In integrative structural biology/hybrid modeling approaches, we integrate structural models of macromolecules and experimental data to obtain faithful representations of the structures underlying the data. For example, in ensemble refinement by reweighting we first generate structural ensembles of flexible and dynamic biological macromolecules in molecular simulations. In a subsequent reweighting step, we refine the statistical weights of the structures to strike a balance between the information provided by simulations and by experimental data. For the "Bayesian inference of ensembles" approach (BioEn), we present two complementary methods to solve the underlying challenging high-dimensional optimization problem. We systematically investigate reliability, accuracy, and efficiency of these methods and integrate molecular dynamics simulations of the disordered peptide Ala-5 and NMR J-couplings. We provide an open-source library free of charge at https://github.com/bio-phys/BioEN/optimize.
Efficient Ensemble Refinement by Reweighting

Jürgen Köfinger, *, † Lukas S. Stelzl, † Klaus Reuter, ‡ César Allande, † Katrin Reichel, † and Gerhard Hummer *, † , ¶

† Department of Theoretical Biophysics, Max Planck Institute of Biophysics, Max-von-Laue-Straße 3, 60438 Frankfurt am Main, Germany
‡ Max Planck Computing and Data Facility, Gießenbachstr. 2, 85748 Garching, Germany
¶ Institute for Biophysics, Goethe University, Max-von-Laue-Straße 9, 60438 Frankfurt am Main, Germany

E-mail: juergen.koefinger@biophys.mpg.de; gerhard.hummer@biophys.mpg.de

Abstract

Ensemble refinement produces structural ensembles of flexible and dynamic biomolecules by integrating experimental data and molecular simulations. Here we present two efficient numerical methods to solve the computationally challenging maximum-entropy problem arising from a Bayesian formulation of ensemble refinement. Recasting the resulting constrained weight optimization problem into an unconstrained form enables the use of gradient-based algorithms. In two complementary formulations that differ in their dimensionality, we optimize either the log-weights directly or the generalized forces appearing in the explicit analytical form of the solution. We first demonstrate the robustness, accuracy, and efficiency of the two methods using synthetic data. We then use NMR J-couplings to reweight a molecular dynamics simulation ensemble of the disordered peptide Ala-5 simulated with the AMBER99SB*-ildn-q force field. After reweighting, we find a consistent increase in the population of the polyproline-II conformations and a decrease of α-helical-like conformations. Ensemble refinement makes it possible to infer detailed structural models also of disordered biomolecules by combining input from experiment and simulation in a balanced manner.

1 Introduction

To infer structures and functions of biological macromolecules, we combine information from diverse experimental and theoretical sources. However, in many experiments the observables reporting on biomolecular structure are averaged over ensembles. Nuclear magnetic resonance (NMR) and pulsed electron paramagnetic resonance (EPR) experiments provide ensemble-averaged high-resolution information about distances (using, e.g., the nuclear Overhauser effect, paramagnetic relaxation enhancement, or double electron-electron resonance (DEER)) and angles (using, e.g., J-couplings and residual dipolar couplings). Small-angle X-ray scattering (SAXS) experiments provide ensemble-averaged information about macromolecular size and shape, and wide-angle X-ray scattering (WAXS) experiments report on secondary structure and fold. Ensemble refinement promises faithful descriptions of the true ensemble of structures underlying the experimental data even for highly dynamic systems.

The conformational diversity of the ensemble can be described in terms of a set of representative reference structures. The relative weights of the ensemble members are then determined by ensemble refinement against experimental data. To regularize this inverse problem one can, e.g., restrict the number of conformers...
as is done in minimal-ensemble refinement or replica simulations; limit the weight changes relative to the reference ensemble as is done in maximum-entropy approaches or in Bayesian formulations; or limit both ensemble size and weight changes. See ref 15 for an in-depth discussion and further references.

The reference ensemble is often defined in terms of a molecular simulation force field, i.e., a classical potential energy function for which one has some confidence that it captures essential features. The experimental data can then be used directly as a bias in molecular dynamics (MD) simulations or a posteriori to reweight an unbiased ensemble in a way that improves the agreement with experiment. Biased simulations improve the coverage of the configuration space but suffer from finite-size effects due to a limited ensemble size in simulations. Reweighting requires good coverage but can handle much larger ensemble sizes. The “Bayesian inference of ensembles” (BioEn) approach makes it possible to combine, if needed, biased sampling and subsequent reweighting to ensure both good coverage of the configuration space and a well-defined, converged ensemble.

Ensemble refinement by reweighting is a computationally challenging optimization problem because the number of structures in the ensemble, usually generated in simulations, and the number of experimental data points provided by experiments can both be large. Simulations can easily create hundreds of thousands of structures. In general, we would like to include as many structures as possible in ensemble refinement, not only to avoid artifacts due to the finite size of the ensemble but also to ensure that we pick up small but significant sub-ensembles. Experiments like NMR, SAXS/WAXS, and DEER can provide thousands of data points. The numbers of pixels or voxels in electron-microscopy projection images or 3D maps, respectively, are of even larger magnitude. More than ten thousand data points are thus common when integrating data from different experimental sources.

With respect to computational efficiency, we also have to take into account that we usually want to perform multiple reweighting runs for different sub-ensembles and sub-sets of the experimental data, while at the same time varying the confidence that we have in the reference ensemble. Consequently, we have to be able to efficiently solve the optimization problem underlying ensemble refinement by reweighting for large numbers of structures and data points.

The paper is organized as follows. In Theory, we derive two complementary numerical methods. We calculate the optimal ensemble by reweighting based on the “ensemble refinement of SAXS” (EROS) method, which is a special case of BioEn. In both methods, constraints are taken into account implicitly such that we can take advantage of efficient gradient-based optimization algorithms. In the first method, we solve for the logarithms of the statistical weights, where \( N \) is the number of structures. In the second method we solve for \( M \) generalized forces, where \( M \) is the number of experimental data points. The efficiency of the two methods depends on \( N \) and \( M \). For both methods, we derive analytical expressions for the gradients that render gradient-based optimization algorithms highly efficient. In Results, we systematically investigate the efficiency and accuracy of these methods using synthetic data. For illustration, we then refine fully atomistic MD simulations of Ala-5 using J-couplings.

2 Theory

We first present the BioEn posterior, whose maximum determines the optimal statistical weights of the structures in the ensemble. We then show that the optimal solution is unique. To be able to apply gradient-based optimization methods to the constrained optimization problem, we recast the posterior as a function of log-weights and as a function of the generalized forces, as already introduced in ref 20. For both formulations, we calculate the respective gradients analytically, facilitating efficient optimization. We focus here on uncorrelated Gaussian noise and present the corresponding expressions for correlated Gaussian noise in Appendix A.
2.1 Background

In the BioEn method,\textsuperscript{20} which is a generalization of the EROS method,\textsuperscript{14} we determine the optimum of the posterior probability as function of the statistical weights $w_\alpha$, where $\alpha$ is the index of the $N$ ensemble members ($\alpha = 1 \ldots N$) given the experimental data,

$$ P(\mathbf{w}|\text{data}) \propto P(\mathbf{w})P(\text{data}|\mathbf{w}). \quad (1) $$

$P(\text{data}|\mathbf{w})$ is the likelihood function, and $\mathbf{w}$ is the vector of weights $w_\alpha$. The prior is given by

$$ P(\mathbf{w}) \propto \exp(-\theta S_{\text{KL}}) = \prod_{\alpha=1}^{N} \left( \frac{w_\alpha^0}{w_\alpha} \right)^{\theta w_\alpha} \quad (2) $$

where

$$ S_{\text{KL}} = \sum_{\alpha=1}^{N} w_\alpha \ln \left( \frac{w_\alpha}{w_\alpha^0} \right) \quad (3) $$

is the Kullback-Leibler divergence.\textsuperscript{20} Both refined weights ($w_\alpha > 0$) and reference weights ($w_\alpha^0 > 0$) are normalized, $\sum_{\alpha=1}^{N} w_\alpha = \sum_{\alpha=1}^{N} w_\alpha^0 = 1$. The parameter $\theta$ expresses the confidence in the reference ensemble. Large values of $\theta$ express high confidence and the optimal weights will be close to the reference weights.

Instead of maximizing the posterior with respect to $w_\alpha$, we can minimize the negative log-posterior given by

$$ L = \theta \sum_{\alpha=1}^{N} w_\alpha \ln \left( \frac{w_\alpha}{w_\alpha^0} \right) - \ln P(\text{data}|\mathbf{w}) \quad (4) $$

The optimization problem is constrained by

$$ 0 \leq w_\alpha \text{ for all } \alpha \quad (5) $$

and

$$ \sum_{\alpha} w_\alpha = 1, \quad (6) $$

i.e., the weights lie in a simplex. For uncorrelated Gaussian errors $\sigma_i$ of the ensemble-averaged measurements $Y_i$ of the observables $i = 1, \ldots, M$, the likelihood is given by

$$ P(\text{data}|\mathbf{w}) \propto \exp \left( -\frac{\chi^2}{2} \right) = \exp \left( -\sum_{i=1}^{M} \frac{(\sum_{\alpha=1}^{N} w_\alpha y_\alpha^\alpha - Y_i)^2}{2\sigma_i^2} \right) \quad (7) $$

Here, $y_\alpha^\alpha$ is the calculated value of observable $i$ for the individual structure $\alpha$. Note that the measurement $Y_i$ can stem from different experimental methods, e.g., from SAXS and NMR, and that $\sigma_i^2 = (\sigma_i^{\text{exp}})^2 + (\sigma_i^{\text{calc}})$ is the sum of uncertainties in the experiment and in the calculation of $y_\alpha^\alpha$.\textsuperscript{4}

The negative log-posterior then becomes

$$ L = \theta \sum_{\alpha=1}^{N} w_\alpha \ln \left( \frac{w_\alpha}{w_\alpha^0} \right) + \sum_{i=1}^{M} \frac{1}{\sigma_i^2} \left( \sum_{\alpha=1}^{N} w_\alpha y_\alpha^\alpha - Y_i \right)^2 \quad (8) $$

The negative log-posterior $L$ corresponds to the EROS free energy $\chi^2 - \theta S$, where $S = -S_{\text{KL}}$ is the negative Kullback-Leibler divergence.\textsuperscript{14} Note that the BioEn and EROS formulations differ by a factor $1/2$ scaling $\chi^2$, which trivially rescales $\theta$.

To solve this optimization problem efficiently, we first show that the negative log-posterior is convex such that there is a unique solution. The gradient of eq 8 is given by

$$ \frac{\partial L}{\partial w_\alpha} = \theta \left( \ln \left( \frac{w_\alpha}{w_\alpha^0} \right) + 1 \right) + \sum_{i} \frac{y_\alpha^\alpha \langle y_i \rangle - Y_i}{\sigma_i^2} \quad (9) $$

where angular brackets indicate the average over the reweighted ensemble, i.e., $\langle y_\alpha \rangle = \sum_{\alpha} w_\alpha y_\alpha^\alpha$. The Hessian is given by

$$ h_{\alpha\gamma} = \frac{\partial^2 L}{\partial w_\alpha \partial w_\gamma} = \frac{\theta}{w_\alpha} \delta_{\alpha\gamma} + \sum_{i} \frac{y_\alpha^\alpha y_i^\gamma}{\sigma_i^2} \quad (10) $$

where $\delta_{\alpha\gamma} = 1$ if $\alpha = \gamma$ and $\delta_{\alpha\gamma} = 0$ otherwise. By casting the Hessian in this form, as a sum of a positive definite diagonal matrix and of dyadic products of vectors, it is straightforward to show that the quadratic form $\mathbf{x}^T \mathbf{h} \mathbf{x}$ is...
positive definite,

$$\sum_{\alpha, \gamma} x_{\alpha} h_{\alpha \gamma} x_{\gamma} = \theta \sum_{\alpha} \frac{x_{\alpha}^2}{w_{\alpha}} + \sum_{i} \left( \frac{\sum_{\alpha} x_{\alpha} y_{i}^{\alpha}}{\sigma_{i}^2} \right)^2 > 0$$

(11)

for $|x|^2 = \sum_{\alpha} x_{\alpha}^2 = 1$. The Hessian is thus positive definite everywhere and the optimal solution is unique.

A possible concern is that the optimal solution is on the boundary of the simplex, i.e., $w_{\alpha} = 0$ for some $\alpha$, because the Kullback-Leibler entropy is bounded. One might then not be able to use gradient-based methods without modification. However, because of the non-analytical character of the logarithm, the partial derivatives of $L$ with respect to every $w_{\alpha}$ diverge to negative and positive infinity at $w_{\alpha} = 0$ and 1, respectively, and are monotonic in between. Therefore, the optimal solution is contained within the simplex, not on its surface.

Another concern is that to find the unique optimal solution, we have to take into account the constraints acting on the weights given by eqs 5 and 6. One could optimize the log-posterior given by eq 8 using algorithms for constrained optimization like LBFGS-B that take advantage of the gradient.

To optimize the log-posterior via log-weights

$$L = \theta \left( \langle g \rangle - \langle G \rangle + \ln \frac{s_0}{s} \right) + \frac{1}{2} \left| \tilde{y} w - \tilde{Y} \right|^2$$

(16)

we have

$$\frac{\partial L}{\partial g_{\mu}} = w_{\mu} \theta (g_{\mu} - \langle g \rangle - G_{\mu} + \langle G \rangle)$$

$$+ w_{\mu} \sum_{i=1}^{M} \frac{(y_{i} - Y_{i}) (y_{i}^{\mu} - \langle y_{i} \rangle)}{\sigma_{i}^{2}}$$

(13)

where $G_{\alpha} = \ln w_{\alpha}^0$ and $g = \sum_{\alpha} w_{\alpha} g_{\alpha}$ indicates the average over the reweighted ensemble. We simplified the expressions by taking advantage of the normalization condition.

Importantly, we need to minimize $L$ only with respect to the $N-1$ variables $g_{\mu} (\mu = 1, \ldots, N-1)$. A starting point of a gradient-based minimization of $L$ could be the normalized prior $w_{\alpha}^0$, corresponding to $g_{\alpha} = \ln (w_{\alpha}^0 / w_{N}^0) = G_{\alpha} - G_{N}$.

In a practical implementation, a procedure to evaluate $L$ and its gradients, called with $g_{\mu} (\mu = 1, \ldots, N-1)$ as arguments, would

1. define $g_{N} = 0$,

2. evaluate $w_{\alpha}$ according to eq 12 for $\alpha = 1, \ldots, N$,

3. evaluate $L$ according to eq 8 or eq 16 below, and

4. evaluate the gradient according to eq 13.

Both $L$ and its gradient can be evaluated efficiently using vector-matrix operations. Given the $g_{\alpha}$, we define $v_{\alpha} = e^{g_{\alpha}}$, $s = \sum_{\alpha} v_{\alpha} = \sum_{\alpha} e^{g_{\alpha}}$, $s_0 = \sum_{\alpha} e^{G_{\alpha}}$, and $w_{\alpha} = v_{\alpha} / s$ (all being efficiently evaluated in vector form). The averages can be calculated as vector dot products:

$$\langle g \rangle = g \cdot w$$

$$\langle G \rangle = G \cdot w$$

(14)

(15)

We then have

$$L = \theta \left( \langle g \rangle - \langle G \rangle + \ln \frac{s_0}{s} \right) + \frac{1}{2} \left| \tilde{y} w - \tilde{Y} \right|^2$$

where $\tilde{y}$ is an $M \times N$ matrix with components $y_{\alpha} = y_{i}^{\alpha} / \sigma_{i}$, and $\tilde{Y}$ is a vector with $M$ components $Y_{i} / \sigma_{i}$ that can be precalculated.

To evaluate the gradient, the averages in eq 13 can be evaluated as dot products. The first part on the right hand side of eq 13 can then be evaluated as in-place vector operation. The second
part can also be evaluated by a combination of matrix-vector multiplication (for \( h_i \)), vector dot products (for the sum over \( i \)), and in-place vector operations (for the different \( \mu \)).

### 2.3 Optimization via Generalized Forces

We showed previously\(^{20}\) that the weights at the maximum of the log-posterior can be expressed in terms of generalized forces

\[
F_k = -\frac{\langle y_k \rangle - Y_k}{\theta \sigma_k^2},
\]

as

\[
w_\alpha = \frac{w_\alpha^0 \exp \left( \sum_j y_j^\alpha F_j \right)}{\sum_\gamma w_\gamma^0 \exp \left( \sum_i y_i^\gamma F_i \right)}.
\]

Note that these forces correspond to Lagrange multipliers in closely related maximum entropy approaches to ensemble refinement.\(^{28,31,32}\) In many practical cases, we have fewer observables than weights, \( M \ll N \). In such cases, one may want to take advantage of eq 18 and minimize \( L \) with respect to the \( M \) generalized forces \( F_k \) instead of the \( N \) weights. By the chain rule, we find for the gradients

\[
\frac{\partial L}{\partial F_k} = \sum_\alpha \frac{\partial L}{\partial w_\alpha} \frac{\partial w_\alpha}{\partial F_k}.
\]

We have

\[
\frac{\partial w_\alpha}{\partial F_k} = w_\alpha (y_k^\alpha - \langle y_k \rangle)
\]

where

\[
\langle y_k \rangle = \sum_\gamma w_\gamma y_k^\gamma
\]

The gradient with respect to the generalized forces becomes

\[
\frac{\partial L}{\partial F_k} = \sum_\alpha \left[ \theta \left( \ln \frac{w_\alpha}{w_\alpha^0} + 1 \right) + \sum_i \frac{y_i^\alpha (\langle y_i \rangle - Y_i)}{\sigma_i^2} \right] \times w_\alpha (y_k^\alpha - \langle y_k \rangle)
\]

In a numerical minimization of \( L \) with respect to the \( M \) generalized forces, one would thus at each iteration step

1. calculate the current weights \( w_\alpha \) from the forces according to eq 18;
2. evaluate \( L \) according to eq 8 or eq 16;
3. evaluate the gradient according to eq 22.

Equations 8, 22, and 18 can be evaluated efficiently by using vector-matrix methods in NumPy etc., using pre-calculated vectors of intermediates. However, for large \( M \times N \), care should be taken to minimize the memory requirements by avoiding \( M \times N \) matrices other than \( y_i^\alpha \).

### 2.4 Optimization Strategies

Small \( \theta \) values are more challenging than large \( \theta \) values because the optimal weights will deviate more from the reference weights. In practice, we usually do not know how to set \( \theta \) a priori. In such cases, we recommend to perform an L-curve analysis.\(^{33}\) In an L-curve or elbow plot, we plot \( \chi^2 \) or the reduced chi-square value \( \chi^2/M \) as a function of the relative entropy \( S \) for the optimal solutions at different \( \theta \) values. The \( \chi^2 \) values will decrease with decreasing relative entropy and we can choose a \( \theta \) value corresponding to the elbow in this plot.

Finding optimal solutions for a series of \( \theta \) values also has the advantage that we can use the more rapidly converging optimal solutions at large \( \theta \) values as starting points for optimizations at smaller \( \theta \) values.

### 3 Methods

#### 3.1 Implementation

With the analytical expressions for gradients in the log-weights and forces formulations derived above, we can take advantage of highly optimized gradient-based optimization methods. The BioEn/optimize package, which can be downloaded from [https://github.com/bio-phys/BioEn/](https://github.com/bio-phys/BioEn/), provides Python and C implementations of the log-posterior and its gradient for both methods and a selection of different gradient-based optimizers and implementations.
The reference implementation is based on Python and on the packages NumPy and SciPy in particular. The log-posterior and its derivatives are written in NumPy notation, and the BFGS minimizer from SciPy is used to compute the minimum. Thanks to the fact that NumPy is typically linked to high-performance mathematical libraries such as MKL, the Python-based implementation is capable of exploiting vectorization and multi-threading on state-of-the-art hardware. On the other hand there is some overhead associated with NumPy related to the use of temporary buffers during expression evaluation.

To improve the performance, we provide C-based implementations of the log-posterior functions and their derivatives, largely avoiding temporary buffers by using explicit loops to implement the expressions. OpenMP directives are used to explicitly leverage vectorization and thread parallelization. The Python interfaces are written in Cython. While these kernels are significantly faster than the NumPy-based code there is still some overhead when the BFGS minimizer from SciPy is used because it is written in plain Python.

To eliminate the bottleneck caused by the SciPy minimizer we have implemented a Cython-based interface to the multidimensional minimizers of the GNU Scientific Library (GSL), i.e., conjugate gradient, BFGS, and steepest descent minimizers. In doing so, the minimization is performed completely in the C layer without any overhead from the Python layer. Additionally, the C implementation of Jorge Nocedal’s Fortran implementation of the limited-memory BFGS algorithm by Naoaki Okazaki (https://github.com/chokkan/liblbfgs) can be used.

A test suite is provided to check the implementations against each other. During code development work we noticed that performing parallel reductions can lead to numerically slightly different results. The reason is that parallel reductions introduce non-deterministic summation orders such that round-off errors vary between runs. Therefore, we also provide parallelized C kernels where we eliminated any non-reproducibility effects.

3.2 Simulation Details

Ala-5 was simulated at pH 2, using the AMBER99SB*-ildn-q force field matching the experimental solution conditions. To describe the protonated C-terminus at a low pH, we took partial charges from the protonated aspartate sidechain. Excess charges were distributed across the C-terminal residue. The simulations of Ala-5 were run for 1 µs using simulation options previously described. J-couplings were calculated as in previous work for the 50000 structures used for the BioEn reweighting. Chemical shifts were calculated with SPARTA+ using MDTraj. MD simulations were analyzed using MDAnalysis.

4 Results

We first investigate the stability and accuracy of the optimization methods using log-weights and generalized forces by applying them to synthetic data. We then refine molecular dynamics simulation ensembles for Ala-5 using J-couplings.

4.1 Accuracy and Performance of Optimization Methods

We investigate how accuracy and efficiency of the log-weights and forces methods depend on the size of the ensemble \( N \) and the number of data points \( M \) using synthetic data. To generate a data set, we drew \( M \) experimental values \( Y_i \) from a Normal distribution, i.e., \( Y_i \sim \mathcal{N}(0,1) \). We generated calculated observables \( y_{ia} \) by drawing Gaussian numbers from \( \mathcal{N}(Y_i + 1, 2) \), where the offset of 1 mimics systematic deviations due to force field inaccuracies. We set the experimental error for all data points to \( \sigma = 0.5 \). For each combination of five \( M \)-values, \( M = 10^2, 316(\approx 10^{2.5}), 10^3, 3162(\approx 10^{3.5}), \) and \( 10^4 \), and nine \( N \)-values, \( N = 10^2, 316(\approx 10^{2.5}), 10^3, 3162(\approx 10^{3.5}), 10^4, 31623(\approx 10^{4.5}), 10^5, 316228(\approx 10^{5.5}), \) and \( 10^6 \),
Figure 1: Scatter plot of the optimal reduced $\chi^2$ and the optimal relative entropy $S$ obtained with the log-weights method (circles) and the forces method for $5 \times 9 = 45$ values of $(M, N)$ and $\theta = 0.01$. For each value of $(M, N)$ we show results for four synthetic data drawn at random as specified in the text. Crosses on top of circles indicate excellent agreement of the two methods. Optimal values for the four data sets for a specific $(M, N)$ can be visually identified as clusters, especially for large $N$.

Figure 2: Cumulative distribution functions of Pearson’s correlation coefficient $r$ of the optimized weights obtained with the log-weights and forces methods with respect to the most optimal weights found in optimizations with different parameters for the LBFGS algorithm. The lowest $r$ values for particular $(N, M)$ combinations are shown in Figure 3.

we generated randomly four sets, giving us $5 \times 9 \times 4 = 180$ data sets in total.

To fully define the optimization problem, we chose uniform reference weights $w^0_\alpha = 1/N$ and a value for the confidence parameter $\theta = 0.01$. The latter expresses little confidence in our reference ensemble, such that the optimal weights will be significantly different from the reference weights, rendering this optimization more challenging than for large values of $\theta$. We minimize the negative log-posterior $L$ given by eq 8 for each data set using the log-weights and forces methods.

The efficiency and accuracy of gradient-based optimization methods depends strongly on their detailed parameterization. Here, we present results for the limited-memory BFGS (LBFGS) algorithm.$^{36,37}$ Due to its memory efficiency, we can refine larger ensembles using more data points compared to other algorithms like BFGS or conjugate gradients. Specifically, we explored the effect of the choice of the line search algorithm and the convergence criteria on the convergence behavior. We found that using the backtracking line search algorithm applying the Wolfe condition$^{45,46}$ in connection with a convergence criterion acting on the relative difference of the log-posterior with respect to a previous value (10 iterations before the current one) strikes the best balance between accuracy, efficiency, and robustness. We used these parameters to obtain the results we show in the following. From all optimal solutions found in our exploration of the parameter space of the LBFGS algorithm, we chose for each data set the solution with the lowest negative log-posterior to compare with. We call these solutions the most optimal solutions in the following.

To characterize the optimization problem for the synthetic data sets considered here, we plot the optimal reduced $\chi^2$ value as a function of the optimal relative entropies $S_{KL}$ in Figure 1. The larger the value of the relative entropy $S_{KL}$, the more the optimal weights differ from the reference weights and the more challenging is the optimization problem. In general, we found that the optimal values for the log-weights and forces methods agree well with each other. Due to the nature of the synthetic data sets, results for individual $(M, N)$ can be visually identified as clusters, especially for large ensemble sizes $N$. Note that for the data sets considered here, the clusters for large $N$ pose more challenging optimization problems because the optimal
weights are further from the initial weights.

The optimal weights obtained with the two methods are highly correlated and correlate excellently with the most optimal weights found in our exploration of parameter space of the LBFGS algorithm. We quantify these correlations using Pearson’s correlation coefficient \( r \), which for two set of weights \( w_\alpha^{(1)} \) and \( w_\alpha^{(2)} \) is given by

\[
\rho = \frac{\sum_\alpha \left( w_\alpha^{(1)} - N^{-1} \right) \sum_\gamma \left( w_\gamma^{(2)} - N^{-1} \right)}{\sqrt{\sum_\alpha \left( w_\alpha^{(1)} - N^{-1} \right)^2 \sum_\gamma \left( w_\gamma^{(2)} - N^{-1} \right)^2}}.
\]

(23)

We find that the cumulative distribution functions of the correlation coefficient for the forces and log-weights methods with respect to the most optimal weights found are strongly peaked at \( r = 1 \) (see Figure 2). For the forces method, 91% of all samples have a correlation coefficient of \( r > 0.99 \). For the log-weights method the peak at \( r = 1 \) is even narrower as 95% of all samples have a correlation coefficient of \( r > 0.99 \). However, the log-weights solutions of fewer than 10 out of 180 samples have a correlation coefficient of \( r < 0.9 \) and thus show poorer correlation with the most optimal weights.

A more detailed analysis of the accuracy shows that the log-weights method performs not as well in cases where the ensemble size is much larger than the number of data points, \( N \gg M \). To quantify the accuracy, we cal-
calculate the difference in log-posterior $\Delta L$ obtained with the forces and log-weights methods to the most optimal negative log-posterior $L^{(\text{opt})}$ found. An average over all samples for given $M$ and $N$ indicates that the forces method performs well (see Figure 3, top left). Only when $M \approx N$ we find occasional small deviations from the most optimal values. The log-weights method performs excellently for $M \gg N$, but not as well where $N \gg M$. This behavior is also reflected in the minimum value of the correlation coefficients over the four random samples at given $M$ and $N$ (see Figure 3, bottom).

![Figure 4: Run times for the log-weights (circles) and forces (crosses) optimization methods as a function of ensemble size $N$ for different numbers of data points $M = 100, 1000, 10000$ (in green, orange, and blue, respectively). Run times have been averaged (bold symbols) over four different synthetic data sets each (light symbols).](image)

For the chosen convergence criterion and line search algorithm, the log-weights method is computationally more efficient than the forces method (Figure 4). We performed benchmark calculations on a single node with two E5-2680-v3 CPUs, 12 cores each, and 64 GB RAM using OpenMP. For the largest system considered, $(N, M) = (10^6, 10^5)$ we used a machine with identical CPUs but 128 GB RAM. In Table 1, we summarize the average run times for the largest ensemble size considered here ($N = 10^6$). For $M = 100$ the log-weights methods is $\sim 20$ times faster than the forces method (30 sec versus 4 min on a single node; see Table 1).

| $M$   | run time [min] | minimal/avg. $r$ |
|-------|----------------|------------------|
|       | log-weights    | forces           | log-weights    | forces           |
| $10^2$ | 0.2            | 4                | 0.63/0.78      | 1.00/1.00       |
| $10^3$ | 1.5            | 7                | 0.98/0.99      | 1.00/1.00       |
| $10^4$ | 48             | 140              | 1.00 /1.00     | 0.98/0.99       |

In conclusion, for the chosen convergence criterion and line search algorithm, optimization using the LBFGS algorithm is stable, efficient, and accurate for both the forces method and the log-weights method. In cases where the ensemble size is much larger than the number of data points, $N \gg M$, the forces method is more accurate but also less efficient. In cases where $N \approx M$, the log-weights method is both more efficient and more accurate than the forces method. The BioEn optimization library has been written to make it easy and straightforward not only to choose from a variety of optimization algorithms, but also to fine tune the chosen optimization algorithms to further improve accuracy and/or efficiency.

### 4.2 Refinement of Ala-5 using J-Couplings

As a realistic example for a biomolecular system, we have conducted BioEn refinement of the disordered peptide penta-alanine (Ala-5) against NMR J-couplings. The Ala-5 model system is simple enough that well converged simulations can be obtained straightforwardly. Nevertheless, it displays much of the complexity encountered in MD simulations of intrinsically disordered proteins with a myriad of shallow free energy minima. Hence, details of the force field matter a lot for such systems and simulations do not provide results at a level routinely achieved for well-ordered proteins. NMR observables such as J-couplings, which report on
dihedral angle equilibria, provide accurate information on disordered systems.\textsuperscript{48}

We assessed the quality of a 1 \( \mu \)s simulation of Ala-5 with the AMBER99SB*-ildn-q force field by comparison to experimental J-couplings.\textsuperscript{38} J-couplings were calculated from the MD trajectory using the Karplus parameters from the original publication\textsuperscript{38} and two sets of Karplus parameters determined from DFT calculations (DFT1 and DFT2).\textsuperscript{49} The DFT2 parameters were used to define the AMBER99SB*-ildn-q force field and hence we initially focused on this set of Karplus parameters.

Even without refinement, the MD simulation gives very good agreement with the experimental J-couplings with \( \chi^2/M \approx 1.0 \) (1.1 and 0.8 for original and DFT1 Karplus parameters, respectively) using the error model of ref 40. For uncorrelated errors, \( \chi^2/M < 1 \) would signify agreement within the experimental uncertainty on average. However, a closer inspection of measured and calculated J-couplings shows that there are systematic deviations. For the \( ^3J_{\text{HNH}} \) and \( ^3J_{\text{HOC}} \) couplings, which report on the \( \phi \)-dihedral angle equilibrium, the simulations predict larger couplings than in experiments, respectively (Figure 5A and C). In addition, for the \( ^2J_{\text{NCA}} \) couplings, which are sensitive to the \( \psi \)-dihedral angle equilibrium, couplings calculated from simulations are all smaller than the experimental couplings (Figure 5G).

With BioEn reweighting, we refined the weights of 50,000 structures from the 1 \( \mu \)s simulation of Ala-5 against 28 experimental J-couplings. Optimizing the effective log-posterior at different values of the confidence parameter \( \theta \) (Figure 6A) we see the expected drop in \( \chi^2 \) as \( \theta \) is decreased. At small values of \( \theta \) we find only marginal improvements in \( \chi^2 \), but start to move away from the reference weights as indicated by a substantial increase in the relative entropy. At \( \theta = 6.65 \), we find a good compromise between reducing \( \chi^2 \) and staying close to the reference weights. The agreement with experiment increased or stayed the same for all J-couplings (Figure S6) except for \( ^3J_{\text{HNC}} \) and \( ^3J_{\text{HNC}B} \) of residue 1 for which the already very good agreement got somewhat worse. The overall improvement demonstrates that the different experiments are consistent with each other. In particular, for the \( ^3J_{\text{HNH}} \) (Figure 5A) and \( ^3J_{\text{HOC}} \) (Figure 5C) couplings, which report on the \( \phi \) dihedral angle, and the \( ^2J_{\text{NC}} \) couplings (Figure 5G), reporting on the \( \psi \) dihedral angles, systematic deviations from the experiment disappear with the refinement. The changes in the weights are associated with an entropy \( S_{KL} \approx 0.5 \) (Figure 6A). The weights of most structures were changed only slightly by the reweighting. In the optimal BioEn ensemble, the most important 2\% of structures constitute \( \approx 50 \% \) of the ensemble (Figure 6B). The weights of these structures approximately double with the refinement. After refinement \( \approx 20 \% \) of the structures contribute negligibly to the ensemble, with weights close to zero. As expected, the optimal weights from the log-weights or generalized forces methods were highly correlated (Figure S1), confirming the equivalence of the two methods to solve the BioEn reweighting problem. The run times of the forces and log-weights optimizations for all \( \theta \) values are comparable, at 42 s and 33 s on a standard workstation, respectively.

The polyproline-II (ppII) conformation at \( \phi \approx -60^\circ \) and \( \psi \approx 150^\circ \) becomes more populated in the optimal ensemble (Figure 7D), irrespective of the choice of Karplus parameters. The shift to ppII is in agreement with the original analysis of the J-couplings for Ala-5,\textsuperscript{38} where it was concluded that the ppII state dominates the conformational equilibrium, and with infrared (IR) spectroscopy.\textsuperscript{50} The same conclusion was drawn from refining Ala-3 MD simulation ensembles against 2D-IR data.\textsuperscript{51} The \( ^3J_{\text{CC}0} \) coupling for residue 2 has been highlighted as potentially spurious by Best et al.\textsuperscript{52} because the reported coupling is atypical for a polyalanine. Leaving out this observable from the BioEn refinement results in an essentially unchanged refined ensemble (Figure S5). Using alternative Karplus parameters to calculate the J-couplings (Figure S2) also leads to a shift to the ppII state (original and DFT1 in Figures S3 and S4, respectively) and in all cases does the ppII state become more favorable at the expense of \( \alpha \)-helical like conformations (Figure S5). For the
Figure 5: Comparison of J-couplings measured by NMR$^{38}$ (black squares) and calculated from MD simulation with the AMBER99SB*-ildn-q force field (red squares) and the optimal BioEn ensemble (blue circles, $\theta = 6.65$). The DFT2 set of Karplus parameters was used to calculate J-couplings.
Figure 6: BioEn optimization for Ala-5. (A) L-curve analysis to determine the optimal value of the confidence parameter θ by plotting χ² as a function of S KL for different values of θ. (B) Cumulative weight summed over rank-ordered w_α for the uniformly distributed reference weights w_0^α (red) and for optimized weights (blue) at θ = 6.65 with S KL ≈ 0.5.

Figure 7: Ala-5 Ramachandran maps. (A) Free energy surface G(ϕ, ψ) = − ln p(ϕ, ψ) from MD simulation with the AMBER99SB*-ildn-q force field averaged over the central residues 2-4. (B) Ramachandran plot for Ala residues outside of regular secondary structure from the PDB. (C) Free energy surface for the optimal BioEn ensemble with DFT2 Karplus parameters. (D) Free energy differences between initial ensemble and the optimal BioEn ensemble.
original Karplus parameters we also find a reduction in \( \beta \)-strand like conformations and an even larger ppII population than for DFT1 and DFT2. While the choice of Karplus parameter model somewhat affects the optimal ensemble, the overall conclusions are robust.

The ensemble refinement improves and preserves the agreement with experimental data not included in the refinement, PDB statistics, and the experimental chemical shifts. The distribution of \( \phi \) and \( \psi \) angles for Ala residues outside of regular secondary structure from the Protein Data Bank,\(^{11}\) while clearly not reflective of the structure of a specific disordered protein in solution, provides a measure of conformational preferences of disordered proteins. Indeed, the BioEn reweighting of the \( \alpha \) and ppII conformations leads to a Ramachandran plot agreeing more closely with the PDB statistics, with a large reduction in the population of left-handed \( \alpha \)-helical conformations as is apparent from Figure 7D, Figure S3 and Figure S4. No information from PDB statistics was included in the refinement and the improved agreement with an independent data set is encouraging.

As a second independent data set, which was not included in the BioEn refinement, we compare the experimental chemical shifts for Ala-5\(^{38}\) to the initial ensemble and the optimal ensemble. The chemical shifts predicted by SPARTA+\(^{41}\) are within the prediction error before and after ensemble refinement (Figure S7). The comparison shows that for Ala-5 (1) chemical shifts cannot be used to refine the ensemble because the initial ensemble already agrees with experiment within the large prediction error and (2) ensemble refinement either improves or leaves unchanged predictions for observables not included the refinement.

The BioEn reweighting leads to a better description of the disordered peptide Ala-5 and highlights the trade-offs inherent even in the most advanced force fields. Current fixed-charge protein force fields underestimate the cooperativity of the helix-coil equilibrium\(^{53}\) because force fields describe the formation hydrogen bonds relatively poorly. To compensate for the lack of cooperativity of helix formation, the formation of \( \alpha \)-helices was favored by the 'star'

correction to the \( \psi \) torsion potential to define a force field balanced between helix and coil conformations. The rebalancing of the AMBER force field\(^{53}\) enabled the folding of both \( \alpha \)-helical and \( \beta \)-sheet proteins.\(^{54}\) Here BioEn reweighting compensates for an adverse effect of the overall very successful rebalancing of the AMBER force field, i.e., the overestimation of the helix content for short peptides such as Ala-5. BioEn reweighting can thus serve as a system specific correction to the force field, which is a promising avenue to tackle systems such as intrinsically disordered proteins where the details of the force field are critical.

## 5 Discussion

Reweighting relies on good coverage of the conformational space such that the true ensemble underlying the experimental data is a sub-ensemble of the simulation ensemble.\(^{55}\) In coarse-grained simulations, sampling is efficient and the free-energy landscapes are smooth such that usually good coverage can be achieved. In atomistic simulations, where sampling is more expensive and the free energy landscape is rougher, we often have to apply enhanced sampling methods to obtain good coverage. Independent of the details of the enhanced sampling method and with or without steering by experimental, one can use binless WHAM\(^{39,56}\) or MBAR\(^{57}\) to obtain the reference weights of the unbiased ensemble, which serve as input for ensemble refinement by reweighting.\(^{20}\)

Here, we demonstrated that even without applying enhanced sampling methods, refinement of fully atomistic trajectories of penta-alanine using J-couplings alleviates deficiencies in the force field and leads to better agreement not only with the NMR data but also with expectations from experimental structures for proteins. These results indicate that ensemble refinement via reweighting is a promising route for highly flexible systems such as nucleic acids\(^{28}\) and intrinsically disordered proteins. For such systems, the number of accessible states can be enormous and consequently even small inaccuracies in the simulation force fields can lead to...
a poor representation of the experimental ensemble.

Importantly, ensembles do not have to be generated by simulations to be amenable to ensemble refinement via reweighting. For example, in the analysis of EPR experiments like DEER, libraries of the rotameric stats of spin labels are used. For a specific residue, one selects from this library all rotameric states that do not have steric clashes with the protein structure. However, the interactions of the spin label with its surroundings can make some rotameric states in this ensemble more preferable than others. To account for this uncertainty, one can perform a BioEn refinement using the DEER data and the ensemble of rotameric states. This procedure has been used recently to resolve Ångstrom-scale protein domain movements.

To integrate experimental results, we often have to take nuisance parameters into account. For refining against SAXS intensities, we have to consider an unknown scaling parameter and often use an additive constant to account for inelastic scattering and, to a first approximation, for differences in the contrast. Using DEER data, we have to determine the modulation depths. We can include such nuisance parameters in the optimization either directly (by minimizing $L$ simultaneously with respect to the weights and nuisance parameters) or iteratively. In the iterative approach, we perform (1) a least-chi-squared fit of the calculated ensemble averages determined by the current weights to the experimental data sets with the corresponding nuisance parameters as fit parameters. We have to perform one fit for every experimental method providing data. (2) With these fitted values of the nuisance parameters we adjust the calculated observables $y^0$. These enter another round of optimization from which we obtain the optimal weights given the values of the nuisance parameters. (3) We use these weights for another round starting with step 1 until convergence is achieved. Note that instead of using least-chi-squared fits, one can also include priors acting on the nuisance parameters in both the direct and iterative formulations.

Interestingly, ensemble refinement by reweighting offers a way to quantify the agreement between simulations and experiment. After reweighting, we can make a quantitative statement of how much we would have had to change the simulated ensemble, expressed by the relative entropy or Kullback-Leibler divergence to be able to obtain agreement with experiment. The quantification of the agreement between simulation and experiment can also be used to identify and correct deficiencies in molecular dynamics force fields. In a perturbative formulation, one can seek force field corrections that capture the weight change.

BioEn accommodates a wide range of error models. With the gradients of the BioEn log-posterior presented here for Gaussian error models, with and without correlation, we already cover a large range of experimental methods. Moreover, in many cases the Gaussian error model can be used to efficiently obtain an initial estimate for the optimal weights. These estimates can then be used as initial weights for an optimization using a more accurate error model but perhaps a less efficient optimization method. The log-weights and forces methods we presented here to calculate the gradients of the BioEn log-posterior are both general and applicable also to non-Gaussian likelihoods with minor adaptations. We provide an open-source implementation at https://github.com/bio-phys/BioEn/optimize at no cost under the GPLv3 license.

Acknowledgement We thank Dr. Sandro Bottaro and Prof. Kresten Lindorff-Larsen for useful discussions. We acknowledge financial support from the German Research Foundation (CRC902: Molecular Principles of RNA Based Regulation) and by the Max Planck Society.

A Gradients of the Log-Posterior for Correlated Gaussian Noise

In the following, we derive expressions for the gradients of the negative log-posterior given by eq 1 and correlated Gaussian noise for the likelihood in log-weights and forces formulations.
For correlated Gaussian noise, the likelihood is given by \( P(\mathcal{y}^{(\text{obs})}_i | \mathbf{w}) \propto \exp(-\chi^2/2) \), with
\[
\chi^2 = \mathbf{x}^\top \mathbf{S}^{-1} \mathbf{x}.
\] (24)

The components of the vector of deviations \( \mathbf{x} \) are given by
\[
x_i = \sum_{\alpha=1}^{N} w_\alpha y_\alpha^i - Y_i = \sum_{\alpha=1}^{N} w_\alpha x_\alpha^i
\] (25)
where we introduced \( x_\alpha^i = y_\alpha^i - Y_i \). \( \mathbf{S} \) is the symmetric covariance matrix of the statistical errors. We denote the \( ij \) elements of its inverse as \( S_{ij}^{-1} \). We may write
\[
\chi^2 = \sum_{i=1}^{M} \sum_{j=1}^{M} S_{ij}^{-1} x_i x_j.
\] (26)

In the log-weights formulation, the gradient of \( \chi^2 \) is given by
\[
\frac{\partial \chi^2}{\partial g_\gamma} = w_\gamma \sum_{i,j} S_{ij}^{-1} \left( x_i^\gamma x_j + x_j^\gamma x_i - 2x_i x_j \right).
\] (27)

where \( \sum_{i,j} = \sum_{i=1}^{M} \sum_{j=1}^{M} \). The gradient of the negative log-posterior with respect to the log-weights for correlated Gaussian noise is then given by
\[
\frac{\partial L}{\partial g_\gamma} = w_\gamma \theta (g_\gamma - \langle g \rangle - G_\gamma + \langle G \rangle) + \frac{w_\gamma}{2} \sum_{i,j} S_{ij}^{-1} \left( x_i^\gamma x_j + x_j^\gamma x_i - 2x_i x_j \right).
\] (28)

For uncorrelated noise the covariance matrix is diagonal, \( \mathbf{S} = \text{diag}\{\sigma_1^2, \ldots, \sigma_M^2\} \), and the gradient of \( \chi^2 \) simplifies to
\[
\frac{\partial \chi^2}{\partial g_\gamma} = 2 \sum_{i} \frac{x_i (x_i^\gamma - x_i)}{\sigma_i^2}
\] (29)
such that we recover eq 13 as expected.

In the forces formulation, we apply the chain rule, eq 19, and calculate first the gradient of \( \chi^2 \) with respect to \( w_\alpha \). Because of the normalization condition \( \sum_{\alpha=1}^{N} w_\alpha = 1 \), we only have \( N-1 \) independent variables. Thus, we set
\[
w_N = 1 - \sum_{\alpha=1}^{N-1} w_\alpha
\] (30)
and write
\[
x_i = \sum_{\alpha=1}^{N-1} w_\alpha x_\alpha^i + \left( 1 - \sum_{\alpha=1}^{N-1} w_\alpha \right) x_N^i.
\] (31)

Using that \( \partial x_i / \partial w_\gamma = x_i^\gamma - x_i^N \) for \( \gamma \neq N \) and \( \partial x_i / \partial w_N = 0 \), we obtain for the gradient of eq 26 with respect to \( w_\gamma \neq N \)
\[
\frac{\partial \chi^2}{\partial w_\gamma} = \sum_{i,j} S_{ij}^{-1} \left( (x_i^\gamma - x_i^N) x_j + (x_j^\gamma - x_j^N) x_i \right)
\] (32)
and \( \partial \chi^2 / \partial w_N = 0 \). For uncorrelated noise, eq 32 simplifies to
\[
\frac{\partial \chi^2}{\partial w_\gamma} = 2 \sum_{i=1}^{M} (x_i^\gamma - x_i^N) x_i.
\] (33)

Next, we derive the gradient of the entropy \( S = \sum_{\alpha=1}^{N} w_\alpha \ln \left( w_\alpha / w_\alpha^0 \right) \), which we rewrite as
\[
S = \sum_{\alpha=1}^{N-1} w_\alpha \ln \frac{w_\alpha}{w_\alpha^0} + w_N \ln \frac{w_N}{w_N^0}.
\] (34)

The derivative of the first term with respect to \( w_\gamma \) is given by
\[
\frac{\partial}{\partial w_\gamma} \sum_{\alpha=1}^{N-1} w_\alpha \ln \frac{w_\alpha}{w_\alpha^0} = \ln \frac{w_\gamma}{w_\gamma^0} + 1
\] (35)
The derivative of the second term with respect to \( w_\gamma \) is given by
\[
\frac{\partial}{\partial w_\gamma} \left( \ln w_N - \ln w_N^0 \right) = - \ln \frac{w_N}{w_N^0} - 1
\] (36)
where we used that \( \partial w_N / \partial w_\gamma = -1 \) and \( \partial \ln w_N / \partial w_\gamma = -1 / w_N \). Thus, we obtain
\[
\frac{\partial S}{\partial w_\gamma} = \ln \frac{w_\gamma}{w_\gamma^0} - \ln \frac{w_N}{w_N^0}
\] (37)
and \( \partial S / \partial w_N = 0 \).

For correlated Gaussian noise, the generalized
forces are given by

\[ F_i = -\frac{1}{\theta} \sum_{j=1}^{M} S_{ij}^{-1} f_j \]  

(38)

where \( f_j = \langle y_j \rangle - Y_j \). These forces enter eq 20 for the weights. To calculate the gradient of \( L \) with respect to the forces we use the chain rule, eq 19, and eq 18. Consequently,

\[
\frac{\partial L}{\partial F_k} = \sum_{\gamma=1}^{N-1} \left[ \theta \left( \ln \frac{w_\gamma}{w_\gamma^0} - \ln \frac{w_N}{w_N^0} \right) + \right. \\
\frac{1}{2} \sum_{i,j} S_{ij}^{-1} \left[ (x_i^\gamma - x_i^N) x_j + (x_j^\gamma - x_j^N) x_i \right] \\
\times w_\gamma (x_k^\gamma - x_k) . \\
(39)
\]

For uncorrelated noise, we recover eq 22, which in the notation used here takes on the form

\[
\frac{\partial L}{\partial F_k} = \sum_{\gamma=1}^{N-1} \left[ \theta \left( \ln \frac{w_\gamma}{w_\gamma^0} - \ln \frac{w_N}{w_N^0} \right) + \right. \\
\frac{1}{2} \sum_{i,j} S_{ij}^{-1} \left[ (x_i^\gamma - x_i^N) x_j + (x_j^\gamma - x_j^N) x_i \right] \\
\times w_\gamma (x_k^\gamma - x_k) . \\
(40)
\]

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Supplementary Information: Efficient Ensemble Refinement by Reweighting

Jürgen Köfinger,*† Lukas S. Stelzl,† Klaus Reuter,‡ César Allande,‡ Katrin Reichel,† and Gerhard Hummer*†¶

†Department of Theoretical Biophysics, Max Planck Institute of Biophysics, Max-von-Laue-Straße 3, 60438 Frankfurt am Main, Germany
‡Max Planck Computing and Data Facility, Gießenbachstr. 2, 85748 Garching, Germany
¶Institute for Biophysics, Goethe University, Max-von-Laue-Straße 9, 60438 Frankfurt am Main, Germany

E-mail: juergen.koefinger@biophys.mpg.de; gerhard.hummer@biophys.mpg.de

1 Refinement of Ala-5 using J-Couplings

Comparison of Optimization using Generalized Forces and Log-Weights

Ensemble refinements using generalized forces and log-weights give very similar results across the full range of the confidence parameter $\theta$. The correlation of the optimal weights for Ala-5 refined against J-couplings is shown in Figure S1A. The DFT2 set of Karplus parameters was used for this comparison. Small deviations are seen at small values of $\theta$. The effective log-likelihoods from optimization with the two methods agree very well over the full range of $\theta$ values (Figure S1B).
**Figure S1**: Comparison of forces and log-weights optimization. (A) Correlation between optimal weights for different values of the confidence parameter $\theta$ using log-weights or generalized forces in the BioEn optimization problem. (B) Effective log-likelihoods from forces and log-weights optimization.

**Effect of the Choice of Karplus Parameters on Ala-5 Ensemble Refinement**

**Figure S2**: Variation of $S_{KL}$ and $\chi^2$ with $\theta$, to determine the optimal value of the confidence parameter $\theta$ for the reweighting of Ala-5 using the original (blue) and DFT1 sets of Karplus parameters (green). Values of $\theta$ of 9.43 and 5.58, respectively, provide a compromise between minimizing $\chi^2$ and small changes to the reference weights for BioEn $S_{KL}$ of about 0.5, as highlighted by squares.

BioEn ensemble refinement produced very similar trends no matter which Karplus parameters were used to calculate the J-couplings. We performed independent Ala-5 ensemble refinement with three different set of Karplus parameters: the empirical parameters\(^1\) (origi-
nal) and two set of parameters obtained from density functional theory\(^2\) (DFT1 and DFT2). For further analysis of optimization with the original and DFT1 parameter sets we picked refined ensembles with \(S_{KL} = 0.5\). Irrespective of which set of Karplus parameters we used to calculate \(J\)-couplings the polyproline-II conformation becomes more populated and the \(\alpha\)-helical like conformation less populated (Figure 3D in main text, Figure S4D and Figure S3D). The changes in the populations of the conformational states, as defined previously,\(^3\) are summarized in Figure S5. The main difference between the results with different Karplus parameters is the reduction in the \(\beta\)-strand like conformations seen when calculating the \(J\)-couplings with the original set of Karplus parameters. The left-handed helical \(\alpha_L\) conformation becomes somewhat less prominent after the refinement. Leaving out the \(^3J_{CC'}\) does not change the trends either (Figure S5).

Figure S3: Free energy surface \(G(\phi, \psi) = -\ln p(\phi, \psi)\) for Ala-5 from BioEn optimization, with \(J\)-couplings calculated with the original Karplus parameters from Graf et al.\(^1\) Free energy surface for AMBER99SB*-ildn-q for the central residues 2-4. (B) Ramachandran map for Ala residues outside of regular secondary structure from the PDB.\(^4\) (C) Free energy surfaces for the optimal BioEn ensemble. (D) Free energy differences between initial ensemble and the optimal BioEn ensemble.
Figure S4: Free energy surface \( G(\phi, \psi) = -\ln p(\phi, \psi) \) for Ala-5 from BioEn optimization, with J-couplings calculated with DFT1 Karplus parameters. Free energy surface for AMBER99SB*-ildn-q for the central residues 2-4. (B) Ramachandran map for Ala residues outside of regular secondary structure from the PDB.\(^4\) (C) Free energy surfaces for the optimal BioEn ensemble. (D) Free energy differences between initial ensemble and the optimal BioEn ensemble.

Figure S5: Populations for the conformational states for the initial ensemble and for the optimal ensembles. Here optimal ensembles for DFT1, DFT2, DFT2 (and excluding \(3J_{CC'}\) coupling) and original Karplus parameters are compared to each other and the original molecular dynamics (MD) simulations.
Agreement for Individual J-Couplings

Comparing the agreement between the simulated ensemble and experiments for individual observables (Figure S6) shows which data points drive the ensemble refinement. Here we focus at ensemble refinement using J-couplings calculated with the DFT2 of Karplus parameters. For $^{3}J_{CC'}$ (Figure S6D), $^{3}J_{HNH_{0}}$ (Figure S6A) and $^{3}J_{HaC'}$ (Figure S6C) couplings the agreement between experiment and simulations improves considerably going to the optimal ensemble at $\theta = 6.65$. $^{3}J_{CC'}$ was measured only for residue 2 of Ala-5 and $\chi^2$ was decreased from $\approx 8$ to $\approx 2$. For $^{3}J_{HaC'}$ the improvement is driven by residue 4 which fits poorly in the initial ensemble, whereas for the other residues the agreement is already very good in the initial ensemble. Some improvement in the fit was obtained for $^{2}J_{NC_{a}}$ (Figure S6G) and $^{3}J_{HNC_{a}}$ (Figure S6H), with $\chi^2$ reduced from 3 to $< 1$ and 2 to $\approx 0.5$ respectively. Only very small changes were seen for $^{1}J_{NC_{a}}$ (Figure S6F). Note that the $^{1}J_{NC_{a}}$ coupling for residue 5 is uninformative in our analysis as evidenced by the flat $\chi^2$ across the full-range of $\theta$ values. The $\psi$ dihedral angle is not defined for the terminal residue and the calculated $^{1}J_{NC_{a}}$ depends on $\psi$ in the current parameterization. For $^{3}J_{HNC_{a}}$ (Figure S6B) and $^{3}J_{HNC_{a}}$ (Figure S6E) the agreement is extremely good to start with and deteriorates somewhat with the refinement. Importantly, as discussed in the main text, the refinement removes systematic offsets for $^{3}J_{HNAS}$, $^{3}J_{HaC'}$ and $^{2}J_{NC_{a}}$.

Chemical Shifts

Experimental chemical shifts for Ala-5 from Graf et al.¹ were compared to the chemical shifts calculated from the initial and reweighted, optimal ensemble Figure S7. The error in the comparison of calculated and measured shifts is dominated by the forward model. Hence the error bars shows the root mean square error for SPARTA+⁵ predictions for the respective nuclei previously determined.
Figure S6: Comparison of the agreement with experiments for J-couplings calculated with the DFT2 set of Karplus parameters. The grey circles indicated the sum of $\chi^2$ for different residues for a type of J-couplings at given value of the confidence parameter $\theta$. The dashed line indicates $\theta = 6.65$ chosen for further analysis.
Figure S7: Ala-5 chemical shifts calculated from the initial and optimal ensemble with the DFT2 Karplus parameters.

References

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