0.1. Hybrid Hamiltonian for the enhanced sampling of protein folding

We define an ideal biasing Hamiltonian $H(B)$ leading to the real underlying partition of the system represented by a free energy landscape (FEL) along suitable restraint vectors and state that the standard forcefield parameter set describes the Hamiltonian $H(A)$ as a reference energy function. Therefore, any property $X$ as a result from the sampling using the Hamiltonian $H(A)$ leads to a corresponding probability $P$ to match a certain value:

$$P_X = f(H(A)),$$

(1)

An additional bias $H(B)$ in the energy space as conventionally applied in enhanced sampling changes the resulting probability $P'$ to match a defined target property $X$ to:

$$P'_X = f(H(A) + H(B)).$$

(2)

The additional bias $H(B)$ is applied along collective variables describing the slowest degrees of freedom of the system, for example in umbrella sampling [1] and related methods such as Metadynamics [2], conformational flooding [3] or local elevation [4]. We recently have introduced a renormalization scheme to solve the problem that the unbiased probability $P(X)$ can be strongly affected by the added bias $H(B)$ [5]. In this scheme, the applied bias is renormalized to the unbiased Hamiltonian $H(A)$ in a way that its magnitude only equals a fraction of the unbiased Hamiltonian $H(A)$ in dependency of two coupling factors $\alpha_{md}$ and $\alpha'$. Therefor, we introduce a renormalization of the unbiased Hamiltonian by the same factor, which results in a new Hamiltonian $H(C)$:

$$H(C) = \frac{H(A)}{1 + \alpha_{md}} + \frac{\alpha' |H(A)|}{|H(B)|} H(B),$$

(3)

where we distinguish between $\alpha_{md}$ as coupling parameter renormalizing the unbiased Hamiltonian and $\alpha'$, which defines the coupling of the bias to the system. Alternatively, the restrained Hamiltonian $H(B)_R$ leads to:

$$H(C)_R = \frac{H(A)}{1 + \alpha_{md}} + \frac{\alpha' |H(A)|}{|H(B)_R|} H(B)_R,$$

(4)

where the new hybrid Hamiltonian $H(C)_R$ consists of a reweighted experimental restraint component $r$. In principle, the total energy remains approximately un-affected, while other properties can be introduced through the bias $H(B)$. In other words, we generate a new probability distribution
$P''(X)$ as a result from adding a particular bias $H(B)$ in dependency of two linear coupling factors $\alpha_{md}$ and $\alpha'$:

$$P'_X = \int \left( \frac{H(A)}{1 + \alpha_{md}} + \frac{\alpha'[H(A)]}{|H(B)|} H(B) \right),$$

(5)

which also affects the probabilities $P''(X^t)$ to find the transition state $X^t$, leading to a behavior which can enhance the sampling:

$$P''_X = \int \left( \frac{H(A)}{1 + \alpha_{md}} + \frac{\alpha'[H(A)]}{|H(B)|} H(B) \right).$$

(6)

Alternatively, the restrained Hamiltonian $H(B)_R$ leads to:

$$P''_{X_R} = \int \left( \frac{H(A)}{1 + \alpha_{md}} + \frac{\alpha'[H(A)]}{|H(B)|} H(B)_R \right),$$

(7)

while the same dependency holds for the probability to find the transition state $X^t$. Through this formalism, we introduce another enhancing fragment $H(B)$ or $H(B)_R$ to the original Hamiltonian $H(A)$, without affecting the total energy of the system. As we show later, we generate an accelerating Hamiltonian $H(B)$ or $H(B)_R$ from the parallel path-increments added to the system in the form of a renormalized fluctuation. The orientation of $H(B)$ is explicitly determined based on a definition of a path-increment and acts on the system through a renormalized fluctuation. That procedure has the advantage that the propagation of the system remains ergodic as long as the coupling factors $\alpha_{md}$ and $\alpha'$ lie within a low range of approximately $< 10^{-4}$ [5]. Considering the fact that bonded interactions in the biomolecular forcefield can contribute with gradients $> 10^4 - 10^5$ kJ/mol/nm, a coupling with a factor with a magnitude of $\alpha_{md} = \alpha' = 10^{-4}$ corresponds to the order of magnitude of typical non-bonded interactions. We mention that we used parameters in the range $\alpha_{md} = \alpha' = 10^{-8}$ for DNA systems using potentials of mean force derived from PDB-data [6].

0.2. Pathway dependent biasing increments: $\nabla H(B)$

We define the bias Hamiltonian $H(B)$ used for the accelerated sampling along multiple path increments $dl_{ik}$ (for pathways $i$ and atom-indices $k$) as well as its modification to its principal components, which is adaptively changed into a bias $H(B)_R$ in dependency of a distance restraint $r$ given by experimental data. We consider that the simulated system in an equilibrium simulation propagates along a pathway with the general condition that the reduced action $L$ as function of momentum $p$ and positions $q$ remains constant [7,8]:

$$L = \int pdq = const. .$$

(8)

Along a time-dependent MD trajectory of a system exposed to non-zero fluctuations of its momenta $dp$, we rewrite the equation 8 as a time-integral:

$$L = \int_{t_0}^{t_1} \left( \frac{dp(t) dq(t)}{dt} \right) dt' = const. ,$$

(9)

where $t_0$ stands for the start and $t_1$ for the end of the simulated MD trajectory. The time-dependent integral is expressed as:

$$L = \int_{t_0}^{t_1} \left( \frac{dp(t)}{dt} dq(t) + p(t) \frac{dq(t)}{dt} \right) dt' = const. .$$

(10)

We then define a differential $dl(t)$ for a microscopic system, in which fluctuations of the momenta $dp(t)$ occur. That local change in $L(t)$ at the time $t$ is defined by:
\[
\frac{dL(t)}{dt} = \frac{d}{dt}(p(t)dq(t)) = \frac{dp(t)}{dt}dq(t) + p(t)\frac{dq(t)}{dt}.
\]  
(11)

We obtain the following differential at time \( t \):
\[
dL(t) = p(t)dq(t) + dp(t)dq(t) = (p(t) + dp(t))dq(t).
\]  
(12)

The expressions in the equations 9 to 12 relate to the standard MD-case with fluctuations in the momenta \( dp(t) \) and displacements \( dq(t) \) at times \( t \). Any arbitrary biasing technique developed to accelerate a MD simulation adds an instantaneous increment \( dL_s(t) \) to \( dL(t) \):
\[
dL'(t) = dL(t) + dL_s(t),
\]  
(13)

where we obtain a modified increment \( dL'(t) \) and additional changes in the momenta resulting from applied bias-energies affect the instantaneous action of a system in order to reach a faster convergence to the statistical average, i.e. the free energy landscape [2,9,10]. In our recent work, we used 2 biasing increments \( dL_s(t) : dL_{ab}(t) \) (adaptive bias MD) and \( dL_{\sigma}(L_{ik}(t)) \) (path-sampling) depending from 2 coupling time intervals \( \tau_1 \) (adaptive bias MD) and \( \tau_2 \) (path-sampling) in which the gradient has been evaluated [11]:
\[
dL_s(t) = dL_{ab}(t) + dL_{\sigma}(L_{ik}(t)).
\]  
(14)

We extend the formalism to the sampling within \( N_R \) multiple biases (and optionally \( N_S \) multiple simulations) and redefine the expression 9 to a multiple sampling in multiple bias-paths along \( N_R \) multiple biases, which the system can undergo simultaneously. The biased simulation then results in the expression of a modified action expression:
\[
L_m(t) = \int_{t=t_0}^{t_1} dL(t)dt' + \int_{t=t_0}^{t_1} \sum_{i} \sum_{k} (dL_{abik}(t) + dL_{\sigma ik}(L_{ik}(t))) dt' = \text{const},
\]  
(15)

where the multiple biasing pathway dependent increment \( dL_{s,m} \) is defined by:
\[
dL_{s,m} = \sum_{i} \sum_{k} (dL_{abik}(t) + dL_{\sigma ik}(L_{ik}(t))),
\]  
(16)

1. Methods

1.1. Renormalization

As given in the equations 3 and 4, we employ a renormalization of the vector \( \nabla H(B) \) or \( \nabla H(B)_R \) to the underlying ensemble [5], derived from multiple biasing increments \( \nabla H(B)_{ab} \) (adaptive bias MD) and \( \nabla H(B)_{\sigma} \) (path-sampling). We apply the relations 3 and 4 for the potential \( H(B) \) or \( H(B)_R \) acting on an individual atom, where the two parameters \( \alpha_{md} \) and \( \alpha' \) allow fluctuations of the coupled bias and the un-biased gradient around an average value:
\[
\alpha_{md} = \beta_{md}' \eta_{md}' \times (1 - \xi),
\]  
(17)

and
\[
\alpha' = \beta' \eta' \times (1 - \xi),
\]  
(18)
where $\eta_{md}'$ and $\eta'$ define the width of the fluctuation and $\xi$ is a random number $\in [0, 1]$. Using the definition:

$$\nabla H(C) = \frac{\nabla H(A)}{1 + \alpha_{md}} + \alpha' |\nabla H(A)| \nabla H(B),$$  \hspace{1cm} (19)

we add the bias $\nabla H(B)$ to the system. As mentioned before, we distinguished between $\alpha_{md}$ to describe the fluctuation of the un-biased MD-gradient and $\alpha'$ for the renormalization of the bias. Using the absolute value of $H(A)$, we can formulate the resulting modified absolute value of the hybrid Hamiltonian $|H(C)|$ of the biased system, which is expressed by:

$$|H(C)| = \frac{1}{1 + \alpha_{md}} |H(A)| + \alpha' |H(A)|$$

$$= \frac{|H(A)|}{1 + \alpha_{md}} \frac{1 + \alpha_{md} + \alpha_{md}^2}{1 + \alpha_{md}}.$$  \hspace{1cm} (20)

If $\alpha_{md} = \alpha'$ and $\alpha_{md}$ on the order of $0 < \alpha_{md} < 0.1$, then $\frac{1 + \alpha_{md} + \alpha_{md}^2}{1 + \alpha_{md}} \approx 1$, and we obtain:

$$|H(C)| \approx |H(A)|,$$  \hspace{1cm} (21)

which means that the energy remains conserved for small $\alpha_{md}$-values. The renormalization to the underlying Hamiltonian leads to a reduction of the added bias $s$ to the norm of the applied un-biased gradient. We summarize that the magnitude of the added bias $s$ derived from multiple path-dependent increments depends from the coupling parameters $\beta'_{md}$, $\beta'$ and the fluctuation parameters $\eta_{md}'$, $\eta'$. Considering the fact that bonded interactions in the biomolecular forcefield can contribute with gradients $> 10^4$ kJ/mol/nm, a coupling with a factor with a magnitude of $\beta'_{md} = \beta' = 10^{-4}$ corresponds to the order of magnitude of typical non-bonded interactions. In general, the renormalization procedure guarantees a sampling in which the occurrence of non-equilibrium configurations is prevented and has the advantage that the applied bias enhances the sampling of the system through the addition of renormalized fluctuations. That means that the underlying Hamiltonian is not overdamped through the application of a bias exceeding the un-biased fluctuations, where essential configurations are potentially not accessed. We tested the influence of the $\alpha$ parameters in simulations of SPC/E and TIP3P water, where we investigated the dielectric and the structural properties of water in dependency of the simulation parameters (see Figure 2 b and the supplementary material: Section - 2. Water simulations. supplementary Table 1, Table 2, Figures S1 and S2).

1.2. Adaptive bias MD

For the adaptive bias MD section of the algorithm, we derive a history dependent bias $\nabla H(B)_{ab}$ of the form:

$$\nabla H(B)_{ab} = \sum_{i} \sum_{k} N_{R} \gamma_{ik}(t) dL_{ab} \gamma(t) = \sum_{i} \sum_{k} N_{R} \gamma_{ik}(t) (p_{k}(t) + dp_{k}(t)) dq_{k}(t).$$  \hspace{1cm} (22)

using a number of $N_{R}$-biases in which the bias is re-evaluated within periods of $\tau_{ik}$ for the bias with an index $i$ and atom $k$. As we introduced in our previous work, we define the corresponding force $F_{k}(t)$ at time $t$, and use time-derivative of $\nabla H(B)_{ab}$:\n
$$\frac{d}{dt} H(B)_{ab} = H(B)_{ab}.$$
\[
\frac{d}{dt} H(B)_{ab} = H(B)_{ab} \\
= \sum_{i}^{N} \sum_{k}^{N} \left[ \gamma_{ik}(t) \frac{d}{dt} [(p_{k}(t) + dp_{k}(t))dq_{k}(t)] \right] \\
+ \gamma_{ik}'(t)(p_{k}(t) + dp_{k}(t))dq_{k}(t) \right]. \tag{23}
\]

As we defined in the equation 8, the added bias has to fulfill the condition that \( \lim_{t \to \infty} \langle dL_{ab}(t) \rangle_t \approx 0 \) in order to sample the system at equilibrium. That also implies that the averages of \( \gamma_{ik} \) have to fulfill that \( \langle \gamma_{ik}'(t) \rangle = 0 \) and \( \langle \frac{d\gamma_{ik}'}{dt} \rangle \approx 0 \). That relation is fulfilled if:

\[
\langle \gamma_{ik}'(t) \rangle = 0 , \tag{24}
\]

which is implemented by:

\[
\langle \gamma_{ik}(t) \rangle = \langle \gamma_{ik}'(t) \rangle = 0 , \tag{25}
\]

where \( \gamma_{ik}' \) stands for the fluctuation range with the dimension of a length \((nm^{-1})\) and \( \xi \) is a normally distributed random number with a weight equal 1. We used a constant value \( \gamma_{ik}' = 10^{-4} \) in all simulations. To enhance sampling along a history-dependent pathway in adaptive bias MD, we employ a coarsening expressed by:

\[
\frac{d}{d\tau_{1}} H(B)_{ab} = \sum_{i}^{N} \sum_{k}^{N} \left[ \gamma_{ik}(t) \frac{d}{d\tau_{1}} [(p_{k}(t) + dp_{k}(t))dq_{k}(t)] \right] \\
+ \gamma_{ik}'(t)(p_{k}(t) + dp_{k}(t))dq_{k}(t) \right]. \tag{26}
\]

By taking into account that \( \frac{d}{d\tau_{1}} \left( \gamma_{ik}'(t)(p_{k}(t) + dp_{k}(t))dq_{k}(t) \right) \approx 0 \) we use that formalism to define the differential over a finite time increments \( \tau_{1} \) to coarse-grain the dynamics and to increase the computational efficiency, which leads to an expression for the corresponding force in adaptive bias MD:

\[
\nabla H(B)_{ab}(\tau_{1}) = \frac{dH(B)_{ab}}{d\tau_{1}}d\tau_{1} \\
= \sum_{i}^{N} \sum_{k}^{N} \left[ \gamma_{ik}(t) \frac{d}{d\tau_{1}} [(p_{k}(t) + dp_{k}(t))dq_{k}(t)] \right] d\tau_{1} \\
+ \frac{d\gamma_{ik}'}{d\tau_{1}} d\tau_{1} \left( \frac{d}{dt} [(p_{k}(t) + dp_{k}(t))dq_{k}(t)] \right) \right]. \tag{27}
\]

1.3. Path sampling

In the path-sampling, we use a definition of the reactive coordinate \( \sigma_{ik}(t) \) to determine the biasing segment \( \nabla H(B)_{\sigma} \) as function of the increment \( dL_{\sigma_{ik}}(L_{ik})(t) \), which we define as [11]:

\[
\sigma_{ik}(t) = dL_{\sigma_{ik}}(L_{ik})(t) \tag{28}
\]
with $L_{ik}(t) = \int_{t_0}^{t} \left( \frac{dp_i(t)}{dt} dq_{ij}(t) + p(t) \frac{dq_{ij}(t)}{dt} \right) dt'$, equal to the path integral reached by integration till time $t$ for the bias with index $i$ and atom $k$. In cartesian coordinates : $\sigma_{ik}(t) = \{L_{x_i}(t), L_{y_i}(t), L_{z_i}(t)\}$ and $L(t) = \{ \int p_{x_i} dx_{ik}, \int p_{y_i} dy_{ik}, \int p_{z_i} dz_{ik}\}$. Along $\sigma_{ik}(t)$ a history dependent bias potential $\Phi_{ik}'$ is added in intervals of $\tau_{2_{ik}}$:

$$\Phi_{ik}' = -\frac{\partial}{\partial \sigma_{ik}} W_{ik} \prod_{l \leq t_0} \prod_{ik} \exp \left( -\frac{|\sigma_{ik} - \sigma_{ik}^{l-\tau_{2_{ik}}}|}{2\delta \sigma_{ik}^{l}} \right),$$

where the height $W_i$ and the width $\delta \sigma_i$ are conventionally parameters chosen to provide computational efficiency and an efficient exploration of the free energy $F(\nabla H(B),r)$. We define the bias component as :

$$\nabla H(B),r(\tau_{2_{ik}}) = \nabla \sigma_{ik}(t) \Phi_{ik}' .$$

That formulation constantly drives the system to explore new configurations along the variable $L_{ik}(t)$ and prevents the system to revisit conformers, which have been sampled previously. In the implementation for an efficient exploration of this space, we note that our algorithm uses the definition [2] :

$$\delta \sigma_{ik} = |t_{b_{ik}}^{t_{b_{ik}}'} - \sigma_{ik}^{t_{b_{ik}}'}| ,$$

where the times $t_{b_{ik}}$ and $t_{b_{ik}}'$ are defined by $\tau_{2_{ik}} = t_{b_{ik}} - t_{b_{ik}}'$. We apply a variable height of each Gaussian added to an individual variable :

$$W_{ik} = W \exp \left( -\Phi_{ik}' \Delta E \times \frac{|\sigma_{ik} - \sigma_{ik}^{t_{b_{ik}}'}|}{\sigma_{ik}^{t_{b_{ik}}'}} \right),$$

where $W$ and $\Delta E$ are constants [12] We applied constant values $W = 0.1$ kJ/mol ($\Delta E = 1000$ kJ/mol) in all simulations. In order to accelerate the sampling along $H(B)$ and $H(B)_{\text{ensemble}}$, we re-evaluate the principal components of $H(B)$ in order to sample the system along its slowest modes of the 2 segments $H(B)_{\text{ensemble}}$ and $H(B)_{\text{ensemble}}$. Therefore, we diagonalize the matrices $dL_{\sigma_{ik}}(L_{ik})$ and $dL_{\sigma_{ik}}(L_{ik})$. The corresponding eigenvectors with the smallest eigenvalues are used to the system [13].

2. Results : Water simulations

In this section, we discuss our results from the simulations of 2 water models : TIP3P and SPC/E, with the sampling algorithm along multiple pathways. We tested the influence of the 2 renormalization parameters $\beta$, $\eta'$ and $\eta_{\text{MD}}$ on the time-dependent relaxation of the total dipole moment of the system. We also varied the parameters $N_R$ and the 2 time-periods $\tau_1$ and $\tau_2$ and tested their influence on the dielectric properties of the 2 water models. As key quantities, we chose the radial distribution function $g(r)$, the dielectric permittivities (static) $\epsilon(0)$ (frequency dependent) $\epsilon(\omega)$ and the diffusion coefficients as a quantitative measure of the properties of water in dependency of the simulation parameters.

As stated in the main text, we consider the effect of the multiple renormalized biases and the renormalization parameters on the dynamical relaxation behavior of a time-dependent quantity $X$ describing a system, such as the relaxation of the time-dependent total dipole moment $M(t)$ of a water system. Any quantity $X(t)$ in an unbiased simulation follows a time-correlation function $A(X)(t)$, which can be described by an expansion to $M$ monoexponential decay processes with periods $\tau_m$, which is defined by the time-behavior of a quantity $X$, rate constants $k_m$ and prefactors $A_m$ :
\[ A(X)(t) = \sum_{m} A_{m0} \exp(-k_{mt}) . \] (33)

In contrast to the equilibrium MD case, a bias coupled to a set of \( N_R \) biases to \( N \) atoms, with \( \tau_1 \) and \( \tau_2 \) leading to an actual acceleration in terms of a change in the time-correlation function. That results in a modified relaxation behavior, affecting all dynamical quantities (That can lead to accelerated folding times, modified diffusion constants and re-orientation kinetics of H-bonds or dipoles in the system. There is also an effect on quantities such as the static and dynamic dielectric properties), which we write as an heuristic equation (as described in our simulation results of the dielectric response of SPC/E and TIP3P water):

\[ A_m(X)(t) = \sum_{m} \left( A'_{m0} \sum_{i} A'_{kim} \right) \exp \left( -\left( k_{m} + \sum_{i} k_{im} \right) t_{rel} \right) , \] (34)

where \( N_R \) stands for the number of renormalized biases and \( k_{im} \) is the rate-constant within each bias with index \( l \), and we note that the time \( t \) changes to a relative time \( t_{rel} \), which scales linear with an acceleration factor \( \rho > 1: t_{rel} = \rho t \), as long as the coupling to the underlying un-biased gradient remains sufficiently low (\( \beta_{ik} \leq 1 \times 10^{-4}, \eta' = \eta_{MD} \approx 25 - \) see Section : Methods, E: ‘Renormalization to the underlying Hamiltonian’) in the enhanced sampling simulation. The relation 34 shows that depending on the magnitude and the parameters (\( \beta, N_R \) and the coupling times \( \tau_1, \tau_2 \)), the Hamiltonian of the system and the time-correlation behavior in the simulation are modified, while the processes depending from the parameters \( A \) and rate constants \( k \) still are described by modified monoexponential time-dependent decays, since the renormalization and the conditions on adaptive bias MD obey the principle of action as described in the equation 5 in the main text. In other words, dynamical quantities such as dielectric quantities related to dipole fluctuations and in general fluctuation-dependent properties related to a linear response of the system can effectively be varied through the choice of the bias-parameters.

2.1. Water simulations : System preparation and analysis

We used SPC/E and TIP3P water simulations in order to test the influence of the new simulation method on the correlation behavior of the system and to validate the property of the algorithm to change dynamic properties while structural properties of water remain approximately unaffected.

For the sampling of dielectric properties of SPC/E and TIP3P water, we filled a box with dimensions \( 1 \times 1 \times 1 \) \( nm^3 \) with 33 water molecules (SPC/E or TIP3P). We applied PME electrostatics with a cut-off of 0.4 nm, while van der Waals interactions have been calculated using a shift function with the same cutoff. We checked the size-dependency of our results in simulations of a system with size of \( 5 \times 5 \times 5 \) \( nm^3 \) filled with 4124 water molecules (PME electrostatics with a real-space cut-off of 0.8 nm, and a cut-off for the Lennard-Jones 12-6 interactions of 0.6 nm). We used the same system and applied identical parameters to test replica exchange MD (REMD) using 4 replicas at 300 K with an exchange interval every 1,000 time-steps. For the REMD simulations, we omitted the determination of frequency dependent dielectric properties. All simulations have been carried out in the \( NVT \)-ensemble using the Nosé-Hoover thermostat (i.e. the Nosé-Hoover equations of motion) with a coupling interval of \( \tau_c = 1 \) ps and a reference temperature of 300 K. The small water-systems have been propagated over 20 ns, while the larger reference systems have been sampled over approximately 5 ns.

For the biased water-simulations, we used a \( \beta \) coupling parameter of \( 5 \times 10^{-4} \), 20 replicas and \( \tau_1=5 \) ps, \( \tau_2=10 \) ps. The variational parameters have been chosen with a width that no variation of \( \beta \)
and $\tau_1$, $\tau_2$ has been performed. The gradient optimization have been switched off. We systematically varied $\eta_{MD}$ and $\eta'$ from 0 to 500.0. For the calculation of frequency dependent properties of the dielectric constant $\epsilon(\omega)$, we calculated the autocorrelation function using the $g_{\text{dipoles}}$ module. We normalized the faster decay of the dipole-autocorrelation function from the biased simulations to the decay in the MD-simulations of SPC/E and TIP3P water using the relative positions of the first zero-value of the autocorrelation functions. In other words, we determined the acceleration factor $\eta'$ of the biased-simulation relative to the un-biased simulation in terms of the decay of the dipole autocorrelation function from 1 to 0 (see Figure 1S), which means that the effective acceleration in the reorientation of the water dipoles introduced by the bias is considered as general acceleration $\eta'$ in the sampling of the underlying system. That procedure leads to the correct frequency dependent dielectric permittivity in comparison with the experiment (see section : Results and Discussion, Water simulations, Dielectric properties).

The static dielectric constant $\epsilon(0)$ have been calculated using the Kirkwood factor $g_k$ as implemented in the gromacs-4.5.5 analysis module $g_{\text{dipoles}}$, using [14] :

$$g_k = \frac{1}{N\mu^2} \left( \frac{\langle M^2 \rangle - \langle M \rangle^2}{\langle M^2 \rangle} \right), \quad (35)$$

and

$$\epsilon(0) = 1 + \frac{4}{3} \pi Ng_k \mu^2/(k_BT). \quad (36)$$

The frequency dependent dielectric constant $\epsilon(\omega)$ has been calculated using the $g_{\text{dielectric}}$ module from GROMACS [15]. That formalism uses a normalized autocorrelation function $\Phi(t)$ :

$$\Phi(t) = \frac{\langle M(0)M(t) \rangle}{\langle M^2 \rangle}, \quad (37)$$

and uses the Fourier-Laplace transform[16] :

$$\frac{\epsilon(\omega) - 1}{2\epsilon_f + \epsilon(0)} = \int_0^\infty \left( - \frac{d\Phi(t)}{dt} \right) e^{-\omega t} dt. \quad (38)$$

For the case of the sampling along $N_R$ multiple biases, we refer to equation 34 and state that the static permittivity can be modified in dependency of the coupling parameters $\eta'$, $\beta$ and the number of multiple biases $N_R$. For the 2 processes as described in the equations 35 and 38, the correlation functions then effectively yield different permittivity values. As starting parameters for the fit to the decay of the autocorrelation function, we used a double exponential function with the fit parameters $A=0.5$, $\tau_1=10$ ps, $\tau_2=1$ ps, $\epsilon(0)$ resulting from the first analysis using the Kirkwood factor, $\epsilon_{RF}=78.5$ and a smoothing over 10 data points.

2.2. Water simulations : Results

2.2.1. Dielectric properties

We determined the static permittivities $\epsilon(0)$ ($\epsilon(\omega)$ at frequency $\omega = 0$) as function of simulation time in biased and unbiased simulations for SPC/E and TIP3P water. Through the investigation of the dependency from the $\eta_{MD}$ and the $\eta'$-parameters on the static permittivity, we determined the property of our algorithm to influence the correlation behavior of the underlying system. We used a static amount of bias-replicas and a constant coupling constant $\beta$ in order to measure the dependency from the fluctuation dependent parameters $\eta_{MD}$ and $\eta'$. Our results are summarized in Table 2.

In the biased simulations of SPC/E water, a parameter of $\eta_{MD}=500$, $\eta'=1.0$ yielded the lowest value of $\epsilon(0)=53$. We reach a maximum value in the static permittivity of $\epsilon(0)=119$ using parameters $\eta_{MD}=100.0$, $\eta'=100.0$, and obtain a permittivity value of $\epsilon(0)=78.4$ using the parameters $\eta_{MD}=30.0$,
Fig. S1. Normalized dipole autocorrelation functions from MD- and biased simulations of SPC/E and TIP3P water. We scaled the decay time of the autocorrelation result of biased SPC/E and TIP3P water to determine the linear acceleration factor $\eta'$ by which the reorientation of the dipole of water is accelerated (see section Methods in the main text). We find that $\eta'=10$ yields an optimal fit to the decay of the unbiased simulation and also leads to a good agreement with the experimental frequency dependent permittivity $\epsilon(\omega)$ (see main text).

We obtained $\eta'=30.0$ independent from the dimension of the system (1 nm$^3$ and 125 nm$^3$). In the MD-simulations of the systems with dimensions $1 \times 1 \times 1$ nm$^3$ and $5 \times 5 \times 5$ nm$^3$, we determine values for the permittivity of $\epsilon(0)=68.7$ and $\epsilon(0)=66.5$ (see Figure 2S c). We mention that the results contain an associated error of approximately $\Delta \epsilon(0) \approx 1-2$.

Using biased simulations of the TIP3P-water model in systems with the same dimensions, we obtain the lowest value for the static permittivity of $\epsilon(0)=55$ using $\eta_{MD}=500, \eta'=1.0$. In contrast, we measure a maximum value in the static permittivity of $\epsilon(0)=92$ with the parameters $\eta_{MD}=1.0, \eta'=1.0$. In the MD-simulations of the $1 \times 1 \times 1$ nm$^3$ and the $5 \times 5 \times 5$ nm$^3$ box we determine values for the permittivity of $\epsilon(0)=93$ and $\epsilon(0)=102$ (see Figure 2S d). We yielded approximately an identical value in a $5 \times 5 \times 5$ nm$^3$ box with the same parameters.

We analyzed the frequency dependent permittivities $\epsilon(\omega)$, and observed a decay in $\epsilon'(\omega)_{spec}$ from a value of 78.4 to 2.69 within the frequency range from 8E-3 to 1 THz in the simulation of SPC/E water using $\eta_{MD}=30.0, \eta'=30.0$. The same function decays 4 times slower in the simulation of TIP3P-water using $\eta_{MD}=110.0, \eta'=1.0$ in the frequency range from 8E-3 to 4 THz. In the unbiased MD-simulation of SPC/E water, we observe a decay of $\epsilon'(\omega)_{spec}$ from 72 to 3.2 in the frequency range from 2.5E-3 to 4 THz, while we measure a decay from 92.6 to 3.1 within the frequency range from 8.7E-3 to 4 THz in the unbiased MD-simulation of TIP3P-water. In the case of $\epsilon''(\omega)$ (i.e. the imaginary part of the frequency dependent permittivity), we observe a strong effect of the bias on the location of the maximum along the frequency axis. In the simulation of SPC/E water using $\eta_{MD}=30.0, \eta'=30.0$, we find a maximum value of $\epsilon''(\omega)_{spec}=39.7$ at 0.11 THz. For TIP3P water simulated with the parameters $\eta_{MD}=110.0, \eta'=1.0$, we find a maximum value of $\epsilon''(\omega)_{spec}=31.47$ at 0.107 THz. The experimental maximum is located at approximately $\epsilon''(\omega)_{spec}=37.24$ and $\epsilon=0.12$ THz [17]. In contrast, in the unbiased MD-simulations of SPC/E and TIP3P-water, we obtain maxima in $\epsilon''(\omega)_{spec}=39.69$ at $\omega=0.031$ THz (SPC/E) and $\epsilon''(\omega)_{spec}=44.5$ at $\omega=0.19$ THz (TIP3P). In the analysis of the REMD simulations, we omitted this analysis for the REMD trajectories of SPC/E and TIP3P water.
Fig. S2. Results from simulations of SPC/E and TIP3P water using MD and the path sampling technique. (a) Final configuration of a SPC/E water simulation after 20 ns of simulation. (b) Site-site radial distribution functions (RDF) (water oxygen - OW, water hydrogen - HW) from biased and unbiased MD simulations of SPC/E and TIP3P-water. (c, d) Static permittivities $\varepsilon(0)$ as function of MD-simulation time using different coupling $\eta_{MD}$ and $\eta'$-parameters in the individual simulations (c) - SPC/E water, (d) TIP3P water. (e) Frequency dependent permittivities $\varepsilon(\omega)$ as function of frequency $\omega$ obtained from biased and un-biased simulations of SPC/E and TIP3P water (SPC/E $\eta_{MD} = \eta' = 30.0$, TIP3P $\eta_{MD}=120$, $\eta=1.0$). (f) Mean-square displacement as function of simulation time from MD- and biased simulations of SPC/E and TIP3P water. Experimental values for the dielectric spectrum ($\varepsilon''(\omega)$) taken from ref. [17]. For the REMD simulations, we used 4 replicas at an identical target temperature (see section Methods) in order to sample the properties of water at 300 K.

2.2.2. Structural and dynamic properties

We analyzed the site-site specific radial distribution functions (RDF) of SPC/E and TIP3P-water obtained from biased and unbiased simulations of the system with a dimension of 125 nm$^3$. We measured the RDFs between water-oxygen, water-oxygen (OW-OW), water-oxygen, water-hydrogen (OW-HW) and water-hydrogen, water-hydrogen (HW-HW). Our results on the structural and dynamic properties are summarized in Table 2.

In unbiased MD-simulations of SPC/E water the RDFs contain maxima ((water oxygen-oxygen)(OW-OW) 0.276 nm (1), 0.45 nm (2) and 0.69 nm (3)), ((water oxygen-hydrogen)(OW-HW) 0.176 nm (1) and 0.326 nm (2)), while biased simulations of the same system using $\eta_{MD}$=40.0 and $\eta'=40.0$ contain maxima ((water oxygen-oxygen)(OW-OW) 0.276 nm (1), 0.45 nm (2) and 0.676 nm (3)), ((water oxygen-hydrogen)(OW-HW) 0.178 nm (1) and 0.314 nm (2)). In unbiased MD-simulations of TIP3P water we find maxima ((water oxygen-oxygen)(OW-OW) 0.278 nm (1), 0.45 nm (2) and 0.698 nm (3)), ((water oxygen-hydrogen)(OW-HW) 0.184 nm (1) and 0.322 nm (2)), while biased simulations of the same system using $\eta_{MD}$=120.0 and $\eta'=1.0$ contain maxima ((water oxygen-oxygen)(OW-OW) 0.272 nm (1), 0.54 nm (2) and 0.802 nm (3)), ((water oxygen-hydrogen)(OW-HW) 0.178 nm (1) and 0.314 nm (2)) (see Figure 2S b). Using the same $\eta$-coupling parameters, the REMD simulations of both SPC/E and TIP3P lead to approximately the same results as in the sequential simulations.

The relative probability densities as the different positions of the maxima are shifted in the biased simulations relative to the unbiased MD-simulations. Between the un-biased and the biased simulation of SPC/E water, we observe shifts (OW-OW) : $\Delta = 0.11, 0.45$ (1.11738)-0.45 nm (1.04048) (un-biased, biased) Δ ≈ 0.07, 0.69 (1.04572)-0.676
Table 1. Results from biased and unbiased MD-simulations of SPC/E and TIP3P water in a comparison with experimental results [17–21]. For the biased simulations, we used 20 bias replicas, a coupling strength $\beta = 5 \times 10^{-4}$, and $\eta_{MD}=30.0$ (SPC/E), 110.0 (TIP3P), $\eta'=30.0$ (SPC/E), 1.0 (TIP3P) which influences the fluctuation strength and quantities connected to the correlation behavior as the static and frequency dependent permittivity ($\epsilon(0)$, $\epsilon(\omega)$). For the REMD simulations, we used 4 replicas at the same target temperature of 300 K (see section Methods), while we used the same coupling parameters and number of bias-replicas. From left to right: Maxima of the site-site specific radial distribution functions averaged over the individual trajectories (water-oxygen, water-oxygen (OW-OW)) (position d(nm), g(r) value in brackets), (water-oxygen, water-hydrogen (OW-HW)).

|                | RDF(OWOW) 1st. d(nm) (g(r)) | RDF(OWOW) 2nd. d(nm) | RDF(OWOW) 3rd d(nm) | RDF (OWHW) 1st. d(nm) |
|----------------|-----------------------------|----------------------|---------------------|-----------------------|
| SPC/E, MD      | 0.276 nm (3.08483)          | 0.45 nm (1.11738)    | 0.69 nm (1.04572)   | 0.176 nm (1.601)      |
| SPC/E, MD $\eta'$=30.0 | 0.274 nm (3.19019)          | 0.45 nm (1.04048)    | 0.676 nm (1.02558)  | 0.176 nm (1.65284)    |
| SPC/E, $\eta_{MD}=30.0$, $\eta'=30.0$, REMD | 0.276 (3.05329)            | 0.45 nm (1.09015)    | 0.69 (1.03992)       | 0.178 nm (1.57627)    |
| TIP3P, MD      | 0.278 nm (2.68578)          | 0.45 nm (1.00191)    | 0.698 nm (1.02366)  | 0.184 nm (1.25235)    |
| TIP3P, $\epsilon_{TIP3P}=120.0$, $\omega=1.0$, REMD | 0.272 nm (2.76855)         | 0.54 nm (1.06742)    | 0.802 nm (1.00734)   | 0.178 nm (1.12709)    |
| TIP3P, $\epsilon_{TIP3P}=120.0$, $\omega=1.0$, REMD | 0.28 nm (2.25373)          | 0.56 nm (1.04699)    | 0.806 nm (1.00143)   | 0.186 nm (1.02549)    |
| Experiment[18] | 0.288 nm (3.09)             | 0.45 nm (1.14)       | 0.673 nm (1.07)     | 0.185 nm (1.38)       |

nm (1.02558) (un-biased, biased) $\Delta \approx 0.07$, (OW-HW) : 0.176 (1.601)-0.176 nm (1.65284) (un-biased, biased) $\Delta \approx 0.05$, and 0.326 (1.58315)-0.326 nm (1.57126) (un-biased, biased) $\Delta \approx 0.01$. In the biased and un-biased simulations of TIP3P water, we measure shifts (OW-OW) : 0.272 nm (2.76855) (un-biased, biased) $\Delta \approx 0.08$, 0.45 (1.00191)-0.54 nm (1.06742) (un-biased, biased) $\Delta \approx 0.06$, 0.698 (1.02368)-0.802 nm (1.00734) (un-biased, biased) $\Delta \approx 0.02$, (OW-HW) : 0.184 (1.25325)-0.178 nm (1.12709) (un-biased, biased) $\Delta \approx 0.1$, and 0.322 (1.44738)-0.314 nm (1.35551) (un-biased, biased) $\Delta \approx 0.09$. In terms of the structural properties, the REMD simulations of both water systems (TIP3P and SPC/E) reach an approximately identical behavior as in the sequential simulations.

In the MD-simulation of SPC/E water, we measure a self-diffusion coefficient $D$ of 1.0866 (± 0.2011) $10^{-5}$ cm$^2$/s. The self diffusion in the enhanced sampling simulation of SPC/E water using $\eta_{MD}=30.0$ and $\eta'=30.0$, equals 2.4436 (± 0.1665) $10^{-5}$ cm$^2$/s. In the MD-simulation of TIP3P water, we measure a self-diffusion coefficient $D$ with a value of 2.3045 (± 0.1547) $10^{-5}$ cm$^2$/s, while the enhanced simulation of the same system using $\eta_{MD}=120.0$ and $\eta'=1.0$ results in a value for $D$ of 17.0078 (± 5.2154) $10^{-5}$ cm$^2$/s (see Figure 2S f). The self-diffusion coefficient observed in the REMD simulations exceeds the values from the sequential runs, where we observe a value of 10.3925 $10^{-5}$ cm$^2$/s for SPC/E water and a value of 50.5028 $10^{-5}$ cm$^2$/s for TIP3P water.

2.2.3. Discussion

SPC/E water and the TIP3P model have been investigated in a number of MD simulation studies [15,17,22–26]. For the self diffusion coefficients of SPC/E water, a coefficient $D$ of 3.02 $10^{-5}$ cm$^2$/s has been determined in simulations, while simulations of TIP3P yielded 6.23 $10^{-5}$ cm$^2$/s [25]. An experimental value of 2.30 $10^{-5}$ cm$^2$/s has been reported for the self-diffusion coefficient of water [19,20]. The self-diffusion coefficient obtained in the biased simulation of SPC/E water is in very good agreement with that experimental value. Maxima in the site-site specific RDFs ((OW-OW) 0.288 nm (1), 0.45 nm (2) and 0.673 nm (3)), ((OW-HW) 0.185 nm (1) and 0.330 nm (2)) have been determined in neutron diffraction experiments [18], while site-site specific RDFs ((OW-OW) 0.278 nm (1), 0.45 nm (2) and 0.685 nm (3) (SPC/E)) ((OW-OW) 0.277 nm (1), 0.45 nm (2) and 0.684 nm (3) (TIP3P)), ((OW-HW) 0.180 nm (1) and 0.327 nm (2) (SPC/E)) ((OW-HW) 0.183 nm (1) and 0.322 nm (2) (TIP3P)) have been measured in simulations [25]. Our results from the biased and un-biased simulations of
SPC/E-water are in good agreement with their results. Investigations on the dielectric properties of water are of specific interest and especially the SPC/E water model has been found to describe frequency dependent dielectric permittivities shifted by -0.1 THz to the experimental maximum of \( \varepsilon''(\omega) \) [17], while a value of 70.7 [23] for the static permittivity value \( \varepsilon(0) \) has been reported. That value is shifted in comparison to the experiment by a value of approximately 8, while the experimental value is at 78.4 at room temperature [21]. In contrast to the SPC/E water model, static permittivities in the range from 93 to 104 have been observed for the TIP3P water model, while its peak for \( \varepsilon'' \) in the dielectric spectrum is shifted by approximately +0.1 THz in relation to the experimental value [17,24]. In terms of the static and frequency dependent permittivity value, our result from the biased simulation of SPC/E water is in good agreement with the experiment. That also holds for the REMD simulation in the case of the static permittivity, while we omitted the frequency dependent analysis due to the dependency of the dynamics of the system from the exchange frequency. The dielectric behavior of water plays an important role in the behavior of biomolecules, while static dielectric constants for biomolecular systems ranging from 10 to 20 have been reported [27,28]. Especially the non-polar region of proteins, which is shielded from the solvent in its folded state, can contain values for \( \varepsilon(0) \) of approximately 5 [29]. However, here it remains questionable to speak of a dielectric constant, which is a macroscopic quantity and might not be reducable to a range below the nm length-scale. Recent studies investigated the effect of ions and the folding of alanine-pentapeptide on the dielectric relaxation behavior of the underlying systems [30,31].

We conclude that the presented algorithm can change the time-dependent properties of SPC/E and TIP3P water through a scaling of \( \eta_{MD} \) and \( \eta' \), as we stated in the main body of the text in the Section: Methods. Especially the parameter \( \eta_{MD} \) can modify the time-dependent decay of the total dipole moment in comparison with the conventional MD propagation. We found that the magnitude \( \eta' \) affects the fluctuations of the applied bias averaged and also can change with the number of parallel replicas, as we find in the REMD simulations. For the TIP3P model, the charge density has a high value that a large value for \( \eta_{MD}=120.0 \) has to be used to scale down the static permittivity to a value of 78. That scaling has a significant effect on the water structure in comparison with the experimental result and shifts the location of maxima in the RDF. In the case of the SPC/E water, a scaling of \( \eta' \) to larger values yields the correct static permittivity value, an approximately correct water-structure in comparison with the experiment as well as a comparatively good agreement in its dielectric relaxation behavior.
behavior with experimental results. As we find in the REMD case, we observe that the combination of REMD with the multiple path-sampling can sample the structural properties of water in terms of the radial distribution function and the static permittivities in the same way as the sequential technique.
Fig. S3. (a, c, e, g) Difference $d_{ij}(t) - d_{ij}(exp)$ of the distance between NMR-distance restraint indexed atoms $ij$ and the experimental value as function of time and the restraint index. (b, d, f, g) Probability of finding the system within the difference of the distance between NMR-distance restraint indexed atoms $ij$ and the experimental value $d_{ij}(t) - d_{ij}(exp)$. (a, b) Results from un-restrained path-sampling simulation in implicit solvent. (c, d) Results from restrained path-sampling simulation in implicit solvent using NMR-restraints. (e, f) Results from un-restrained path-sampling simulation in explicit solvent. (g, h) Results from restrained path-sampling simulation in explicit solvent using NMR-restraints.
Fig. S4. Cluster populations from the cluster analysis of the simulations in implicit solvent (unrestrained).
Fig. S5. Cluster populations from the cluster analysis of the simulations in implicit solvent (NMR restraints).
**Fig. S6.** Cluster populations from the cluster analysis of the simulations in explicit solvent (unrestrained).
Fig. S7. Cluster populations from the cluster analysis of the simulations in explicit solvent (NMR restraints).
Fig. S8. (a, c) Average and minimal distance values along 11 distance restraints. (b, d) Population densities of each of the 11 distance restraints as function of the difference between the distance and the chemical restraint distance value as given in ref. [32]. (a, b) Results from restrained simulations in implicit solvent. (c, d) Results from restrained simulations in explicit solvent.
3. Restraint parameters

- Experimental NMR-NOESY restraint data used for the restraint simulations of the TrpCage miniprotein from ref. [33] (restraint-index, residue number 1, atom-name, residue number 2, atom-name, distance (Å)):

  1  2LEU  H  1ASN  HA  2.500  
  2  1ASN  HB2  2LEU  H  2.900  
  3  3TYR  H  2LEU  H  2.500  
  4  4ILE  H  2LEU  H  3.300  
  5  5GLN  H  2LEU  HA  2.500  
  6  5GLN  HB2  2LEU  HA  2.900  
  7  2LEU  HA  3TYR  H  2.900  
  8  2LEU  HB2  3TYR  H  2.900  
  9  4ILE  H  3TYR  H  2.000  
 10  3TYR  HB2  3TYR  HA  2.000  
 11  6TRP  H  3TYR  HA  2.900  
 12  6TRP  HB2  3TYR  HA  2.900  
 13  6TRP  HE3  3TYR  HA  2.900  
 14  19PRO  HB2  3TYR  HA  3.300  
 15  19PRO  HD2  3TYR  HA  2.900  
 16  19PRO  HG2  3TYR  HA  2.900  
 17  19PRO  HG2  3TYR  HB2  2.900  
 18  3TYR  HA  4ILE  H  2.900  
 19  4ILE  HB  4ILE  H  2.000  
 20  5GLN  H  4ILE  H  2.500  
 21  6TRP  H  4ILE  H  3.300  
 22  4ILE  HB  4ILE  HA  2.900  
 23  7LEU  H  4ILE  HA  2.500  
 24  4ILE  HA  5GLN  H  2.900  
 25  4ILE  HB  5GLN  H  2.500  
 26  5GLN  HB2  5GLN  H  2.000  
 27  7LEU  H  5GLN  H  3.300  
 28  8LYS  H  5GLN  HA  2.500  
 29  8LYS  HB2  5GLN  HA  2.900  
 30  6TRP  H  5GLN  HB2  2.900  
 31  5GLN  HA  6TRP  H  2.900  
 32  5GLN  HA  6TRP  H  2.900  
 33  6TRP  HB2  6TRP  H  2.000  
 34  6TRP  HE3  6TRP  H  3.300  
 35  7LEU  H  6TRP  H  2.500  
 36  6TRP  HD1  6TRP  HA  2.500  
 37  9ASP  H  6TRP  HA  3.300  
 38  9ASP  HB2  6TRP  HA  3.300  
 39  6TRP  HA  6TRP  HB2  2.000  
 40  6TRP  HD1  6TRP  HB2  2.500  
 41  16ARG  HB2  6TRP  HD1  2.900  
 42  16ARG  HD2  6TRP  HD1  2.900  
 43  16ARG  HG2  6TRP  HD1  2.900  
 44  18PRO  HA  6TRP  HD1  3.300  
 45  16ARG  H  6TRP  HE1  3.300
|   | 16ARG   | 6TRP   | HE1   | 2.900 |
|---|---------|--------|-------|-------|
| 47| 17PRO   | 6TRP   | HE1   | 2.900 |
| 48| 18PRO   | 6TRP   | HE1   | 2.900 |
| 49| 12PRO   | 6TRP   | HZ2   | 2.000 |
| 50| 17PRO   | 6TRP   | HZ2   | 3.300 |
| 51| 18PRO   | HD2    | 6TRP  | 2.900 |
| 52| 12PRO   | HG2    | 6TRP  | 2.900 |
| 53| 7LEU    | HG     | 6TRP  | 3.300 |
| 54| 7LEU    | HA     | 6TRP  | 2.900 |
| 55| 7LEU    | H      | 6TRP  | 2.500 |
| 56| 7LEU    | HG     | 6TRP  | 2.000 |
| 57| 6TRP    | 7LEU   | H     | 2.900 |
| 58| 7LEU    | HB2    | 7LEU  | 2.000 |
| 59| 7LEU    | HG     | 7LEU  | 2.500 |
| 60| 8LYS    | H      | 7LEU  | 2.500 |
| 61| 9ASP    | H      | 7LEU  | 3.300 |
| 62| 7LEU    | HB2    | 7LEU  | 2.900 |
| 63| 9ASP    | H      | 7LEU  | 3.300 |
| 64| 10GLY   | H      | 7LEU  | 3.300 |
| 65| 11GLY   | H      | 7LEU  | 2.000 |
| 66| 11GLY   | HA2    | 7LEU  | 2.900 |
| 67| 7LEU    | HA     | 8LYS  | 2.900 |
| 68| 7LEU    | HB2    | 8LYS  | 2.500 |
| 69| 8LYS    | HB2    | 8LYS  | 2.000 |
| 70| 9ASP    | H      | 8LYS  | 2.000 |
| 71| 10GLY   | H      | 8LYS  | 3.300 |
| 72| 8LYS    | HA     | 9ASP  | 2.900 |
| 73| 8LYS    | HB2    | 9ASP  | 2.900 |
| 74| 9ASP    | HB2    | 9ASP  | 2.000 |
| 75| 10GLY   | H      | 9ASP  | 2.000 |
| 76| 11GLY   | H      | 9ASP  | 3.300 |
| 77| 9ASP    | HA     | 10GLY | 2.900 |
| 78| 11GLY   | H      | 10GLY | 2.000 |
| 79| 10GLY   | HA2    | 11GLY | 2.900 |
| 80| 11GLY   | HA2    | 11GLY | 2.000 |
| 81| 12PRO   | HB2    | 12PRO | 2.500 |
| 82| 13SER   | H      | 12PRO | 2.900 |
| 83| 13SER   | H      | 12PRO | HB2  | 3.300 |
| 84| 6TRP    | HH2    | 12PRO | HA   | 3.300 |
| 85| 13SER   | H      | 12PRO | HA   | 2.900 |
| 86| 14SER   | H      | 12PRO | HA   | 3.300 |
| 87| 15GLY   | H      | 12PRO | HA   | 3.300 |
| 88| 13SER   | HB2    | 13SER | H    | 2.500 |
| 89| 14SER   | H      | 13SER | H    | 2.000 |
| 90| 13SER   | HA     | 14SER | H    | 2.900 |
| 91| 15GLY   | H      | 14SER | H    | 2.000 |
| 92| 14SER   | H      | 14SER | HB2  | 2.000 |
| 93| 15GLY   | H      | 14SER | HB2  | 3.300 |
| 94| 13SER   | HA     | 15GLY | H    | 2.900 |
| 95| 14SER   | HA     | 15GLY | H    | 2.900 |
Experimental chemical crosslinking restraint data used for the restraint simulations of the Killer Cell Lectin-like Receptor Subfamily B Member 1A from ref. [32] (restraint-index, residue number 1, atom-name, residue number 2, atom-name, distance (Å)):

1 166LYS N 179LYS N 20.000
2 179LYS N 196LYS N 20.000
3 148LYS N 196LYS N 20.000
4 146LYS N 196LYS N 20.000
5 125LYS N 212LYS N 20.000
6 146LYS N 148LYS N 12.000
7 147GLU N 148LYS N 12.000
8 121ASP N 125LYS N 12.000
9 123ASP N 125LYS N 12.000
10 143ASP N 146LYS N 12.000
11 176ASP N 196LYS N 12.000
12 176ASP N 179LYS N 12.000

Restraint data set used for the restraint simulations of the Killer Cell Lectin-like Receptor Subfamily B Member 1A from ref. [32] (restraint-index, residue number 1, atom-name, residue number 2, atom-name, distance (Å)). We note that we considered each distance-value to be significantly lower than the chemical cross-linking distances as described before (7.5 Å). The distance value only affects the sign of the overlapping segment of the path-increment dL in contrast to harmonic potentials:

1 212LYS N 121ASP N 7.500
2 212LYS N 121ASP H 7.500
3 212LYS N 121ASP CA 7.500
4 212LYS N 121ASP HA 7.500
5 212LYS N 121ASP CB 7.500
6 212LYS N 121ASP HB1 7.500
7 212LYS N 121ASP HB2 7.500
8 212LYS N 121ASP CG 7.500
9 212LYS N 121ASP OD1 7.500
10 212LYS N 121ASP OD2 7.500
11 212LYS N 121ASP C 7.500
12 212LYS N 121ASP O 7.500
|   |   |   |   |   |
|---|---|---|---|---|
| 13 | 125LYS | N | 123ASP | N | 7.500 |
| 14 | 125LYS | N | 123ASP | H | 7.500 |
| 15 | 125LYS | N | 123ASP | CA | 7.500 |
| 16 | 125LYS | N | 123ASP | HA | 7.500 |
| 17 | 125LYS | N | 123ASP | CB | 7.500 |
| 18 | 125LYS | N | 123ASP | HB1 | 7.500 |
| 19 | 125LYS | N | 123ASP | HB2 | 7.500 |
| 20 | 125LYS | N | 123ASP | CG | 7.500 |
| 21 | 125LYS | N | 123ASP | OD1 | 7.500 |
| 22 | 125LYS | N | 123ASP | OD2 | 7.500 |
| 23 | 125LYS | N | 123ASP | C | 7.500 |
| 24 | 125LYS | N | 123ASP | O | 7.500 |
| 25 | 212LYS | N | 125LYS | N | 7.500 |
| 26 | 212LYS | N | 125LYS | H | 7.500 |
| 27 | 212LYS | N | 125LYS | CA | 7.500 |
| 28 | 212LYS | N | 125LYS | HA | 7.500 |
| 29 | 212LYS | N | 125LYS | CB | 7.500 |
| 30 | 212LYS | N | 125LYS | HB1 | 7.500 |
| 31 | 212LYS | N | 125LYS | HB2 | 7.500 |
| 32 | 212LYS | N | 125LYS | CG | 7.500 |
| 33 | 212LYS | N | 125LYS | HG1 | 7.500 |
| 34 | 212LYS | N | 125LYS | HG2 | 7.500 |
| 35 | 212LYS | N | 125LYS | CD | 7.500 |
| 36 | 212LYS | N | 125LYS | HD1 | 7.500 |
| 37 | 212LYS | N | 125LYS | HD2 | 7.500 |
| 38 | 212LYS | N | 125LYS | CE | 7.500 |
| 39 | 212LYS | N | 125LYS | HE1 | 7.500 |
| 40 | 212LYS | N | 125LYS | HE2 | 7.500 |
| 41 | 212LYS | N | 125LYS | NZ | 7.500 |
| 42 | 212LYS | N | 125LYS | HZ1 | 7.500 |
| 43 | 212LYS | N | 125LYS | HZ2 | 7.500 |
| 44 | 212LYS | N | 125LYS | HZ3 | 7.500 |
| 45 | 212LYS | N | 125LYS | C | 7.500 |
| 46 | 212LYS | N | 125LYS | O | 7.500 |
| 47 | 146LYS | N | 143ASP | N | 7.500 |
| 48 | 146LYS | N | 143ASP | H | 7.500 |
| 49 | 146LYS | N | 143ASP | CA | 7.500 |
| 50 | 146LYS | N | 143ASP | HA | 7.500 |
| 51 | 146LYS | N | 143ASP | CB | 7.500 |
| 52 | 146LYS | N | 143ASP | HB1 | 7.500 |
| 53 | 146LYS | N | 143ASP | HB2 | 7.500 |
| 54 | 146LYS | N | 143ASP | CG | 7.500 |
| 55 | 146LYS | N | 143ASP | OD1 | 7.500 |
| 56 | 146LYS | N | 143ASP | OD2 | 7.500 |
| 57 | 146LYS | N | 143ASP | C | 7.500 |
| 58 | 146LYS | N | 143ASP | O | 7.500 |
| 59 | 196LYS | N | 146LYS | N | 7.500 |
| 60 | 148LYS | N | 146LYS | N | 7.500 |
| 61 | 196LYS | N | 146LYS | H | 7.500 |
| 62 | 148LYS | N | 146LYS | H | 7.500 |
113 148LYS N 147GLU CD 7.500
114 148LYS N 147GLU OE1 7.500
115 148LYS N 147GLU OE2 7.500
116 148LYS N 147GLU C 7.500
117 148LYS N 147GLU O 7.500
118 196LYS N 148LYS N 7.500
119 196LYS N 148LYS H 7.500
120 196LYS N 148LYS CA 7.500
121 196LYS N 148LYS HA 7.500
122 196LYS N 148LYS CB 7.500
123 196LYS N 148LYS HB1 7.500
124 196LYS N 148LYS HB2 7.500
125 196LYS N 148LYS CG 7.500
126 196LYS N 148LYS HG1 7.500
127 196LYS N 148LYS HG2 7.500
128 196LYS N 148LYS CD 7.500
129 196LYS N 148LYS HD1 7.500
130 196LYS N 148LYS HD2 7.500
131 196LYS N 148LYS CE 7.500
132 196LYS N 148LYS HE1 7.500
133 196LYS N 148LYS HE2 7.500
134 196LYS N 148LYS NZ 7.500
135 196LYS N 148LYS HZ1 7.500
136 196LYS N 148LYS HZ2 7.500
137 196LYS N 148LYS HZ3 7.500
138 196LYS N 148LYS C 7.500
139 196LYS N 148LYS O 7.500
140 179LYS N 166LYS N 7.500
141 179LYS N 166LYS H 7.500
142 179LYS N 166LYS CA 7.500
143 179LYS N 166LYS HA 7.500
144 179LYS N 166LYS CB 7.500
145 179LYS N 166LYS HB1 7.500
146 179LYS N 166LYS HB2 7.500
147 179LYS N 166LYS CG 7.500
148 179LYS N 166LYS HG1 7.500
149 179LYS N 166LYS HG2 7.500
150 179LYS N 166LYS CD 7.500
151 179LYS N 166LYS HD1 7.500
152 179LYS N 166LYS HD2 7.500
153 179LYS N 166LYS CE 7.500
154 179LYS N 166LYS HE1 7.500
155 179LYS N 166LYS HE2 7.500
156 179LYS N 166LYS NZ 7.500
157 179LYS N 166LYS HZ1 7.500
158 179LYS N 166LYS HZ2 7.500
159 179LYS N 166LYS HZ3 7.500
160 179LYS N 166LYS C 7.500
161 179LYS N 166LYS O 7.500
162 196LYS N 176ASP N 7.500
163 179LYS  N  176ASP  N  7.500
164 196LYS  N  176ASP  H  7.500
165 179LYS  N  176ASP  H  7.500
166 196LYS  N  176ASP  CA  7.500
167 179LYS  N  176ASP  CA  7.500
168 196LYS  N  176ASP  HA  7.500
169 179LYS  N  176ASP  HA  7.500
170 196LYS  N  176ASP  CB  7.500
171 179LYS  N  176ASP  CB  7.500
172 196LYS  N  176ASP  HB1  7.500
173 179LYS  N  176ASP  HB1  7.500
174 196LYS  N  176ASP  HB2  7.500
175 179LYS  N  176ASP  HB2  7.500
176 196LYS  N  176ASP  CG  7.500
177 179LYS  N  176ASP  CG  7.500
178 196LYS  N  176ASP  OD1  7.500
179 179LYS  N  176ASP  OD1  7.500
180 196LYS  N  176ASP  OD2  7.500
181 179LYS  N  176ASP  OD2  7.500
182 196LYS  N  176ASP  C  7.500
183 179LYS  N  176ASP  C  7.500
184 196LYS  N  176ASP  O  7.500
185 179LYS  N  176ASP  O  7.500
186 196LYS  N  179LYS  N  7.500
187 196LYS  N  179LYS  H  7.500
188 196LYS  N  179LYS  CA  7.500
189 196LYS  N  179LYS  HA  7.500
190 196LYS  N  179LYS  CB  7.500
191 196LYS  N  179LYS  HB1  7.500
192 196LYS  N  179LYS  HB2  7.500
193 196LYS  N  179LYS  CG  7.500
194 196LYS  N  179LYS  HG1  7.500
195 196LYS  N  179LYS  HG2  7.500
196 196LYS  N  179LYS  CD  7.500
197 196LYS  N  179LYS  HD1  7.500
198 196LYS  N  179LYS  HD2  7.500
199 196LYS  N  179LYS  CE  7.500
200 196LYS  N  179LYS  HE1  7.500
201 196LYS  N  179LYS  HE2  7.500
202 196LYS  N  179LYS  NZ  7.500
203 196LYS  N  179LYS  HZ1  7.500
204 196LYS  N  179LYS  HZ2  7.500
205 196LYS  N  179LYS  HZ3  7.500
206 196LYS  N  179LYS  C  7.500
207 196LYS  N  179LYS  O  7.500

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