atom at the heme center. Finally, the whole structure was subjected to 200 steps of Powell conjugate gradient minimization, without any restraints. During this step, all ionizable residues were assigned their standard protonation state (positive Lys; neutral Tyr, Cys, His; negative Asp, Glu). Histidine residues were assigned to the N_{δ} tautomer. The heme force field was the same as for myoglobin.

Energy matrix calculations were done for the entire tetramer with NEA. With FDB, two separate calculations were done, where either the α_1 or the β_1 subunit was treated with FDB, while the rest of the tetramer was treated with NEA. Although the heme itself contains two propionate groups that can titrate, in principle, the force field does not include a description of the protonated state, and rotamer libraries for the propionate groups are not available. Therefore, for simplicity, we kept the whole heme and the proximal linked histidine fixed (like the backbone), with the propionates in their ionized, deprotonated state. Because of this simplified model, we do not report the computed titration behavior of nearby α H45, since it donates a hydrogen bond to one of the heme propionates. The experimental pK_a reported for this histidine [90] may also be questionable. The reported uncertainty is large [57] and the value (5.25 in the deoxy state) is downshifted with respect to the typical His range of \approx 6.5; this appears inconsistent with its short, 2.8 Å distance from the propionate oxygen in the crystal structures (2DN2, 2DN3), determined at a pH of 6.5.

2.9 The experimental pK_a set

Table 1 indicates the number of titratable groups in each protein, and the number for which experimental values are available. We consider acid/base changes for Asp, Cys, Glu, His, Lys, and Tyr side chains. The only titratable Cys residues in our dataset are in hemoglobin. In the simulations, we do not allow the chain termini to change protonation states; they occupy the ionized state that is usual at neutral pH. For myoglobin, we ignore the experimental p K_a 's reported for the His24–His119 pair, since these side chains are closely coupled, sharing a proton over a broad pH range and their titration cannot be described by two acid/base constants (three are needed [22]). This leaves us with 11 myoglobin side chains with experimental p K_a values.

The four Hb subunits include 150 titratable groups in all: 44 lysines, 34 histidines, 30 aspartic acids, 24 glutamic acids, 12 tyrosines and 6 cysteines (not counting arginines, C-, N- terminal groups, histidines that ligate the heme iron, or heme propionates). Only for 13 of the histidines (and their 13 symmetry mates) are experimental values available, both for oxy and deoxy Hb. They are located at the protein surface and their pK_a values were determined experimentally using proton NMR spectroscopy, at 29 °C, with a 100 mM NaCl concentration [91]. One of them is α H45, whose titration we do not predict (see above). The other six histidines are buried. α H87 and β H92 ligate the heme and cannot be protonated. Two are "distal" to the heme, and close to the CO ligand: α H58 and β H63. The last two are α H103 and α H122, whose pK_a values lie outside the experimental pH range, pH = 4 to 9. Overall, for hemoglobin, we compare our predictions to 12 His pK_a's in two protein forms.

2.10 Bohr effect in hemoglobin

For hemoglobin, we compute the acid and alkaline Bohr effects; namely, the release of protons upon oxygenation of the protein, at low or high pH. To compute the number ΔH_i of protons released by side chain i upon oxygenation at a given pH, we subtract the deoxy and oxy protonation probabilities:

$$\Delta H_i = \frac{1}{1 + 10^{pH - pK_{a,i}^{\text{deoxy}}}} - \frac{1}{1 + 10^{pH - pK_{a,i}^{\text{oxy}}}}$$
(16)

This function has a single peak at the position $pH = (pK_{a,i}^{deoxy} + pK_{a,i}^{oxy})/2$, whose height is

$$\Delta H_i(\text{peak}) = \frac{10^{\delta_i/2} - 1}{10^{\delta_i/2} + 1}$$
 (17)

where $\delta_i = pK_{a,i}^{oxy}$ - $pK_{a,i}^{deoxy}$. Both Hb dimers contribute to the Bohr effect. While Hb is symmetric in solution, there are differences between the dimers in the computed pK_a values, due to the crystal structures employed. With FDB, we only obtained pK_a values for one α subunit and one β subunit (see above); we assumed the other two subunits had the same pK_a values. For the NEA computations, $oxy-\alpha_2H20$ titrates outside the simulated pH range; for the Bohr effect calculation, we assumed its pK_a was the same as that of $oxy-\alpha_1H20$.

2.11 Monte Carlo in sequence space

To generate designed PDZ sequences with Proteus, we perform a Monte Carlo simulation where one copy of the folded protein is explicitly represented. The unfolded state is included implicitly, by propagating the simulation with the energy function $E_M = E_f - E_u$, where E_f and E_u are the energies of the folded and unfolded states, respectively. The structure of the unfolded form is not specified; the energy is assumed to be independent of the particular unfolded structure, and to have the additive form:

$$E_u(S) = \sum_{i=1}^n E_u(t_i) \tag{18}$$

where the sum is over the protein residues and $E_u(t_i)$ is a free energy associated with side chain type t_i in the unfolded state. The $E_u(t)$ values associated with each amino acid type were optimized empirically, so that the simulations give an overall amino acid composition that matches the natural PDZ domains in Pfam. The detailed empirical optimization procedure is described elsewhere [92]; the resulting values are reported in Table 3. The folded state energy is obtained from the energy matrix, precomputed as for the acid/base calculations above. Residue pairs were minimized for 15 steps (instead of 30 for the acid/base calculations) and surface coefficients optimized for CPD were used (Table 3). Calculations were done using either the NEA or the FDB method for the GB solvent.

The simulations used a Replica Exchange MC procedure, where eight simulations were run in parallel at different temperatures, and the instantaneous conformations were exchanged randomly according to a Metropolis criterion [93]. Temperatures ranged from 0.125 to $2 \, kT$ units. Gly and Pro residues present in the wildtype protein were not allowed to mutate, and positions that did mutate could not change into Gly or Pro. Simulations were run for 750 million steps (per replica). The 10,000 lowest energy sequences and conformations sampled by any replica were kept for analysis.

2.12 Rosetta sequence generation

Monte Carlo simulations were also done using version 2015.38.58158 of Rosetta (freely available online) [23], using the command:

fixbb -s prot.pdb -resfile prot.res -nstruct 10000 -ex1 -ex2 -linmem_ig 10 where the ex1 and ex2 options activate an enhanced rotamer search for buried side chains,

the last option (linmem_ig) corresponds to on-the-fly energy calculation, and default parameters were used otherwise. Gly and Pro residues present in the wildtype protein were not allowed to mutate, and positions that did mutate could not change into Gly or Pro (as with the Proteus design simulations). Simulations were run for each PDZ domain until 10,000 unique low energy sequences were identified, corresponding to run times of about 5 minutes per sequence on a single core of a recent Intel processor, for a total of 10 hours (per protein) using 80 cores. This was comparable to the cost of the Proteus calculations with NEA (energy matrix plus MC).

2.13 Designed sequence characterization

Designed sequences were compared to the Pfam alignment for the PDZ family, using the Blosum40 scoring matrix and a gap penalty of -6. This matrix is appropriate for comparing rather distant homologs (CPD and Pfam sequences in this case). Each Pfam sequence was also compared to the Pfam alignment, which allowed for comparison between the designed sequences and a typical pair of natural PDZ domains. The Pfam alignment was the "RP55" alignment, consisting of 12,255 sequences. Similarities were computed separately for 14 core residues and for the entire protein.

Designed sequences were submitted to the Superfamily library of Hidden Markov Models [94, 95], which attempts to classify sequences according to the Structural Classification Of Proteins, or SCOP [96]. Classification was based on SCOP version 1.75 and version 3.5 of the Superfamily tools. Superfamily executes the hmmscan program, which implements a Hidden Markov model for each SCOP family and superfamily. The hmmscan program was executed using an E-value threshold of 10^{-10} and a total of 15,438 models to represent the SCOP database.

3 Results

3.1 Acid/base performance for nine test proteins

We first describe the overall performance of the Native Environment Approximation (NEA) and the Fluctuating Dielectric Boundary method (FDB) for nine test proteins (Table 1). We performed pH scans from 0 to 15, to obtain the titration curves of all ionizable sites and we fitted each curve to the modified Hill equation to obtain the pK_a. Calculations were done for a protein dielectric constant $\epsilon_p = 4$. We focus on the pK_a

shifts relative to the appropriate model compound. The rms deviations from experimental values are reported in Table 4 for each protein and in Table 5 for each amino acid type. Notice that some of the experimental measurements only provide an upper or a lower bound for the pK_a . In these cases (15 residues out of 149), if the computed value is compatible with the bound, we consider the deviation to be zero. If the computed value is outside the experimental interval, the deviation is computed with respect to the limit of the interval that is closest to the computed value. This convention is consistent with some previous pK_a benchmark studies. The individual experimental and computed pK_a shifts are compared in Fig. 2.

The overall rms deviation from the experimental values is 1.1 with FDB and 1.2 with NEA. The mean unsigned errors are 0.8 (FDB) and 0.9 (NEA). The correlation between computed and experimental pK_a shifts is 0.7 with FDB and 0.6 with NEA. Thus, there is a small, but systematic improvement with FDB. For each individual side chain i, we denote $\delta E_i = E_i^{\text{NEA}} - E_i^{\text{NEA}}$ the difference between the unsigned NEA and FDB pK_a errors. If δE_i is positive, FDB gives a better result than NEA, and vice versa. The large differences, $|\delta E_i| > 1$, are evenly distributed between the two methods (5 positive and 5 negative values). The intermediate differences are mostly in favor of FDB, with 37 δE_i values between 0.5 and 1, versus 12 between -1 and -0.5, and 20 δE_i values between 0.2 and 0.5, versus 15 between -0.5 and -0.2.

The largest individual FDB errors are 2.5 or 2.6 units for three residues; there are four NEA errors that are larger (Table 4). The largest FDB errors occur for individual Tyr, His, Asp, and Glu residues. On average, the largest rms deviations with FDB are for Asp and Tyr (1.3 units); the largest rms deviations with NEA are for Asp and Glu. FDB does better for all types except Tyr. Notice that FDB uses a single, mean solvation radius B_I for the side chain (residue-GB), which is a more significant approximation for Tyr than for smaller side chains. Table 4 also reports mean errors obtained with the empirical PropKa method [97, 98], which are smaller on average (overall rms deviation from experiment of 0.8) than with FDB. The Null model, which assumes all pK_a shifts are zero, gives an rms deviation of 1.1, like FDB. For positions with large pK_a shifts (one unit or more), FDB does about as well as PropKa (Table 5), slightly better than NEA, and much better than the Null model.

Illustrative titration curves are shown in Figure 3 for the OMTKY3 protein. The numerical populations obtained from the MC simulations follow closely the theoretical, sigmoidal curves, which is an indication of sampling quality. We also show the titration

behavior of a coupled histidine pair in myoglobin, His24 and His119, which bind a single proton together at low pH, with the proton dividing its time between the two side chains (Figure 3). This picture differs from the published interpretation of the NMR experiments, where only a single pK_a was reported and assigned entirely to His119 [99].

3.2 Performance and pK_a error analysis for hemoglobin

With NEA, all titratable residues change their protonation state between pH = 0 and 15 except α_1 H45 in the deoxy form and α_2 H20, α_1 H58, α_2 H58, β_1 H63, β_2 H63, α_1 Y42, α_1 K60, α_1 K61, β_1 K66, α_2 E23, α_2 Y24, α_2 K56, α_2 K60, α_2 K61, α_2 C104, β_2 K59 in the oxy form. Small differences are seen between the two dimers. The mean unsigned difference between the predicted pK_a's of the two dimers is 0.3, with a few sizeable differences (5 of more than 0.5 units, out of 30). Oxy- α 20 titrates in one dimer but not the other. Below, we use the titrating His for our error statistics, excluding the non-titrating symmetry mate.

Overall, the NEA method performs well. We achieved a good agreement with experiment for a large group of 24 histidines (two forms of Hb with 12 independent histidines per tetramer), with an rms deviation of 0.9. The errors are similar to our previous study, where 27 histidine pK_a's in 12 proteins were predicted with an rms deviation of 1.0 using $\epsilon_p = 4$ [22]. The individual computed and experimental pK_a's are reported in Table 6. The pK_a shifts are compared to experiment in Fig. 4. The data set includes three residues per α/β dimer with experimental pK_a shifts of over 1 unit: β H97, β H143 and β H146. 10 of the 12 His residues (per α/β dimer) are predicted with the correct pK_a shift direction in the deoxy form, and 8 of 12 in the oxy form. The large pK_a errors (unsigned errors of 0.7 or more) are all positive, meaning the positive His form is overstabilized.

The FDB model performs slightly better than NEA, with an overall rms deviation of 0.7 from experiment. Of the ten residues with NEA pK_a errors of 0.7 or more, nine are improved with FDB and one is slightly worse (oxy- β 116, error increased from 1.9 to 2.1 units). The large errors are all positive, as with NEA. The experimental Hill coefficients of the individual His titration curves are fairly well-reproduced, with an rms deviation of 0.3.

3.3 Bohr effect in hemoglobin

We consider next the acid and alkaline Bohr effects; namely, the release of protons from Hb upon oxygenation of the protein, at low or high pH. To compute the number ΔH_i of protons released by side chain i upon oxygenation at a given pH, we subtract the fractional populations of the protonated state in the deoxy and oxy forms, Eq. (16). $\Delta H_i(pH)$ has a single peak at pH(peak) = $(pK_{a,i}^{deoxy} + pK_{a,i}^{oxy})/2$. The peak height is

$$\Delta H_i(\text{peak}) = \frac{10^{\delta_i/2} - 1}{10^{\delta_i/2} + 1} = \frac{\ln 10}{4} \,\delta_i + O(\delta_i^2) \tag{19}$$

where $\delta_i = \mathrm{pK}_{a,i}^{\mathrm{deoxy}}$ - $\mathrm{pK}_{a,i}^{\mathrm{oxy}}$ and the rightmost form is a power expansion where $O(\delta_i^2)$ is of order δ_i^2 . The peak has the same sign as δ_i , as expected; e.g., if $\mathrm{pK}_{a,i}^{\mathrm{deoxy}}$ is large and $\mathrm{pK}_{a,i}^{\mathrm{oxy}}$ small $(\delta_i > 0)$, the group releases a proton upon oxygenation at an intermediate $\mathrm{pH}\ (\Delta H_i(\mathrm{peak}) > 0)$. The area of the peak is found to be $A_i = \delta_i$.

From oxygenation measurements, there are a maximum of 1.96 protons released upon oxygenation at pH = 7.1 (alkaline Bohr effect) and a maximum of 1.40 protons absorbed at pH = 5.2 (acid Bohr effect) [90]. The total effect is well accounted for by the contributions of the His side chains for which experimental pK_a 's are available. We focus on these 12 histidines (we exclude α H45, which interacts closely with a titratable heme propionate; see Methods). The individual His peaks are described in Table 7. The proton release curve due to the 12 histidines is shown in Fig. 5, which includes the experimental, FDB and NEA results. The experimental curve has a negative peak at pH = 5.2 (1.3 protons absorbed) and a positive peak at pH = 7.1 (1.9 protons released). The FDB curve has the alkaline peak at the correct position but with an overestimated height (2.9 protons released). The overestimate is due to small errors (about 0.2–0.4 protons each) in four His peaks ($\alpha 20$, $\alpha 112$, $\beta 77$, $\beta 146$), which are well within the mean error level of the individual pK_a's and the δ_i values. The computed pK_a's are too high for β 116 and β 117 and the peaks are too high, but the height errors have opposite signs and cancel out. The NEA behavior is reported in Table 7 and Fig. 5. The alkaline peak height is somewhat better than FDB but the peak is much too broad, extending down to the acid range and merging with the acid peak to form a single broad peak.

At acidic pH, five histidines (excluding α H45) have experimental peaks below pH = 6.5, absorbing a maximum of 1.3 protons. The acidic peak is absent from the FDB curve. The main experimental contributors are α 89 (+0.61 protons), and β 143 (-0.93 protons). α 89 is well-reproduced by FDB (with a slight pH upshift), but β 143 releases 0.36 protons.

Results with NEA are distinctly worse, with a signel broad positive peak extending down to pH = 4. Overall, the main error in the Bohr effect with FDB is the lack of proton absorption by β 143 at acidic pH, due to small β 143 pK_a errors (0.3–0.5 units) in the oxy and deoxy forms.

3.4 Complete redesign of three PDZ domains

Proteus design simulations used Replica Exchange Monte Carlo with eight replicas and 750 million steps per replica, at thermal energies kT that ranged from 0.125 to 2 kcal/mol. All positions (except Gly and Pro) were allowed to mutate freely into all amino acid types except Gly and Pro. The simulations were done with a protein dielectric constant of four. The unfolded energies and surface energy coefficients employed are reported in Table 3. The 10,000 sequences with the lowest energies among those sampled by any of the MC replicas were retained for analysis, along with 10,000 sequences designed with Rosetta. The sequences were analyzed by the Superfamily fold recognition tool [95, 100] (Table 8). For all three proteins, all 10,000 Rosetta sequences were classified as PDZ domains by Superfamily. With Proteus and NEA, 98.7% of the sequences were classified as PDZ domains. With the FDB method, there is a clear improvement, as all 10,000 of the Proteus sequences were correctly classified, for all three proteins, with match lengths and E-values that are significantly better than with NEA, and comparable to Rosetta.

Fig. 6 shows the Blosum40 similarity scores between designed and natural sequences, for each PDZ protein, both for core positions and the entire protein. With Rosetta, the designed sequences have scores that are similar to the Pfam sequences themselves, especially for the core positions. With Proteus and the NEA, the similarities for the protein core are excellent, as observed previously [92]. In the core, we are far from the protein/solvent boundary so that an approximate NEA treatment is not damaging. For the entire protein, the overlap with Pfam scores is less good, especially for the NHERF case, where the Proteus sequences overlap with the bottom of the peak of natural scores. Switching to the FDB method, there is a slight decrease for syntenin in the overall similarity to Pfam, and a significant decrease for the core residues. For DLG2, there is a significant improvement with FDB and for NHERF, there is a very large improvement, about 50 Blosum points. For all three proteins, the overall similarities with FDB overlap with the lower half of the Pfam peak and are close to the Rosetta similarities, a significant improvement over NEA.

4 Concluding discussion

Simple continuum dielectric models are important tools to understand and engineer electrostatic interactions in proteins. Some variants use a fixed protein conformation, while others use an all-atom molecular dynamics exploration of protein conformations. The present variant is in between, with a fixed protein backbone but explicit exploration of side chain rotamers. This level of detail is common in protein design applications, but less so for acid/base applications, especially in combination with a GB solvent model. With this level of detail, the protein/solvent dielectric boundary fluctuates over time due to thermal motion and reorganization in response to protonation changes. In previous studies, these dielectric shape fluctuations were neglected, and a mean, effective dielectric boundary was used instead. Here, we have applied and tested an improved variant, where the protein shape is rigorously accounted for in the GB interaction energies at all times. In this way, we achieved several goals.

Our first goal was to implement the more rigorous GB method within the Proteus software and model framework. This framework uses a precalculated energy matrix to allow very efficient Monte Carlo simulations of proteins, where rotamers, protonation states, and side chain types can all vary [45, 93]. The present implementation is about four times as costly as the NEA method; its cost is reduced by half compared to an earlier FDB implementation [37]. The present implementation is also much friendlier and more flexible than the earlier one. In particular, it makes it straightforward to use FDB for some residue pairs and NEA for others, further reducing the cost.

A second goal was to verify that the more rigorous FDB method leads to more accurate acid/base and protein design calculations, which is indeed the case. We observed a small but significant improvement for acid/base constants in nine proteins. The rms error and correlation with experiment improved and there were fewer large errors, compared to NEA. The rms error with FDB was 1.1 for the present test set, the mean unsigned error was 0.8, and the performance for residues with large pK_a shifts was rather good. The performance was not as good as some more empirical models, such as PropKa or Rosetta [21] or the simplest PB models. These last models employ a large protein dielectric constant and therefore behave much more like the Null model. Here, good performance was obtained with a model that is physically plausible, including a low value (ϵ_P =4) of the protein dielectric constant, which is used to account for the protein's electronic polarizability and motions of the protein backbone [101]. This low value is in line with experimen-

tal measurements on protein powders and acetamide crystals, and with MD simulation analyses, as reviewed earlier [4]. It is also more consistent with the use of atomic charges taken from molecular mechanics force fields, which are optimized for use with a low solute dielectric constant. Notice that errors introduced by the NEA method tend to scale with the inverse of the protein dielectric constant [22, 50], so that studies with a low ϵ_P can be done with much more confidence using FDB instead of NEA.

There was also a distinct improvement with FDB for the design of three PDZ domains. There was a slight performance decrease for syntenin, compared to NEA, but a significant improvement for DLG2 and a very large improvement for NHREF. Fold recognition tests with the Superfamily library of Hidden Markov Models gave better results for all three proteins using FDB. The good design results were obtained with a low protein dielectric of four, in contrast to earlier work where a larger value of around eight worked best [92].

Several aspects of the model could be analyzed and improved further. Treatment of ionic strength might improve performance [22, 102]. The GB parameters and surface energy coefficients could be optimized further, since previous parameterization did not use "Residue-GB" [46]. The structural model could perhaps be improved by including several backbone conformations, for example by extracting models from a molecular dynamics simulation. Monte Carlo hops between backbone conformations could then be combined with rotamer and protonation or mutation moves [103].

By eliminating one source of error in our MC method, we expect to obtain not only increased accuracy but an improved understanding of other sources of error. The method described here can be used as a complement to fast, empirical acid/base predictors like PropKa, since the cost of a full titration scan is about two hours on a small computer cluster. While this is far more expensive than PropKa, it is still fast enough to perform as a routine, preliminary analysis of a protein of interest. Since the performance for large pK_a shifts is similar to PropKa, our method can provide a useful additional check for such residues. Finally, by implementing FDB in our protein design software Proteus, we improved the design performance and also make it easier to perform protein design applications that combine changes in protonation states with side chain mutations. Protonation changes can also be modelled in combination with the binding/unbinding of a ligand [104], two processes that are often coupled [19].

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Table 1: Acid/base test proteins

	PDB	number of	titratable	experimental	acomputed
Protein	code	residues	side chains	pK_a 's	pK_a 's
BPTI	4pti	58	12	12	12
OMTKY3	2gkr	56	13	$13 \ (3)^b$	13
protein G	1pga	56	19	17 (4)	16
barnase	1a2p	110	29	14 (1)	14
lysozyme	2lzt	129	18	19	18
myoglobin	1a6k	151	54	12 (2)	11
SNase	$3 \mathrm{bdc}$	141	44	17(2)	17
RNase H1	1goa	155	43	24 (3)	24
deoxy Hb	2dn2	287×2	75×2	12	12
oxy Hb	2dn3	287×2	75×2	12	12
Total			532	$155 \ (15)^b$	$149 \ (15)^b$

^aWe exclude titratable side chains at the chain termini (6 side chains).

 $^{{}^}b\mathrm{In}$ parentheses: number of cases where only an upper or lower bound is available.

Table 2: Model compound protonation free energies ΔG_X

Protonation	Exp.				$-kT \times$		$2.303 kT \times$	ΔG_X
reaction	pK_a	E_X	E_{XH}	ΔE_X	$\ln \frac{n_{XH}}{n_X}$	$\delta g_X^{ m conf}$	$(pK_a^X - 7)$	(pH = 7)
$Asp^- o AspH$	3.9	-9.8	-1.2	8.6	-0.42	0.12	-4.28	4.02
$\mathrm{Glu^-} \to \mathrm{GluH}$	4.3	-7.2	0.2	7.4	-0.42	0.07	-3.73	3.32
${\rm Tyr}^-\to {\rm TyrH}$	9.8	-5.4	1.1	6.5	-0.42	0.08	+3.87	10.03
$\mathrm{Cys}^- \to \mathrm{CysH}$	8.6	-11.4	7.6	19.0	-0.66	-0.04	+2.21	20.51
$\mathrm{Lys^0} \to \mathrm{Lys^+}$	10.4	6.3	4.6	-1.7	0	0	+4.70	3.00
$\mathrm{His}_\delta \to \mathrm{His}^+$	6.5	21.5	25.3	3.8	0	0	-0.69	3.11
$\mathrm{His}_{\epsilon} \to \mathrm{His}^+$	7.0	20.7	25.3	4.6	0	0	0.00	4.60

In kcal/mol. Individual terms are defined in Eq. (14). E_X and E_{XH} are from earlier work [22]. We set kT to 0.6 kcal/mol.

Table 3: Unfolded state energies $E_u(t)$ for CPD

	F	DB	N	EA				
Residues	Buried	Exposed	Buried	Exposed				
ALA	0.00	0.00	0.00	0.00				
CYS	-1.06	-1.64	-0.89	-2.57				
THR	-4.84	-6.68	-5.31	-8.07				
SER	-4.45	-5.24	-5.55	-6.55				
ASP	-14.55	-18.82	-17.26	-22.06				
GLU	-14.52	-18.21	-16.12	-20.68				
ASN	-14.02	-17.80	-16.38	-20.41				
GLN	-13.14	-16.61	-14.00	-18.41				
$^a\mathrm{HIS}^+$	12.86	10.98	11.21	10.72				
${}^a{ m HIS}_\epsilon$	10.85	8.12	10.63	6.95				
$^a\mathrm{HIS}_\delta$	10.41	7.37	15.17	6.15				
ARG	-51.37	-54.76	-53.40	-57.36				
LYS	-8.24	-11.35	-8.20	-12.34				
ILE	5.50	3.06	6.76	3.44				
VAL	-0.05	-1.66	0.43	-2.19				
LEU	0.00	-2.94	0.52	-3.72				
MET	-2.85	-3.09	-1.61	-3.21				
PHE	0.17	-3.18	1.86	-2.68				
TRP	-1.94	-5.53	-0.23	-7.67				
TYR	-5.91	-10.14	-5.10	-10.90				

Unfolded state energies $E_u(t)$ (kcal/mol) for each amino acid type t, optimized with NEA or FDB; there are distinct values for positions that are buried or exposed (in the folded structure). Calculations were done with a protein dielectric constant of four and the following surface energy coefficients (cal/mol/Å²):
-60 (hydrophobic), 30 (aromatic), -120 (polar), -110 (ionic). Buried positions are those with a fractional solvent accessible surface area of less than 0.30; other positions are exposed. ^aHis protonation states.

Table 4: Deviation between experimental and computed pK_a 's per protein

	number	NEA	NEA	FDB	FDB	FDB	PropKa
Protein	of pK_a 's	$\max(\text{position})^a$	rmsd	rmsd	mue^b	$\max(\text{position})^a$	rmsd^c
BPTI	12	-1.6 (Y35)	0.8	0.8	0.6	-2.5 (Y35)	0.4
OMTKY3	13	-2.4~(K29)	1.1	0.9	0.7	-1.8 (Y31)	0.6
protein G	16	-1.8 (D47)	0.8	0.8	0.5	-2.0 (D47)	0.6
barnase	14	2.5 (D101)	1.1	1.0	0.8	2.0 (D101)	1.2
lysozyme	18	-3.4 (D66)	1.4	1.1	0.8	-2.8 (E35)	0.5
myoglobin	11	1.7 (Y146)	0.9	1.1	0.9	2.1 (Y103)	0.8
SNase	17	2.7 (D19)	1.5	1.3	1.1	-2.3 (E135)	1.3
RNase H1	24	3.0 (D148)	1.5	1.4	1.0	-2.6(H114,D134)	0.8
deoxy Hb	12	$2.1 \; (H\beta 117)$	0.9	0.7	0.5	$1.1~(\mathrm{H}\beta1$	
oxy Hb	12	$1.9 \ (H\beta 116)$	1.1	0.8	0.6	$1.8~(\mathrm{H}\beta11$	
All	149		1.2	1.1	0.8		0.8
			1				

 $[^]a$ Maximum error and corresponding residue(s). b Mean unsigned error. c From earlier work [22].

Table 5: Deviation between experimental and computed $\mathbf{p}\mathbf{K}_a$'s per residue class

	number	rmsd	rmsd	
	of pK_a 's	NEA	FDB	PropKa^a
Asp	38	1.3	1.2	0.9
Glu	37	1.4	1.2	0.9
His	41	0.9	0.7	1.0
Lys	17	1.0	0.8	0.6
Tyr	15	0.8	1.2	0.6
shift<1	95	1.1	1.0	0.6
$1 \le \text{shift} < 2$	32	1.1	1.0	1.0
$2 \leq \text{shift}$	7	1.8	1.6	1.5

 $^{^{}a}$ From earlier work [22].

Table 6: pK_a shifts for histidines in oxy/deoxy hemoglobin

_	deoxy					oxy			
	Exp.	Exp.	NEA	FDB	Exp.	Exp.	NEA	FDB	
Residue	pK_a	shift	shift	shift	pK_a	shift	shift	shift	
α 20	7.02	0.52	$1.25[0.7]^a$	$0.88[0.4]^a$	7.08	0.58	$1.00[0.4]^a$	$0.75[0.2]^a$	
α 45	5.25	-1.25			6.12	-0.38			
α 50	7.14	0.64	1.00[0.4]	0.62[0.1]	6.90	0.40	0.50[0.1]	0.00[0.4]	
α 58			0.88	1.58					
α 72	7.47	0.97	1.67[0.7]	1.5[0.5]	7.27	0.77	1.83[1.0]	1.50[0.7]	
α 89	6.80	0.30	0.75[0.5]	0.94[0.6]	6.25	-0.25	0.12[0.4]	0.16[0.4]	
α 103			1.62	1.62			0.62	0.75	
α 112	7.49	0.99	1.00[0.1]	1.25[0.3]	7.53	1.03	0.62[0.5]	1.00[0.0]	
α 122			-0.50	-1.34			-0.12	-1.12	
β 2	6.17	-0.33	-0.25[0.1]	-0.50[0.2]	6.39	-0.11	0.62[0.7]	0.42[0.5]	
β 63			1.25	1.88					
β 77	7.46	0.96	1.88[0.9]	1.75[0.8]	7.79	1.29	1.88[0.6]	1.75[0.5]	
β 97	8.01	1.51	1.38[0.1]	1.75[0.2]	7.75	1.25	1.31[0.1]	1.75[0.5]	
β 116	6.35	-0.15	1.00[1.1]	1.00[1.1]	6.13	-0.37	1.62[1.9]	1.75[2.1]	
β 117	6.43	-0.07	2.00[2.1]	1.44[1.5]	6.39	-0.11	1.16[1.3]	1.00[1.1]	
β 143	4.70	-1.80	-0.75[1.0]	-1.25[0.6]	5.57	-0.93	-1.25[0.3]	-1.25[0.3]	
β 146	7.93	1.43	1.25[0.2]	1.25[0.2]	6.42	-0.08	-1.25[1.7]	-0.38[0.3]	
α	rms de	eviations:	0.5	0.4			0.8	0.5	
β	$\operatorname{rms}\mathrm{d}\epsilon$	eviations:	1.0	0.8			1.0	0.8	

 $[^]a\mathrm{Values}$ in brackets are unsigned errors.

Table 7: Bohr effect contributions from individual residues

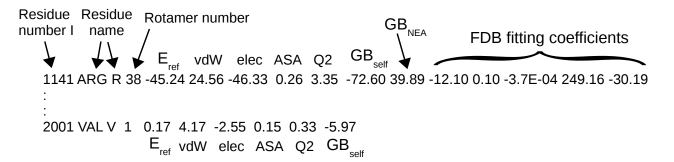
	Exp	Experiment		FDB		NEA
Residue	рН	protons	рН	protons	рН	protons
β H143	5.1	-0.93	5.2	0.36	5.5	0.56
β H116	6.2	0.26	8.0	-1.04	7.8	-0.68
$\beta \mathrm{H2}$	6.3	-0.25	6.5	-0.89	6.7	-0.93
β H117	6.4	0.05	7.8	0.93	8.1	0.90
α H89	6.5	0.61	7.0	0.65	6.9	0.69
α H50	7.0	0.27	6.8	0.56	7.3	0.56
α H20	7.1	-0.07	7.4	0.29	7.6	0.29
β H146	7.2	1.40	7.0	1.64	6.6	1.84
α H72	7.4	0.23	8.0	0.00	8.3	-0.18
$\alpha \mathrm{H}112$	7.5	-0.05	7.6	0.29	7.3	0.43
β H77	7.6	-0.37	8.9	-0.05	8.4	0.00
β H97	7.8	0.30	7.9	0.29	7.8	0.08

Table 8: Fold recognition of designed sequences by Superfamily

	Design	^a Match/seq	b Superfamily	c Superfamily	b Family	cFamily
Protein	model	length	E-value	success $\#$	E-value	success $\#$
NHERF	Proteus, NEA	62/91	3.2e-3	9857	1.0e-2	9857
syntenin	Proteus, NEA	70/82	2.8e-3	9879	3.6e-3	9879
DLG2	Proteus, NEA	83/97	1.7e-3	9876	3.2e-3	9876
NHERF	Proteus, FDB	80/91	8.5e-14	10000	8.9e-3	10000
syntenin	Proteus, FDB	70/82	2.8e-6	10000	2.7e-3	10000
DLG2	Proteus, FDB	88/97	3.3e-12	10000	2.0e-3	10000
NHERF	Rosetta	79/91	1.3e-13	10000	2.2e-3	10000
syntenin	Rosetta	76/82	7.3e-13	10000	1.8e-3	10000
DLG2	Rosetta	86/97	1.3e-9	10000	9.6e-4	10000

 $[^]a$ The average match length for sequences recognized by Superfamily and the total sequence length. b Average E-values for Superfamily assignments to the correct SCOP superfamily/family. c The number of designed sequences (out of 10000 tested) assigned to the correct SCOP superfamily/family.

Diagonal matrix elements



Off-diagonal matrix elements

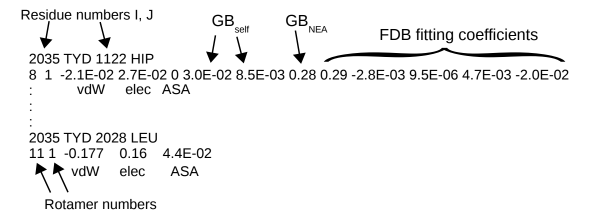


Figure 1: Energy matrix elements. For diagonal elements (above), we show an example of an FDB element and an NEA element (one rotamer each). The individual energy components are labelled. ASA labels the surface energy term. Although the Arg1141 position is treated with FDB, the matrix includes the NEA estimate of the GB contribution. The quantity labeled Q2 is the total squared charge $\sum_{i \in I} q_i^2$, needed to compute the solvation radius B_I (Eq. 6). The five rightmost quantities are the fitting coefficients c_n^{II} (Eq. 9). For off-diagonal elements (below), we also show an FDB and an NEA pair (one rotamer combination each). For the FDB pair, the GB self-energy contributions E_{IJ}^{self} and E_{JI}^{self} (Eq. 3) are both stored (the GB self energy is non-symmetric).

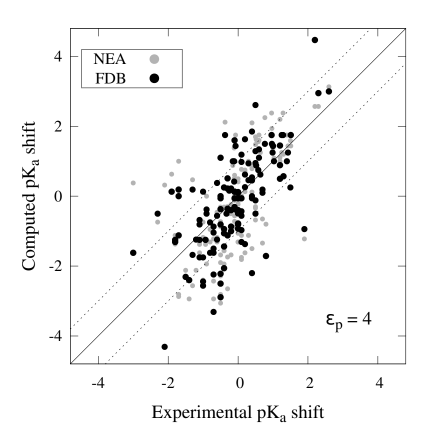


Figure 2: Comparison between computed and experimental pK_a shifts.

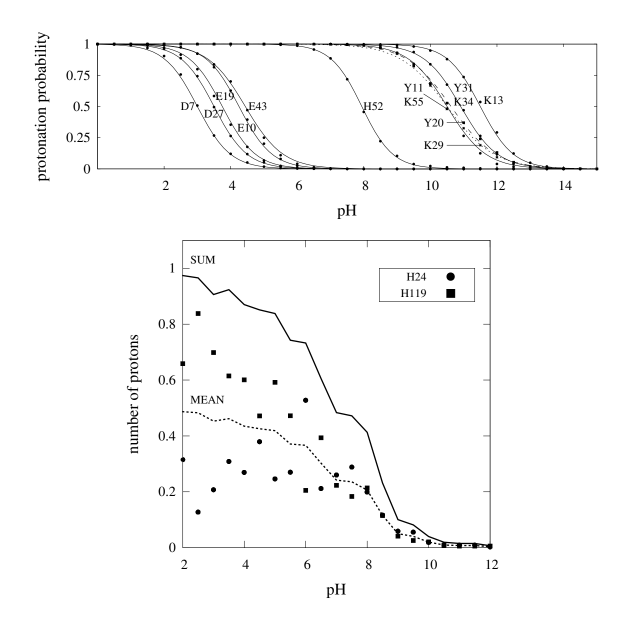


Figure 3: Above: OMTKY3 Titration curves. Below: titration of a coupled histidine pair in myoglobin (FDB results). The pair binds one proton at low pH. "MEAN" indicates the mean number of protons per histidine (0.5 at low pH).

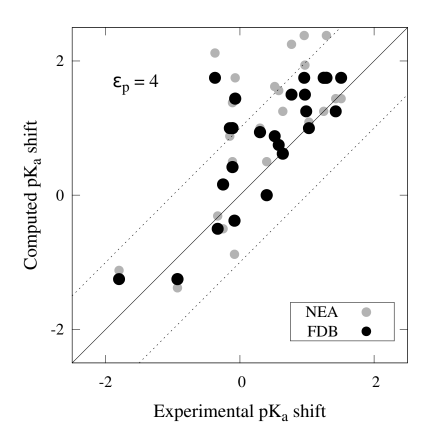


Figure 4: Comparison between computed and experimental pK_a shifts for hemoglobin.

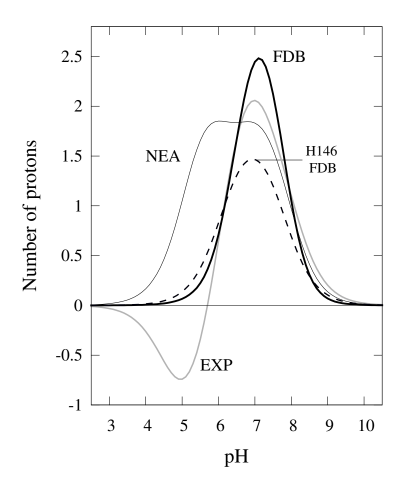


Figure 5: Bohr effect in hemoglobin due to histidine residues (excluding H45; see text). Both FDB and NEA predictions are shown. The separate contribution of β His146 is shown.

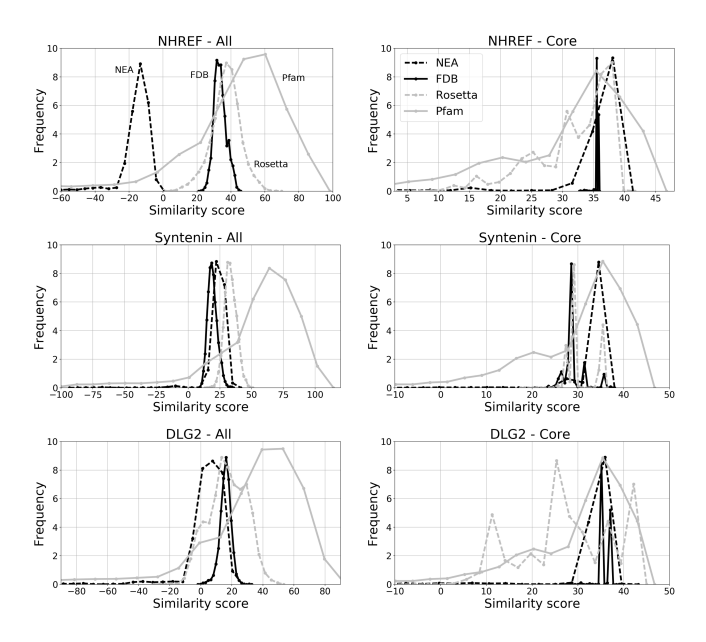


Figure 6: Histogram plots showing similarity scores for designed PDZ sequences. Similarity scores for the entire protein (left) or 14 core positions (right). Values are shown for Proteus(NEA), Proteus(FDB), Rosetta, and Pfam sequences (all compared to the Pfam RP55 alignment).