Software News and Updates

Wordom: A User-Friendly Program for the Analysis of Molecular Structures, Trajectories, and Free Energy Surfaces

MICHELE SEEBER, ANGELO FELLINE, FRANCESCO RAIMONDI, STEFANIE MUFF, RAN FRIEDMAN, FRANCESCO RAO, AMEDEO CAFLISCH, FRANCESCA FANELLI
1 Dipartimento di Chimica, University of Modena and Reggio Emilia v. Campi 183, 41125 Modena, Italy
2 Dulbecco Telethon Institute (DTI), University of Modena and Reggio Emilia, 41125 Modena, Italy
3 Department of Biochemistry, University of Zurich, Winterthurstrasse 190, CH-8057 Zurich, Switzerland
4 Freiburg Institute for Advanced Studies (FRIAS), University of Freiburg, Albertstr. 19, 79104 Freiburg, Germany

Received 11 June 2010; Revised 20 August 2010; Accepted 5 September 2010
DOI 10.1002/jcc.21688
Published online 29 November 2010 in Wiley Online Library (wileyonlinelibrary.com).

Abstract: Wordom is a versatile, user-friendly, and efficient program for manipulation and analysis of molecular structures and dynamics. The following new analysis modules have been added since the publication of the original Wordom paper in 2007: assignment of secondary structure, calculation of solvent accessible surfaces, elastic network model, motion cross correlations, protein structure network, shortest intra-molecular and inter-molecular communication paths, kinetic grouping analysis, and calculation of mincut-based free energy profiles. In addition, an interface with the Python scripting language has been built and the overall performance and user accessibility enhanced. The source code of Wordom (in the C programming language) as well as documentation for usage and further development are available as an open source package under the GNU General Purpose License from http://wordom.sf.net.

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Key words: structural/dynamics analysis program; free energy landscape; elastic network model; protein structure network; communication paths

Introduction

Wordom is a program aimed at fast manipulation and analysis of individual molecular structures and molecular conformation ensembles. Its development started in 2003 and the relative publication appeared in 2007.1

A number of programs are already available to analyze molecular structures and dynamics. These include: (a) the most common molecular simulation and analysis packages, like CHARMM, Gromacs, and Amber; (b) a number of molecular viewers, like VMD and Pymol; (c) command-line oriented analysis programs and script suites, like MMTSB, carma, and pcazip; and (d) packages that provide environments for structural analysis, like Bio3D, MTK, or Biskit. In this panorama, Wordom was originally conceived as a simple command-line utility to quickly access data in common structural data files. Basic manipulation tools were then implemented, which paved the way for the adoption of a modular framework to easily add analysis routines. At the time of the first publication, novel analysis modules already formed the bulk of Wordom’s code, and others have been added since then.

Some of the new modules (Table 1), such as secondary structure assignment (SSA), surface area calculations, and elastic network models (ENM), implement tools that are already available in some form in other software packages or web servers. However, their use on trajectory files is either cumbersome or unpractical. Indeed, programs for SSA and surface computation are widespread, but most of them can only deal with a single structure file at a time, thus making the handling of multiconformation files complex and time-consuming.
Table 1. New Features in Wordom Since the Original Publication.$^1$

| Module                          | Label$^3$ | Function                                                | Reference |
|---------------------------------|-----------|---------------------------------------------------------|-----------|
| Secondary Structure Assignment  | SSA       | Assignment of secondary structure based on geometric criteria | 15, 16    |
| Molecular Surface               | SURF      | Calculation of solvent accessible, solvent excluding and van der Waals surfaces; surface correlation along a trajectory | 17, 18    |
| Elastic Network Model           | ENM       | Calculation of elastic network models on a protein structure | 19–24     |
| Cross Correlation               | CORR      | Correlations of atomic displacements along a trajectory | 25–27     |
| Protein Structure Network       | PSN       | Calculation of network of amino acid interactions | 28–31     |
| PSN Path                        | PSN-path  | Path calculation within protein structure network | 32, 33    |
| Clustering                      | CLUS      | Clustering according to conformation similarity | 34–36     |
| cut-based Free Energy Profile   | cFEP      | Computation of a one-dimensional free energy profile that preserves barriers between free energy basins | 39, 46    |
| Kinetic Grouping Analysis       | KGA       | Determination of free energy basins based on kinetic behavior | 40        |

$^3$Abbreviation/acronym used in the text.

New Tools in Wordom

**Secondary Structure Assignment**

The SSA module is able to evaluate the secondary structure of a peptide or protein using two methods, DLIKE or DCLIKE, derived from the DSSP$^{15, 16}$ and DSSPcont$^{15, 16}$ algorithms, respectively. These two approaches are considered two standards in the field of secondary structure assignments. DSSPcont is a consensus-based DSSP assignment, in which the whole DSSP procedure is run 10 times with different values of the energy cutoff that defines an hydrogen bond (H-bond)$^{15, 16}$. Assignments are then weighted according to the cutoff and a consensus is given as the final output. DSSP and DSSPcont assignments are generally comparable.

Both algorithms have been rewritten from scratch since the DSSP license does not allow free reuse of the code. The output is a simple string where the $n^{th}$ character corresponds to the secondary structure of the $n^{th}$ amino acid. There are eight possible letters in the secondary structure “alphabet”: H, G, I, E, B, T, S, and L, standing for α helix, 3_10 helix, π helix, extended, isolated β-bridge, hydrogen bonded turn, bend, and unstructured loop, respectively.$^{15, 16}$ No extra information such as that included in the typical DSSP output is given, since the SSA module is meant to be used for a quick analysis of the secondary structure profile along a trajectory, rather than for a complete and throughout characterization of a single structure.
Comparisons between the secondary structure assignments by Wordom and by the DSSP program are shown in Table 2. The agreement is good, i.e., 92%, considering also that most discrepancies do not concern exchanges between helices and strands. The higher speed of the SSA module compared to DSSP shows itself on trajectory files (see Table 3). In fact, whereas the SSA module can compute the secondary structure along a trajectory very fast, DSSP works on single frame files previously extracted from the trajectory. Thus, the better performance of Wordom must be ascribed, at least in part, to the lack of input/output operations associated with handling each molecule conformation as a standalone file (see Table 3). The speedup is more pronounced when dealing with small systems, e.g., peptides.

Contrarily to DSSP, because Wordom is conceived to operate on the results of simulations, the structure files must contain all the atoms that contribute to the backbone H-bonds. Therefore, structures derived directly from the protein data bank (PDB), especially the X-ray structures that miss hydrogen atoms or entire residues, must be completed before submission to the SSA module.

### Molecular Surface Calculation, Correlation, and Clusterization

Wordom computes different kinds of molecular surfaces using two different algorithms: an exact analytical method developed by Hu and coworkers (i.e., ARVO algorithm)\(^{17}\) and a fast numerical method developed by Pascual-Almir and coworkers (i.e., GEPO\(_{\text{L}}\) algorithm).\(^{18}\) ARVO calculates the solvent accessible surface area by expressing the molecular surface as a sum of double integrals around an energy minimum. Within this technique, the protein structure is described by a reduced subset of atoms (usually C\(_{\text{α}}\) atoms)\(^{20}\) whose coordinates can be derived either from structure determinations (crystallography, NMR) or from molecular simulations. The interactions between particle pairs are given by a single term Hookean harmonic potential.\(^{19}\) The total energy of the system is given by

\[
E = \sum_{i\neq j} k_{ij} (d_{ij} - d_{ij}^0)^2 
\]

where \(d_{ij}\) and \(d_{ij}^0\) are the instantaneous and equilibrium distances between C\(_{\text{α}}\)-atoms \(i\) and \(j\), respectively, whereas \(k_{ij}\) is a force constant, whose definition varies depending on the type of ENM used. The second derivatives of the harmonic potential are stored in a \(3N \times 3N\) Hessian matrix (H), whose diagonalization gives a set of \(3N\)-6 nonzero-frequency eigenvectors and associated eigenvalues.

Two alternative versions of ENM have been implemented. In the first version, termed "linear cutoff-enn," the force constant is equal to 1 for pairwise interactions between C\(_{\text{α}}\)-atoms \(i\) and \(j\), respectively, whereas \(k_{ij}\) is a force constant, which is constant (with a default value of 40 Kcal/mol \(\cdot\) \(\text{Å}^2\)).\(^{21}\) The total energy of the system is thus described by the simple Hamiltonian:

\[
E = \sum_{i\neq j} k_{ij} (d_{ij}^0 - d_{ij}^0)^2
\]

where \(d_{ij}\) and \(d_{ij}^0\) are the instantaneous and equilibrium distances between C\(_{\text{α}}\)-atoms \(i\) and \(j\), respectively, whereas \(k_{ij}\) is a force constant, whose definition varies depending on the type of ENM used. The second derivatives of the harmonic potential are stored in a \(3N \times 3N\) Hessian matrix (H), whose diagonalization gives a set of \(3N\)-6 nonzero-frequency eigenvectors and associated eigenvalues.

Two alternative versions of ENM have been implemented. In the first version, termed "linear cutoff-enn," the force constant is equal to 1 for pairwise interactions between C\(_{\text{α}}\)-atoms lying within a cutoff distance chosen by the user, and adjacent C\(_{\text{α}}\)-atoms are assigned a force constant equal to 10.\(^{20}\) In the second one, termed "Kovacs-ENM,"\(^{21}\) the force constant depends on the distance of the interacting particles:

\[
k_{ij} = C \left( \frac{d_{ij}^0}{d_{ij}} \right)^6
\]

where \(C\) is constant (with a default value of 40 Kcal/mol \(\cdot\) \(\text{Å}^2\)).\(^{21}\)

### Table 2. Comparison Between the Secondary Structure Assignments Made by Wordom (SSA Module, DLIKE Option) and Those Made by the DSSP Program.\(^8\)

| E | B | T | S | L | H | G | I | Total |
|---|---|---|---|---|---|---|---|-------|
| 2103 | 31 | 18 | 21 | 85 | 2 | 0 | 0 | 2260 |
| 11 | 58 | 6 | 6 | 38 | 1 | 7 | 0 | 127 |
| 16 | 1 | 638 | 51 | 32 | 6 | 3 | 0 | 747 |
| 12 | 0 | 5 | 656 | 13 | 0 | 2 | 0 | 688 |
| 44 | 3 | 9 | 6 | 1351 | 3 | 0 | 0 | 1416 |
| 0 | 1 | 11 | 2 | 7 | 951 | 17 | 0 | 989 |
| 1 | 0 | 39 | 0 | 3 | 1 | 163 | 0 | 207 |
| 0 | 0 | 2 | 0 | 0 | 0 | 0 | 0 | 2 |
| **Total** | **2187** | **94** | **728** | **742** | **1529** | **964** | **192** | **6436** |

\(^8\)The test set consists of 29 proteins (2CCY, 1ECA, 2IFO, 1TPM, 1HRE, 1PHT, 2POR, 3BCL, 2HLA, 1CDQ, 1AFC, 1MSA, 1VMO, 1HXN, 1NSC, 2BBK, 3AAH, 1TSP, 2PEC, 1PPK, 1STD, 4TIM, 1BR, 1INT, 1PYA, 2DNJ, 1PLQ, 1BNH, and 1PYP) selected as representatives of common folds.\(^{30}\) Results have been pooled together for each program and compared. Each element \(ij\) of the matrix reports the number of residues assigned by Wordom and by DSSP to be in conformation \(i\) and \(j\), respectively.

### Table 3. Speed (in Seconds) Comparison of Secondary Structure Computations.

| #Residues | #Frames | DSSP\(_{\text{DCD}}^a\) | DSSP\(_{\text{PDB}}^b\) | Wordom\(_{\text{SSA}}^c\) |
|-----------|---------|----------------|----------------|----------------|
| 316       | 10,000  | 1460           | 920            | 640            |
| 16        | 10,000  | 238            | 155            | 0.35           |

\(^a\)A script extracted each single frame by mean of Wordom and called DSSP on the extracted frame.

\(^b\)A script called DSSP on the already-extracted frames.

\(^c\)Calculation through the Wordom SSA module.
The structural perturbation method (SPM) has been recently described as a technique useful to characterize allosteric wiring diagrams in the context of the ENM lowest frequency modes.\textsuperscript{22} According to this methodology, amino acid positions that are relevant to protein dynamics are searched by perturbing systematically all the springs that connect the Ca-atom to the other atoms and then measuring the residue-specific response of such perturbations in the context of a given mode $m$. The perturbation response is computed as:

$$\delta \omega_m = v_m^T \cdot \delta H \cdot v_m$$  \hfill (3)

where $v_m$ is the eigenvector of mode $m$, $v_m^T$ is its transpose, and $\delta H$ is the Hessian matrix of the perturbation to the energy of the elastic network:

$$\delta E = \frac{1}{2} \sum_{i \neq j} \delta k_{ij} (d_{ij} - d_{ij}^0)^2$$  \hfill (4)

The response $\delta \omega_m$ is proportional to the elastic energy of the springs that are connected to the $i$th residue when they are perturbed by an arbitrary value (0.1), thus defining the most critical nodes for the dynamics of a given mode. The number of modes used for the computation is specified by the user (from 1 up to 3N-6). It is also possible to generate, for each analyzed mode, a pdb file containing the values of $\delta \omega_m$ in the $\beta$-factor field (Fig. 1).

Theoretical $\beta$-factors can be computed inside the ENM module, by the formula\textsuperscript{23}

$$B_n^T = \frac{8\pi^2 kT}{3} \frac{N}{\sum_{i=1}^{N} \lambda_m} \sum_{m=1}^{N} v_m^2$$  \hfill (5)

where $\lambda_m$ is the $n$th element of eigenvector $m$, $\lambda_m$ is the associated eigenvalue, $k$ is the Boltzmann constant, and $T$ is the temperature in K.

Cross correlations between theoretical and experimental $\beta$-factors can be also computed according to the following equation:

$$CC = \frac{\sum_{i=1}^{N} \beta_i^0 \beta_i^T}{\sqrt{\sum_{i=1}^{N} \beta_i^0 \beta_i^0} \cdot \sqrt{\sum_{i=1}^{N} \beta_i^T \beta_i^T}}$$  \hfill (6)

where $\beta_i^0$ and $\beta_i^T$ are the theoretical and experimental $\beta$-factors, and $\bar{\beta}^0$ and $\bar{\beta}^T$ are the theoretical and experimental $\beta$-factor average over all atoms, respectively. The number of modes used for the computation is specified by the user (from 1 up to 3N-6).

Moreover, involvement coefficients $I$ between the ENM modes and the displacement vector between a given structure/frame $T$ and a reference structure $R$ can be computed according to the following equation:

$$I_m = \frac{\sum_{n=1}^{3N} v_{mn} \Delta r_i^T}{\sum_{n=1}^{3N} \sum_{n=1}^{3N} v_{mn} \Delta r_i^T}$$  \hfill (7)

where $\Delta r_i = r_i^T - r_i^R$ and $r_i^R$ is the $i$th coordinate in the two conformers and $v_{mn}$ is the $n$th element of eigenvector $m$.\textsuperscript{24} By default, the computation is done for all 3N-6 modes, and only the values of $I$ greater than an arbitrary threshold (i.e., 0.2) are output.
Cross Correlation

Wordom implements two different algorithms to calculate correlations of atomic displacements along an MD trajectory. One algorithm, called dynamic cross-correlation (DCC), is a simple and well-established method based on the calculation of the normalized covariance of atom/residue position vectors. DCC values are computed as:

\[ C_{ij} = \frac{(r_i - \bar{r}_i)(r_j - \bar{r}_j)}{\sqrt{(r_i^2 - \bar{r}_i^2)(r_j^2 - \bar{r}_j^2)}} \]  (10)

where \( i \) and \( j \) may be atoms or centroids of atoms grouped by residue, and \( r_i \) and \( r_j \) are the corresponding position vectors. DCC represents the extent of atom/residue displacement correlation within a range that goes from 1.0 to \(-1.0\); where 1.0 indicates completely correlated (same period and phase) and \(-1.0\) completely anti-correlated (same period and opposite phase) displacements.

The second algorithm, called linear mutual information (LMI), is computationally more expensive (see Table 4) than DCC but overcomes some limitations of the DCC algorithm and is able to estimate correlations between perpendicular motions. LMI values are computed as:

\[ I_{lin}(x_i, x_j) = \frac{1}{2} \left( \ln |C_i| + \ln |C_j| - \ln |C_{ij}| \right) \]  (11)

where \( i \) and \( j \) may be atoms or residues, \( C_{ij} \) is the pair-covariance matrix, and \( C_i \) and \( C_j \) are marginal covariance matrices.

For a selection of 360 atoms within 10000 trajectory frames, the DCC and LMI methods took, respectively, 4'' and 63'' on the same processor (Table 4). The relative correlation matrices are shown in Figure 2.

Protein Structure Network

In recent times, the concept of PSN has been explored, giving more insights into the global properties of protein structures. The representation of protein structures as networks of interactions between amino acids has proven to be useful in a number of studies, such as protein folding, residue contribution to the protein–protein binding free energy in given complexes, and prediction of functionally important residues in enzyme families. All these aspects pertain to the issue of intra-molecular and inter-molecular communication.

Wordom implementation of PSN analysis is based on the work and algorithms described in the relevant papers by the Vishveshwara and coworkers. PSN is constructed from the atomic coordinates of residues, which represent the nodes of the network. Two nodes are connected by an edge if the percentage of interaction between them is greater than or equal to a given Interaction Strength cutoff.

\[ I_{ij} = \frac{n_{ij}}{\sqrt{N_i N_j}} \times 100 \]  (12)

where \( n_{ij} \) is the number of interactions between nodes \( i \) and \( j \), and \( N_i \) and \( N_j \) are the degrees of nodes \( i \) and \( j \), respectively.
Figure 2. Cross-correlation matrix of the atomic fluctuations of the Gαi1-subunit Ca-atom and the geometrical center of GTP. The regions below and above the matrix main diagonal concern the DCC and LMI correlation methods, respectively. DCC correlation values go from \(-1.0\) (fully anti-correlated motions) to 1.0 (fully correlated motions), whereas LMI correlation values go from 0.0 (fully uncorrelated motions) to 1.0 (fully correlated motions).

where \(I_{ij}\) is the interaction percentage of nodes \(i\) and \(j\), \(n_{ij}\) is the number of side-chain atom pairs within a given distance cutoff (4.5 Å as a default), and \(N_i\) and \(N_j\) are, respectively, the normalization factors (NF) for residues \(i\) and \(j\), which take into account the differences in size of the different nodes and their propensity to make the maximum number of contacts with other nodes in protein structures. The NFs for the 20 amino acids in our implementation were taken from the work by Kannan and Vishveshwara. Novel NFs for nonamino acid nodes can be introduced as well by the user. In this respect, the current version of the module holds also the NFs for retinal, guanine nucleotide di- and tri-phosphates (GDP and GTP, respectively), and Mg\(^{2+}\). In detail, retinal NF (i.e., 170.13) was computed as the average number of contacts done by the molecule in a dataset of 83 crystallographic structures concerning the different photointermediate states of bacteriorhodopsin, bovine rhodopsin, sensory rhodopsin, and squid rhodopsin. The NFs for GDP and GTP (i.e., 220.19 and 274.78, respectively) were derived from datasets of 55 and 69 G proteins, respectively. Finally, the NFs for Mg\(^{2+}\) concern GTPases and is 14.65 and 22.01 in the GDP- and GTP-bound states (i.e., based upon 41 GTPase structures) and GTP-bound states (i.e., based upon 68 GTPase structures). \(I_{ij}\) are calculated for all node pairs excluding \(j = i \pm n\), where \(n\) is a given neighbor cutoff (2 as default), and each node pair with an \(I_{ij}\) value greater than or equal to a given \(I_{\text{min}}\) cutoff is connected by an edge. Different networks can be achieved by probing a range of \(I_{\text{min}}\) cutoffs. At high \(I_{\text{min}}\) cutoffs, only nodes with high number of interacting atom pairs will be connected by edges, indicative of stronger inter-residue interactions. At a given \(I_{\text{min}}\) cutoff, those nodes that realize more than a given number of edges (4 as default) are called hubs. The percentage of interaction of a hub node is

\[
I_i = \frac{n_{ij}}{N_i} \times 100
\]

where \(I_i\) is the hub interaction percentage of node \(i\), \(n_{ij}\) is the number of side-chain atom pairs within a given distance cutoff and \(N_i\) is the normalization factor of residue \(i\). Node inter-connectivity is used to highlight cluster-forming nodes, where a cluster is a set of connected amino acids in a graph. Node clusterization procedure is such that nodes are iteratively assigned to a cluster if they can establish a link with at least one node in such cluster. A node not linkable to existing clusters initiates a novel cluster and so on until the node list is exhausted. The size (defined as the number of nodes) of the largest cluster is used to calculate the \(I_{\text{critic}}\) value. \(I_{\text{critic}}\) is defined as the \(I_{\text{min}}\) at which the size of the largest cluster is half the size of the largest cluster at \(I_{\text{min}} = 0.0\). At \(I_{\text{min}} = I_{\text{critic}}\) weak node interactions are discarded, emphasizing the effects of stronger interactions on PSN properties.

The Wordom implementation of PSN analysis allows the user to: (a) modify all the involved cutoffs (i.e., distance, neighbor, hub); (b) make residue selections; (c) set \(I_{\text{min}}\) ranges; and (d) set,
when dealing with a trajectory, the fraction of frames for which a PSN property is defined as stable. Furthermore, Wordom computes all network properties described in the relevant papers by Vishveshwara’s group (i.e., interaction strength for all node pairs, stable node interactions, hub frequencies, cluster compositions, and dimensions).\textsuperscript{28, 29}

An original feature of Wordom is the hub correlation analysis, a simple but effective method to highlight correlations in the propensity of two nodes to behave as hubs along an MD trajectory (Fig. 3).

The results of an application of the PSN module to a 10,000 frame trajectory of the $G_{\alpha_1}$-subunit complexed with GTP are shown in Figures 4A and 4B. The relative CPU time required for such a demonstrative calculation is reported in Table 4.

**Search for Communication Paths**

As an extension of the PSN analysis tool, Wordom can calculate the shortest non-covalently connected path(s) between two residues of interest in a single structure or in a trajectory (Fig. 4C), by combining PSN node inter-connectivities and residue correlated motions, as described in the relevant paper by the Gosh and Vishveshwara.\textsuperscript{32} Path search through the PSN-path module uses Dijkstra’s algorithm\textsuperscript{33} to traverse PSN inter-connectivities, and to find the shortest paths in each frame. It consists in: (a) search for all shortest paths between selected residue pairs based upon the PSN connectivities and (b) selection of paths that contain at least one residue correlated with either one of the two extremities (i.e., the first and last amino acids in the path). Once the shortest paths have been found, their frequencies, i.e., the number of frames containing the selected path divided by the total number of frames in the trajectory, are computed, which helps selection of the most meaningful paths. Steps (a) and (b) of path search employ the outputs from the PSN and CORR modules, respectively. The Wordom implementation allows the user to tune several parameters of the path-search routine (i.e., minimum interaction strength cutoff between nodes, lowest accepted residue correlation cutoff, minimum length and frequency of paths). Either the DCC or LMI methods can be chosen as a source of residue correlations.
Clusterings

The original RMSD- and DRMS-based clustering module allowed the choice of three different algorithms: leader-like, \(^{34}\) hierarchical, \(^{35}\) and quality threshold-like (QT-like). \(^{36}\) QT-like differs from the original QT algorithm in the check performed to assess whether a conformation belongs to a cluster or not. The original QT builds a perspective cluster for each frame by comparing it with all others and adding conformations progressively farther away from the starting frame until each new addition is within the chosen threshold with respect to all previously added frames. The largest of all these perspective clusters is then taken as the first cluster, its members are taken out of the conformation population and the procedure is run again until all conformations are either in a cluster or isolated. In contrast, QT-like builds the clusters only checking that newly added conformations are within the threshold with respect to all previously added frames. The largest of all these clusters within a distance lower than the threshold were labelled as isolated. In contrast, leader-like clustering is less accurate than the QT or the Hierarchical algorithms since the frame belongs to the same cluster as the preceding frame(s) is quite high. The accuracy of the new option is only slightly lower than the “comparison with all clusters” approach, but the execution is faster than the original “stop at first viable cluster” option. Leader-like clustering is less accurate than the QT or the Hierarchical algorithms since it compares each frame only with the clusters that have been already found along the trajectory, thus making the outcome dependent on the frame order.

More relevant improvements concern the Hierarchical and QT-like algorithms. Indeed, they have been both modified so that the original memory requirements have been almost halved. Furthermore, the actual clustering algorithms massively use multithreading in the CPU-intensive computation of the inter-frame distances (RMSD or DRMS). Also, the distance matrix can now be saved for later use, so that, if clustering with different threshold values is desired, the distance-computing step needs to be performed only once. Finally, the original two-pass clustering has been improved as well. In detail, after a first clustering run on a subset of frames, a second pass can be run that assigns each considered frame to the nearest cluster found in the first run. In the original version, frames with no new cluster centers, so that new clusters can be found and populated in the second run. This improves the overall accuracy and allows for a smaller portion of the total data set to be used in the first run.

Figure 4. Results of PSN and PATH analyses on a 10 ns MD trajectory of GTP-bound Ger\(_{1\alpha}\)-subunit. (A) Ca\(_\alpha\)-atoms of the 27 stable hub residues, at \(I_{\text{min}} = 3.0\%\), are represented as cyan spheres. The GTP molecule, which is a stable hub as well, is shown as a red sphere centered on the C\(_4\) ribose atom. Nodes are considered as stable hubs if they are involved in at least four connectivities at a given \(I_{\text{min}}\) (3.0% in this case) in more than 50% of the trajectory frames. (B) The 90 nodes that contribute to the largest cluster at \(I_{\text{min}} = 3.0\%\) are shown as green spheres centered on the Ca\(_\alpha\)-atoms. The GTP molecule, which participates as well in such cluster, is shown as a red sphere centered on the C\(_4\) ribose atom. (C) Representation of the most frequent shortest communication path (i.e., frequency = 46%). The amino acids that participate in the path are shown as magenta spheres centered on the Ca\(_\alpha\)-atoms, whereas GTP, which participates in the path as well, is shown as a red sphere centered on the C\(_4\) ribose atom. The two apical residues in this path are A152 and I222, located, respectively, in the \(\alpha\)-helical and Ras-like domains.
Figure 5. Complex network analysis of free energy landscapes. (A) Conformation space network. Nodes and links are protein conformations (i.e., microstates, see main text) and direct transitions sampled during the MD simulation, respectively. Node size is proportional to the population of the microstate, whereas link width is proportional to the transitions frequencies, i.e., larger link widths indicate more frequent transitions. Densely connected regions of the network represent rapidly interconverting microstates that belong to the same free energy basin (highlighted by a shaded circle). (B) Simplified example of a two state system. The free energy barrier between the two macro-states is represented by a region of minimum flow in the network (identified by a minimum-cut). (C) Cut-based free energy profile (cFEP). The free energy is projected onto the partition function-based reaction coordinate \( Z \), a projection that preserves the barriers as it takes into account all possible pathways to a reference microstate. The solid vertical line indicates the correspondence between the minimum-cut and the highest free energy barrier.

When accuracy is paramount, the QT-like algorithm is probably the most appropriate, being more accurate than the leader-like one and significantly faster (with comparable accuracy) than the Hierarchical algorithm (Table 4). Yet, in spite of the improvements, it remains considerably memory-hungry. Therefore, when dealing with big data sets (>1M–10M frames, depending on the available computing power and memory), with which it is impossible to consider all frame–frame distances, the user can choose to either use QT on a subset of frames and then run a two-pass clustering, or to opt for the leader-like algorithm.

**Determination of Free Energy Basins and Barriers**

Wordom has two distinct modules, cFEP and KGA, devoted to the identification of (meta)stable states sampled by MD simulations. The key idea of cFEP and KGA is to group conformations not according to structural criteria, but mainly according to equilibrium kinetics. In this way, an analysis of the MD trajectory in terms of free energy basins, i.e., basins of attraction of the dynamics, is provided. The main advantage of cFEP with respect to KGA is the information on the height and location of the free-energy barriers along the cumulative partition function, which can be used to identify the transition state structure(s). On the other hand, the KGA procedure does not require the use of a reaction coordinate to determine the free-energy basins.

For both cFEP and KGA procedures, MD snapshots (i.e., Cartesian coordinate sets) need to be finely clusterized and assigned to a discrete set of microstates. Clusterization can be done according to atomistic, i.e., RMSD-based clustering, or to coarse-grained representations such as secondary structure strings. Both RMSD-based and coarse-grained clustering have proven to be good discretization methods of MD trajectory snapshots into a set of microstates that describe large conformational changes (see Ref. 40, 48, and 53–56 for examples in protein folding). Application to large proteins requires more sophisticated clustering procedures like principal component space.

**Min-cut-Based Free Energy Profile**

The cFEP module refers to a rigorous method introduced by Krivov and Karplus for determining a one-dimensional free energy profile that preserves the barriers between free energy basins; given the barriers, free energy basins can be determined. The method uses the relative partition function, which is a reaction coordinate that takes into account all possible pathways to a reference state (e.g., the folded state).

The cFEP algorithm is based on a network description of the conformational dynamics. Each microstate (see above) represents a node of the conformation space network and a link is made if...
Figure 6. Network description of MD and evaluation of kinetic distance. The high-dimensional free-energy surface is coarse-grained into nodes of a network. The figure shows a schematic illustration of the transition network of a $\beta$-sheet peptide where nodes represent microstates and links represent direct transitions sampled along the MD simulation(s). The size of the nodes and links is proportional to the statistical weight of the microstates and number of transitions, respectively. The cFEP method implemented in Wordom requires a reference microstate. In this simplified illustration, the reference microstate is the large red sphere in the center of the folded state (which is the $\beta$-sheet structure, i.e., the basin on the left). The kinetic distance of each node from the reference microstate can be evaluated in Wordom by the folding probability ($p_{\text{fold}}$) or the mean first passage time (mfpt). The kinetic distance is rendered by the continuous coloring from red (folded, i.e., $p_{\text{fold}} = 1$ or $\text{mfpt} = 0$) to blue (unfolded, i.e., $p_{\text{fold}} = 0$ or $\text{mfpt} = \infty$).

A direct transition between two microstates is observed during the timeseries in a time step of a given size (see Fig. 5).59

The cFEP module implemented in Wordom is a precise and fast approximation of the minimum-cut method.60, 61 The free energy is projected as a function of the partition function relative to a reference node.39, 46 With this method, microstates are ranked according to their kinetic proximity with respect to a reference microstate (Fig. 6). The relative partition function is used as the progress coordinate, and the free energy barriers are determined as a function of it, either based on the probability of reaching the folded state before unfolding ($p_{\text{fold}}$)46 or on the mean first passage time (mfpt)39 to a selected node (both calculated analytically from the transition matrix). The $p_{\text{fold}}$ implementation, which requires a target node, is appropriate to find barriers between two well-defined basins, which are specified by the user through the assignment of $p_{\text{fold}} = 1$ to the representative node of one basin, and $p_{\text{fold}} = 0$ to the representative node of the other. On the other hand, the mfpt-based method is more suitable for free energy profiles relative to a single target basin (e.g., for unfolding profiles), for which the representative node of the target basin is assigned $\text{mfpt}_{\text{target}} = 0$.

**Kinetic Grouping Analysis**

The free energy basins are determined by KGA on the basis of kinetic behaviors (fast relaxation at equilibrium) along an MD simulation.40 The KGA method is based on a network description of the conformational dynamics.

Two protein microstates are grouped in a basin if, along the MD trajectory, they interconvert frequently within a short commitment time $\tau_{\text{commit}}$, which represents a typical relaxation time within basins. The principle behind this approach is that if two conformations interconvert rapidly, they are not separated by a barrier and therefore belong to the same basin. The $\tau_{\text{commit}}$ is a characteristic of the investigated system. It is an important (user-chosen) parameter of KGA and defines the resolution with which basins are isolated. A short $\tau_{\text{commit}}$ will group structures only locally or if the free energy
surface is very smooth. A longer $\tau_{\text{commit}}$ is more generous and might group sub-basins, isolated by a shorter $\tau_{\text{commit}}$, into larger basins. The log–log plot of the distribution of first passage times to the native microstate (or a representative microstate of another basin) usually reflects two timescales: the inter- and intra-basin relaxation times (see Fig. 7 of Mufl and Caflisch40). The barrier that separates the two regimes can give a good indication for the relaxation time.

The KGA module allows for isolation of either all relevant basins at once or of a single basin. In the first case, for a fixed commitment time $\tau_{\text{commit}}$, a matrix with interconversion (commitment) probabilities $p_{\text{commit}}$ between any pair of microstates can be calculated in principle, and microstates with $p_{\text{commit}} \geq 0.5$ are grouped together. Because the computational cost of all-against-all calculations increases quadratically, in practice one selects a subset of highly populated microstates (e.g., the 500 most populated microstates), calculates the $p_{\text{commit}}$-matrix and divides them into basins. In a post-processing step, all other microstates are assigned commitment probabilities to the isolated basins; finally, all microstates having a $p_{\text{commit}} \geq 0.5$ to a given basin are assigned to it. Otherwise, the microstates remain unassigned. Both the heavy-microstate calculation and the post-processing are done by Wordom in the same function. On the other hand, if only one basin is of interest or if the relaxation times within basins lay on different timescales, it is better to choose an appropriate $\tau_{\text{commit}}$ for each basin separately and then calculate the commitment probability ($p_{\text{commit}}$) according to it. In this way, it is possible to isolate basins one-by-one. In this case, the user has to run the procedure a number of times equal to the number of basins that need to be isolated. In addition to the $\tau_{\text{commit}}$, a representative microstate of each basin (usually the most populated microstate in the basin) has to be specified. Finally, all microstates that commit to the representative microstate of a basin with probability $p_{\text{commit}} \geq 0.5$ are considered as part of that basin.

**Python Bindings**

Using the SWIG (simple wrappers and interface generator)62 tools, a python module has been written that gives access to most of Wordom’s input/output functions and structures in the python environment via a simple import command. Basic analysis functions (e.g., RMSD, distances, atoms selections) are also exposed to the python environment. The availability of Wordom’s input/output functions allows scripts to operate directly on molecular data, whereas access to Wordom’s analysis functions makes it easy to compute properties on molecules or whole trajectories, and to further process the output without writing full-fledged C code or resort to temporary files. It is also practical to write the prototype of an analysis module in python and then convert it to C to enhance its performance, as has been done for some of the recently added modules.

**Conclusions**

Wordom is a user-friendly program for manipulating and analysing data from structural studies and molecular simulations. The latest release represents a significant improvement and enrichment of the original version published in 2007,1 as it provides new analysis tools that are unique to Wordom. These include new procedures for efficient structural analysis such as dynamic PSN and shortest communication path modules, which are effective tools to infer amino acids essential for stability and function as well as to unravel intra-molecular and inter-molecular communication mechanisms. Other novelties are user-friendly methods for determining free energy basins and barriers using the network of transitions sampled by MD simulations. With these new tools, Wordom can be used to analyze the free energy surface and therefore investigate the thermodynamics and kinetics of complex molecular processes, e.g., the reversible folding of structured peptides (Fig. 6).

Improvements include also the implementation of an interface with the popular scripting language Python.

Like the original version, this version of Wordom is released under the general purpose license (GPL), which allows anybody to download, modify, and redistribute both source code and binary files. This license has been adopted in order to foster diffusion and encourage contributions to the code.

**Acknowledgments**

The authors thank M. Cecchini, G. Settanni, and A. Vitalis for helpful discussions, Y. Valentini for technical help, S. Krivov for suggesting the modification to the leader clustering algorithm, and P. Schütz for providing Figure 6.

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