Sensitivity of peptide conformational dynamics on clustering of a classical molecular dynamics trajectory

Christian H. Jensen, Dmitry Nerukh, and Robert C. Glen
Unilever Centre for Molecular Science Informatics, Department of Chemistry, University of Cambridge, CB2 1EW Cambridge, United Kingdom

(Received 10 July 2007; accepted 8 January 2008; published online 21 March 2008)

We investigate the sensitivity of a Markov model with states and transition probabilities obtained from clustering a molecular dynamics trajectory. We have examined a 500 ns molecular dynamics trajectory of the peptide valine-proline-alanine-leucine in explicit water. The sensitivity is quantified by varying the boundaries of the clusters and investigating the resulting variation in transition probabilities and the average transition time between states. In this way, we represent the effect of clustering using different clustering algorithms. It is found that in terms of the investigated quantities, the peptide dynamics described by the Markov model is sensitive to the clustering; in particular, the average transition times are found to vary up to 46%. Moreover, inclusion of nonphysical sparsely populated clusters can lead to serious errors of up to 814%. In the investigation, the time step used in the transition matrix is determined by the minimum time scale on which the system behaves approximately Markovian. This time step is found to be about 100 ps. It is concluded that the description of peptide dynamics with transition matrices should be performed with care, and that using standard clustering algorithms to obtain states and transition probabilities may not always produce reliable results. © 2008 American Institute of Physics [DOI: 10.1063/1.2838980]

I. INTRODUCTION

There are many methods which seek to simulate the folding of a peptide or protein. They range from very coarse-grained approaches like the HP model to models with atomic detail like molecular dynamics. While the coarse-grained method gives results which can be useful as guidelines when designing proteins, they do not describe exactly how a specific protein folds. To do this, a model with the detail of molecular dynamics is needed. However, for the system sizes of interest, the computational task of performing a molecular dynamics simulation which shows protein folding is unfeasible. Therefore, there have been developments of algorithms which modify standard molecular dynamics to allow for simulations of these larger systems. These methods range from modifying the potential energy landscape of the protein, to simulating several replicas of the same system at different temperatures, to constructing Markov models from a large number of molecular dynamics simulations.

A method which combines several molecular dynamics simulations by using clustering and a Markov model for the state transitions has recently been proposed. Using this method, it is possible to reconstruct the overall dynamics of a peptide from thousands of individual simulations. This is done by counting the number of transitions between the different states from all the simulations. The Markov model can be described by a state vector \( \nu \) which holds probabilities for the different configurations and a transition matrix \( T \). Given that the system has state vector \( \nu_t \) at time \( t \), the state vector at time \( t+\Delta t \) can be calculated as \( \nu_{t+\Delta t} = \Delta t \).

A source of error in this approach could be the clustering of configurational states. In the present paper, we investigate how the state transition probabilities and folding dynamics vary with slightly different clustering. The total number of clusters is kept constant and only the boundaries between clusters are varied. This is done to try and mimic the effect of different clustering algorithms. The investigation is carried out on a small peptide, ensuring that possible transitions are sufficiently sampled. To the best of our knowledge, there is no systematic analysis of the sensitivity of the clustering to the resulting dynamical characteristics. However, we have found that the results are sensitive to the clustering. First, if the clustering is done in dihedral space, that could lead to the appearance of nonphysical sparsely populated states, resulting in a variation in average transition times of up to 814%. This shows a likely effect of clustering incorrectly. Second, if the clusters are defined correctly, the sensitivity of the average transition times to the variation in the boundaries is up to 46%. This shows the likely variation in results obtained with a clustering algorithm that performs well.

Another source of error in the transition matrix approach is whether the transitions between the states can be described accurately with a Markov model. It is found that at short time scales, the transitions do not have a Markovian behavior; however, at longer time scales, they become Markovian. This is in line with previous work reported in the
The problem is addressed by choosing a sufficiently long time scale in the construction of the Markov model.

II. METHODS

In this investigation, we analyze a molecular dynamics trajectory. The simulation was performed using the software package GROMACS 3.2. The system examined was the four residue peptide valine-proline-alanine-leucine (VPAL) solvated in 874 water molecules. The peptide is shown in Fig. 1. The simulation box was 3.0 × 3.0 × 3.0 Å³. The force field was 53αβ. This is optimized for bimolecular systems interacting with water. Periodic boundary conditions were used. The temperature was kept at 300 K using the thermostat of Berendsen et al. Atomic positions were recorded every 0.5 ps. The integration algorithm was a Verlet type and the integration step was 0.002 ps. The system was equilibrated before it was sampled for 500 ns. This produced a total of 10^8 data points.

In our investigation, we need to be able to vary the clusters. Therefore, the clustering is done by choosing dihedral angles as cutoff angles between the different regions. We only use the two central pairs of dihedrals because the terminal residues are too flexible and do not define the overall structure of the molecule. The initial clustering is represented by the solid lines in Fig. 2. The dotted lines represent the interval in which the cutoff angles are varied. By varying each angle, in turn, it is possible to investigate the transition matrix as a function of different cutoff angles. Each angle is varied ±0.5 rad around the initial cutoff. By plotting the variation in the transition matrix elements with the dihedral angle cutoff positions, it is possible to inspect how sensitive the transition matrix is to clustering. By the method given in Sec. II A, it is also possible to calculate how the variation in clustering affects the average transition time. The latter is a clear physical measure which characterizes the folding routes directly. It can also be used to describe the folding pathways when there are multiple initial and final states.

To apply the Markov model transition matrix approach, we need to find the time scale at which the systems behavior is Markovian. The Markovian assumption is that \( \nu_{t+\Delta t} = T_{\Delta t} \nu_t \), where \( T_{\Delta t} \) is the transition matrix constructed for a time step of \( \Delta t \). For a transition matrix constructed at a time step of \( n \Delta t \), we must have \( T_{n\Delta t} = T_{\Delta t}^n \), where \( n = 1, 2, 3, \ldots \). By expanding each transition matrix in eigenvalues and eigenvectors, it can be shown that a necessary condition for the Markovian assumption to be valid is that \( \lambda_{n\Delta t,i} = \lambda_i^n \), where \( \lambda_i \) denotes an eigenvalue and \( i \) runs over the number of eigenvalues. From this we find that \( \lambda_{n\Delta t,i} \) has to be constant for \( n = 1, 2, 3, \ldots \). This constant is the eigenvalue of a transition matrix with a time step of \( \Delta t \), which does satisfy the Markovian assumption. Given an eigenvalue, it is possible to calculate a decay time (e.g., the half-life) for the corresponding eigenvector. Using the constant eigenvalue, we, therefore, get that the time \( \tau_i = -[n\Delta t/\ln(\lambda_{n\Delta t,i})] \) has to be constant if the Markovian description is correct. To find the time scale at which the system’s behavior is Markovian, we can, therefore, construct transition matrices for the time steps \( n \Delta t, n = 1, 2, 3, \ldots \), and calculate \( \tau_i \) for each matrix. The time step at which \( \tau_i \) for all \( i \) becomes constants is the time step at which the system’s behavior is Markovian.

A. Calculating the average transition time

To calculate the average transition time of a Markov model, we need to define initial and final states. Each of these can either be one state or a set of states. Assuming that we have a set of initial states \( I \) and a set of final states \( F \), the average transition time can be written as

\[
t_{IF} = \sum_{n=1}^{\infty} n P_{IF}(n).
\]

Here \( P_{IF}(n) \) is the probability for all paths of length \( n \) which start in \( I \) and end on \( F \). We assume that the Markov process is described by a transition matrix \( T \) and that there is a total of \( N \) states. The first problem in the calculation is to find an expression for \( P_{IF}(n) \). By introducing \( \tilde{T}, L, o, \) and \( v \), we construct the following algorithm:

- Form the transition matrix \( T \), remove the rows and columns for all states in \( F \) to form a new matrix \( \tilde{T} \). This new matrix will have a dimension of \((N-d)\times(N-d)\)

- Form the matrix \( L \), which is of dimension \( d \times (N-d) \) and holds the matrix elements of \( T \) that give the probabilities for entering \( F \) from all other states.

- Form the row vector \( o \), which is of dimension \( 1 \times d \) and holds 1’s in all places.
Form the vector $v$ of dimension $(N-d) \times 1$. The elements of $v$ must describe the initial distribution of states in $I$. If each starting state is equally likely, then their elements must be equal. For the states not in $I$, the initial value in $v$ must be zero. The total sum of all elements in $v$ must be 1.

Using the quantities given above, $P_{IF}(n)$ can be written as $oL\tilde{T}_n^{-1}v$ (an explanation is given in the Appendix). Let us assume that $\tilde{T}$ has eigenvectors $e_i$ with corresponding eigenvalues $\lambda_i$. We then expand $v$ in this basis. This gives $v = \sum \alpha_i e_i$. The average transition time [Eq. (1)] can then be written as

$$t_{IF} = \sum_{n=1}^{\infty} n P_{IF}(n) = \sum_{n=1}^{\infty} noL\tilde{T}_n^{-1}v$$

$$= \sum_{n=1}^{\infty} noL\tilde{T}_n^{-1} \sum_{i} \alpha_i e_i = \sum_{n=1}^{\infty} \sum_{i} noLa_i \lambda_i^{-1} e_i \quad (2)$$

$$= \sum_{i} \left( \sum_{n=1}^{\infty} n \lambda_i^{-1} \right) \alpha_i oLe_i = \sum_{i} \frac{\alpha_i}{(1-\lambda_i)^2} oLe_i.$$

III. RESULTS

In our investigation, we partition the configurational space of the peptide in six different locations (Fig. 2). In the

FIG. 3. The average RMSD for the molecular configurations from different clusters compared to a representative conformation from cluster number 1. The error bars indicate the standard deviation.

FIG. 4. (Color online) The average conformations of the VPAL molecule in the different states. Comparing to the clusters in Fig. 2, the states correspond to (1) $A_1A_2$, (2) $B_1A_2$, (3) $A_1C_2 + B_1C_2$, (4) $A_1B_2$, and (5) $B_1B_2$. 

Author complimentary copy. Redistribution subject to AIP license or copyright, see http://jcp.aip.org/jcp/copyright.jsp
plot for proline, we see that the two cutoff lines means that there are two states. In the alanine plot, there are four cutoff lines, which give three different states. This gives a total of six different states for the peptide. However, because one of the states found in this way is very sparsely populated, we remove this state to form a total of five states. The average conformations in these states can be seen in Fig. 4. To investigate if this clustering is correct, we have compared it to clustering using root mean square deviation (RMSD). This is done by taking a representative configuration for each cluster and calculating the RMSD of all the configurations in each cluster. For cluster number 1, the result is shown in Fig. 3. It can be seen that the RMSD is smallest for configurations which are also in cluster number 1, and that this cluster is well separated from the other clusters. Similar results are obtained when using the other clusters. Therefore, clustering using cutoff angles in dihedral space is comparable to clustering using RMSD.

Using the states shown in Fig. 4 allows the calculation of a transition matrix. This is done by simply counting the number of transitions between the states in the molecular dynamics trajectory. This gives a frequency matrix which holds the number of transitions. By normalizing the columns in this matrix to unity, the transition matrix is obtained. To determine an appropriate time step to take when building the transition matrix, we need to find the time step at which the system behaves in a Markovian manner. To to this, we follow the procedure given in Sec. II. Transition matrices are constructed with varying time steps. For each matrix, the $\tau_i$'s are calculated for all $i$. The result of this can be seen in Fig. 5. When the system’s behavior is Markovian, the $\tau_i$’s should be constant. From about 50 ps, it can be seen that the values become approximately constant; however, we chose a time step of 100 ps to make sure that our system’s behavior is sufficiently Markovian.

In Eq. (3), the transition matrix for the initial clustering with a 100 ps time step is given. It can be seen that once in a state, there is a high probability of staying there in the next time step. From the transition probabilities, it is possible to trace out the transition paths of the highest probabilities. These paths will be the conformational routes that the pep-

FIG. 5. (Color online) The variations in the $\tau_i$’s (see text). Each curve corresponds to an eigenvalue. The curve for the eigenvalue 1 is not shown as this gives an infinite $\tau$ value.

FIG. 6. The range of transition probabilities for the different matrix elements as the clustering is varied. $k$ is the matrix element index defined as $k=5(i-1)+j$, where $i$ is the row number and $j$ the column number. The range of the variation has been magnified five times for clarity.
tide will most likely follow during transitions. It is what is commonly known as the folding path. In Fig. 6, the variation in transition probability between all pairs of states can be seen for the six different variations in cutoff angles. For some elements, these variations are substantial. However, the variation of a single transition probability does not describe what happens to the peptide as a whole. Therefore, to describe the sensitivity of the folding path of a peptide, it is desirable to have a measure which describes how variations in the probabilities affect the folding path. This is exactly what is achieved by calculating the average transition time between states:

\[
T_{100 \text{ ps}} = \begin{bmatrix}
0.8972 & 0.1006 & 0.0345 & 0.3502 & 0.0650 \\
0.0407 & 0.7215 & 0.0055 & 0.0324 & 0.2129 \\
0.0240 & 0.0136 & 0.9496 & 0.1289 & 0.0519 \\
0.0295 & 0.0182 & 0.0091 & 0.3491 & 0.1475 \\
0.0087 & 0.1461 & 0.0013 & 0.1394 & 0.5228
\end{bmatrix}
\]

(3)

In Fig. 7, the variation in average transition time between all pairs of states can be seen for the six different variations in cutoff angles. It is clear that the variation is more significant compared to the variation of the transition matrix elements. This is because the variation in average transition time describes the variations in the folding path as a whole and not just a single transition. Since a deviation in cutoff angle from the initial cutoff angle will typically mean that clusters are connected by more transitions, the average folding time, between states, will generally tend to decrease. This causes a typical bell shaped variation in the average transition time as a function of the variation in cutoff angle. For the VPAL peptide, we assume the unfolded state to be state 1 and the folded state, where the terminal residues of the peptide form a salt bridge, to be state 5. The average transition times between these two states are shown in Fig. 8.

In Fig. 7, the average transition time between states is also shown for a transition matrix constructed with a time step of 0.5 ps (red in the figure). As can be seen from Fig. 5, this is not a correct description of the system since it does not have a Markovian behavior at this time scale. However, it is still interesting to note that on this time scale, the average transition times seem to be more sensitive to the clustering than at the longer time scale.

The transition probabilities for transitions directly between these two states are almost zero. This means that the variation in average transition time is caused by the variations of the transition probabilities between the intermediate states. The variation in average transition time between these two states is about 46%, which is significant. In the case where the sparsely populated state was included as a state on its own, the variations in average transition time to and from
this state was up to 814%. Examples of these large variations are shown in Fig. 9. It can be seen that the variation mostly affected the average folding time between a few states. This is because the main path for transitions between other states does not include the scarcely populated state. For the VPAL peptide, it can also be seen that \( t_{51} \) is generally larger than \( t_{15} \), which means that the folded state is more stable than the unfolded state.

For a larger peptide, the variation can be expected to be smaller, because there are many more paths by which the peptide can fold. However, assuming a given peptide has a specific initial and final states can reach 46%. When the transition matrix is constructed with a time step of 0.5 ps, the variation in average transition time between representative initial and final states can reach 46%. When the transition matrix is constructed with a time step of 0.5 ps (i.e., a non-Markovian time step), this variation increases to 100%.

For a case where the initial clustering was miscalculated by inclusion of the nonphysical sparsely populated states, we found the variations in average transition times between some of the states to be as much as 814%. The choice of clustering is a difficult one. On the one hand, if one chooses to use only clusters which are highly populated, the transition probabilities and average transition times will not be as sensitive. However, this may also mean that important information about the folding path is lost.

IV. CONCLUSIONS

When constructing Markov models from molecular dynamics simulations, care must be taken. First, it is important that the Markov model is constructed with a sufficiently large time step so that the dynamics of the system are as close to Markovian as possible. In our investigation, we found that the transitions’ behavior is sufficiently Markovian at 100 ps time step. However, for the purpose of construction of reliable models, we also found that this is not enough to ensure an accurate description of the dynamics. In particular, we have found that transition probabilities and, hence, average transition times are sensitive to the specific clustering. By varying the boundaries between clusters, we found that the variation in average transition time between representative initial and final states can reach 46%. When the transition matrix is constructed with a time step of 0.5 ps (i.e., a non-Markovian time step), this variation increases to 100%.

For a case where the initial clustering was miscalculated by inclusion of the nonphysical sparsely populated states, we found the variations in average transition times between some of the states to be as much as 814%. The choice of clustering is a difficult one. On the one hand, if one chooses to use only clusters which are highly populated, the transition probabilities and average transition times will not be as sensitive. However, this may also mean that important information about the folding path is lost.

ACKNOWLEDGMENTS

The work is supported by Unilever and the European Commission (EC Contract No. 012835-EMBIO).

APPENDIX: CALCULATION OF \( P_{ir}(n) \)

To illustrate how \( P_{ir}(n) \) is calculated, let us consider a three-state system. Let the initial state be 1 and the final state be 3. The transition matrix for the system is given as

\[
T = \begin{bmatrix}
a_{11} & a_{12} & a_{13} \\
a_{21} & a_{22} & a_{23} \\
a_{31} & a_{32} & a_{33}
\end{bmatrix}
\]

First, we form the matrices \( \bar{T} \), \( L \), \( o \), and \( v \):

\[
\bar{T} = \begin{bmatrix}
a_{11} & a_{12} \\
a_{21} & a_{22}
\end{bmatrix},
L = [a_{31} \ a_{32}],
\ o = [1],
\ v = \begin{bmatrix} 1 \\ 0 \end{bmatrix}.
\]

For \( n=1 \), we get

\[
P_{31}(1) = oL\bar{T}v = a_{31}.
\]

Since \( P_{31}(1) \) is the probability to go from state 1 to state 3 in one step, there is only one possible path 1-3. The probability for this is simply \( a_{31} \). For \( n=2 \), we get

\[
P_{31}(2) = oL\bar{T}^2v = a_{31}a_{11} + a_{32}a_{21}.
\]

There are two possible paths 1-1-3 and 1-2-3. The probability for each of these is \( a_{31}a_{11} \) and \( a_{32}a_{21} \), respectively. The sum of these, therefore, gives the total probability. For \( n=3 \), we get

\[
P_{31}(3) = oL\bar{T}^3v = a_{31}a_{11}a_{11} + a_{31}a_{11}a_{21} + a_{32}a_{21}a_{11} + a_{32}a_{22}a_{21}.
\]

In this case, there are four possible paths from state 1 to state 3. These are 1-1-1-3, 1-2-1-3, 1-1-2-3, and 1-2-2-3. \( P_{31}(3) \) is the sum of the probabilities for each of these paths.

115107-7 Peptide conformational dynamics

