Fractal Analysis of Protein Potential Energy Landscapes

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The fractal properties of the total potential energy $V$ as a function of time $t$ are studied for a number of systems, including realistic models of proteins (PPT, BPTI and myoglobin). The fractal dimension of $V(t)$, characterized by the exponent $\gamma$, is almost independent of temperature and decreases with time, more slowly the larger the protein. Perhaps the most striking observation of this study is the apparent universality of the fractal dimension, which depends only weakly on the type of molecular system. We explain this behavior by assuming that fractality is caused by a self-generated dynamical noise, a consequence of intermode coupling due to anharmonicity. Global topological features of the potential energy landscape are found to have little effect on the observed fractal behavior.

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I. INTRODUCTION

The energy landscape of proteins near the folding temperature is often described as complex. That is, the landscape is expected to be rugged and composed of many minima and maxima, separated by barriers of varying heights, the result of frustration due to conflicting interactions between different segments of the protein. The complex nature of the underlying energy landscape has been demonstrated using ergodic measures and by exact enumeration of minima and transition states of a tetrapeptide. The goal of the present work is to investigate this energy landscape under a variety of conditions using tools from fractal analysis. The trajectories needed for analysis were generated using “MOIL”, a molecular dynamics (MD) program which allows realistic simulations of small proteins using semi-empirical potentials. Using MOIL, the total potential energy $V(t)$ was calculated as a function of time, for several proteins of choice (PPT – Pancreatic Polypeptide, BPTI – Bovine Pancreatic Trypsine Inhibitor and Myoglobin) at constant temperature. For comparison, we also simulated poly-alanine and NaCl. The $V(t)$ curves so obtained under a variety of choices of parameters, were then subjected to a fractal analysis. The main purpose was to correlate the resultant fractal dimensions with the topology of the energy landscape, and to understand the connection to the relevant physical parameters. Both MOIL and the method used to calculate the fractal dimension of the $V(t)$ curves are described in Sec.1.

There are several reasons for using a fractal approach in the analysis of the $V(t)$ curves. Firstly, there is an intimate connection between “roughness” and fractality: much is known about the generation of rough surfaces using fractal algorithms and conversely, on the analysis of rough surfaces in fractal terms. In this sense the notion of the roughness of the energy landscape of proteins naturally calls for a fractal analysis, which can quantify the degree of roughness. Secondly, fractal dimensions often show up as “universal exponents”, and are related to the critical exponents of phase transition theory. Conversely, there is a certain universality in protein dynamics, in the sense that most proteins exhibit a folding transition and multi-exponential relaxation times. This suggests a common feature in their energy landscapes. While the rapid transition to folded states in small proteins is best rationalized by the existence of a dominant native basin of attraction (NBA) or a folding funnel, the relaxation phenomenon might well be related to the roughness of the energy landscape, and hence to its fractal dimension. A “universal” fractal dimension would be a strong characterization of what it is that is common to different energy landscapes. Thirdly, establishing that the energy landscape of a protein is indeed fractal would allow one to relate the subject of protein dynamics to a whole body of work on dynamics on fractals, e.g., diffusion on a fractal substrate.

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The main problem in trying to characterize the roughness of a high-dimensional energy landscape, such as that exhibited by a protein, is the proper choice of a reaction coordinate. In this work we have chosen time for this purpose. The advantage is that time is a universal coordinate (unlike any of the spatial coordinates), and one can think of the protein potential energy dynamics in terms of this coordinate as a random walk taking place on a rough substrate. Indeed, the rate of sampling of conformation space in complex many-body systems has been shown to be a diffusive process in the energy space. The universality of the time-coordinate, however, can also be a drawback, in that it is not very sensitive to the global topology of the energy landscape, but rather to the local structure of minima. Indeed, as will be presented in detail in Sec.III, our results appear to contain information mostly on the role of anharmonicity in determining the fractal features of the \( V(t) \) curves. As such, the results reported here reflect more on the general properties of dynamics in anharmonic potentials than on protein dynamics per se. These and other issues will be discussed and analyzed in detail in Sec.IV. Conclusions and a summary are presented in Sec.V. Finally, to establish a simple reference model, an analytical solution for the fractal dimension of the \( V(t) \) curve in the Rouse model is presented in the appendix.

II. METHODS

A. MOIL Simulations

The MOIL program has been described extensively in the literature. In a nutshell, it takes as input the x-ray structure of a given protein from the Brookhaven Protein Data Base (PDB), minimizes this structure, and runs molecular dynamics. MOIL makes it possible to simulate the energetics, dynamics and thermodynamics of systems that vary in size, up to several thousands of atoms. It accounts for a number of different energy contributions: van der Waals and electrostatic (non-covalent); bonds, angles, torsions and improper torsions (covalent):

\[
V = \sum V_{vdW} + \sum V_{electrostatic} + \sum V_{bond} + \sum V_{angle} + \sum V_{torsion} + \sum V_{improper}. \tag{1}
\]

The sums are over all interacting pairs of particles, and the components are given as follows:

\[
V_{vdW} = \frac{a_i a_j}{r_{ij}^{12}} - \frac{b_i b_j}{r_{ij}^6}, \tag{2}
\]

\[
V_{electrostatic} = \frac{q_i q_j}{\epsilon r_{ij}}, \tag{3}
\]

where \( q_i \) is the charge on atom \( i \) and \( \epsilon \) is the dielectric constant.

\[
V_{bond} = k_{ij} (r_{ij} - \langle r_{ij} \rangle)^2, \tag{4}
\]

where \( k_{ij} \) is the bond force constant. \( \langle x \rangle \) denotes the equilibrium value of coordinate \( x \).

\[
V_{angle} = \kappa_{ijk} (\theta_{ijk} - \langle \theta_{ijk} \rangle)^2, \tag{5}
\]

where \( \kappa_{ijk} \) is the angle force constant and \( \theta_{ijk} \) is the angle between two sequential bonds, i.e., bonds connecting atoms \( i, j \) and \( j, k \).

\[
V_{torsion} = \sum_{m=1}^{\infty} \alpha_{m,ijkl} \cos^m (\phi_{ijkl}). \tag{6}
\]

Here \( i, j, k, l \) are four sequentially connected atoms; \( \phi_{ijkl} \) is the angle between the planes defined by atoms \( i, j, k \) and \( j, k, l \); \( \alpha_{m,ijkl} \) is a constant which depends on the types of atoms \( i, j, k, l \) and on the periodicity of the rotation \( m \) (see Ref.[5] for details). Finally,

\[
V_{improper} = \mu_{ijkl} (\Phi_{ijkl} - \langle \Phi_{ijkl} \rangle)^2. \tag{7}
\]

Here \( j, k, l \) are all connected to atom \( i \) and \( \Phi_{ijkl} \) is the angle between the planes defined by atoms \( i, j, k \) and \( j, k, l \).

In the present work, we considered only the time dependence of the total potential energy \( V \) due to all these terms. The time series for a number of systems were generated using numerical simulations employing either the canonical
or microcanonical ensembles. As is customary, all simulations were preceded by a short initial annealing (heating) period. In the “equilibration mode” (used mainly for testing), the kinetic energies of all particles were rescaled in order to preserve the constant temperature (canonical ensemble). In the more important “no-equilibration mode” (default unless mentioned otherwise), no such rescaling was attempted and the system was allowed to reach thermal equilibrium at the specified temperature (microcanonical ensemble).

The exponents characterizing the fractal nature of $V(t)$ reported below derive from averages over groups of 25 runs differing only in the initial random velocity distribution. A number of tests were performed to ensure that our results are not artifacts of the particular integration schemes used. We checked that using more than 25 runs did not significantly affect our results statistically. Between groups of 25 but for fixed molecule-type, the runs differed, e.g., in temperature, addition of a water solvation shell, treatment of the terminus hydrogens, bond-freezing, and deletions of residues. Most runs started at a temperature of $T_i = 1\mathrm{K}$ and heated in 1K per time-step increments to the final temperature $T_f$, at which the rest of the simulation was run. The total duration of the simulations, once $T_f$ was reached, was usually 10 or 25 psec, corresponding to 20 000 or 50 000 time-steps respectively, of 0.5 fsec each. Exceptions to these conditions include some shorter and longer runs, with some variation in time-step size. They will be discussed in detail below.

**B. Fractal Analysis**

The $V(t)$ curves were subject to a standard fractal analysis, with the purpose of calculating their fractal dimension

$$\gamma = 2 - \alpha, \quad (8)$$

where $\alpha$ is the Hölder exponent, defined in the next section.

**1. General Theory**

Given any set of points $E = \{(x, y)\},$ one can estimate their fractal dimension by a number of (in principle) equivalent procedures. A popular one is the so-called “Minkowski Sausage” where one draws a circle of radius $R$ centered at each point in $E,$ and calculates the area $A(R)$ of the union (“sausage”) of all such circles. The fractal dimension is then found as

$$\gamma(E) = \lim_{R \to 0} \left[ 2 - \frac{\log[A(R)]}{\log(R)} \right] \approx \text{slope}\{\log(1/R), \log[A(R)/R^2]\} \quad (9)$$

where the second (approximate) equality follows by assuming that the first equality holds for all $R$ and multiplying it throughout by $\log(1/R).$ This assumption is equivalent to a linear relationship between $\log(1/R)$ and $\log[A(R)/R^2],$ so that $\gamma$ is found as the slope of the corresponding log-log plot. If such a linear relationship is not found, it is taken in practice as an indication that $E$ is not a fractal set. Conversely, if at least one order of magnitude of linearity is observed, the convention is to consider $E$ a fractal set in that range.

It should be noted that rigorously speaking the curve of a *function* is generally not a self-similar fractal, but rather a *self-affine* one. The latter refers to those cases where the axes have different units, so that the scaling might be different in the two directions. Indeed, a (deterministic) self-affine curve satisfies the following scaling relation:

$$h(t) = b^{-\alpha}h(b t), \quad (10)$$

which holds only on average for random processes. The “Hölder”, or self-affinity exponent $\alpha,$ is $2 - \gamma$ [Eq.(8)] if the curve is subjected to the Minkowski analysis of Eq.(9). An example is the Weierstrass–Mandelbrot function: $h(t) = \sum_{n=-\infty}^{\infty} b^{-n\alpha}(1 - \cos b^n t),$ which satisfies Eq.(10) and whose fractal dimension is $2 - \alpha.$ An equivalent characterization of statistically self-affine fractals is through the correlation function, which satisfies:

$$C_h(t) = \langle [h(t + t_0) - h(t_0)]^2 \rangle \sim t^{2\alpha}. \quad (11)$$

This relation is expected to hold for durations shorter than a typical correlation time $t_c,$ found from:

$$t_c = \int dt \frac{t \Gamma(t)}{\int dt \, \Gamma(t)}, \quad (12)$$

where $\Gamma(t) = \langle h(t_0)h(t + t_0) \rangle - \langle h(t_0) \rangle^2.$
2. Analysis of Time Series

In the present case the (self-affine) set under consideration is: \( E = \{(t, V(t))\}_t \), and the fractal analysis we applied is a numerically well-behaved version of Eq.(1); the "\( \epsilon \)-variation method". A full description of the method is beyond the scope of this paper; for full details see Ref.[15]. Briefly, Dubuc et al.[15] introduced a method particularly well suited for the evaluation of the self-affinity exponent. They demonstrated that their method has the most stable local exponent in comparison to a variety of other methods, such as box-counting and power-spectrum. In their notation, the \( \epsilon \)-variation of a function \( f \) is \( V(\epsilon, f) = \int_0^1 v(x, \epsilon) dx \), where the \( \epsilon \) oscillation \( v(x, \epsilon) \) is: \( v(x, \epsilon) = \sup_{x' \in R_\epsilon(x)} f(x') - \inf_{x' \in R_\epsilon(x)} f(x') \), and where \( R_\epsilon(x) = \{ s \in [0, 1] : |x - s| < \epsilon \} \). The corresponding log-log plot for the calculation of the self-affinity exponent is \( \{ \log(1/\epsilon), \log(V(\epsilon, f)/\epsilon^2) \} \), with the exponent given by 2–slope. We applied this analysis in order to extract the self-affinity exponent (which we refer to simply as the "fractal dimension" and denote by \( \gamma \)) of our \( V(t) \) curves.

III. RESULTS

In this section we will present the findings from our numerical simulations. A discussion will be postponed to the next section. We will start by presenting a sample of "raw" simulation results, then move on to a detailed presentation of each of the three proteins, compare them amongst each other, perform a comparison with several test cases, estimate the effect of time-scales of different processes, and finally study the role of anharmonicity vs harmonicity. All our MD simulations started out from the folded state (as determined by X-ray), which we subsequently minimized using the Powell algorithm. The protein was then heated from 1K in increments of 1K at every time-step, with a random initial velocity distribution, until the desired final temperature was reached. Fractal analysis was performed on data from the constant temperature period only. Dynamics on all three proteins was run for 25ps, with a time-step of 0.5ps (with a few exceptions, to be detailed below). Fractal analysis was performed on the full 25ps trajectories and on the first 10ps of each such trajectory. This process was repeated for a wide selection of temperatures in the range 1K-800K, with a focus on the room temperature regime.

A. “Raw” Results

A graph of a typical \( V(t) \) curve is displayed in Fig.1 for BPTI at 310K. We only present the time series after the temperature is stabilized, i.e., the first 0.4ps during which annealing from 1K occurs are omitted. The noisy data are strikingly similar to "fractional Brownian motion" and clearly suggest the use of fractal-type analysis. This is done in Fig.2, where the corresponding \( \epsilon \)-variation analysis is shown. A straight line can be fitted with confidence to the \( \epsilon \)-variation data over some 1.5 orders of magnitude (decades). The linear regression coefficient in this case is: \( R^2 = 0.998 \), where 1 indicates a perfect straight line. The fractal dimension (slope in Fig.2) is: \( \gamma = 1.762 \pm 0.004 \), where the uncertainty is due to the linear regression. All log-log plots look strikingly similar to the one in Fig.2 and have a similar range of linearity. The \( R^2 \) coefficient rarely went below 0.99. Fig.3 presents all 25 fractal dimensions \( \gamma \) obtained from the runs for BPTI under the abovementioned conditions. The dashed line is the average fractal dimension, which in this case yields: \( \gamma = 1.75 \pm 0.02 \). The uncertainty in this number reflects both the individual linear regression errors on each data point, as well the standard deviation about the average. The procedure illustrated in Figs.1-3 was repeated throughout this work. Thus every \( \gamma \) data point in the figures to follow is the result of an average over 25 fractal dimensions obtained from independent runs differing only in the initial velocity distribution.

B. Myoglobin

Fig.4 shows the results of our simulations of myoglobin. Every \( \gamma \) point was obtained as discussed above, so the error bars are due to both the averaging over the 25 data points and the linear regression error on each of these points. The dashed (solid) line connects the \( \gamma \) values resulting from the 25ps (10ps) runs. The inset shows the room temperature results. In the 10ps case \( \gamma \) appears to have a shallow minimum around 10K, which is however absent in the full 25ps simulations. The error bars are in fact too large to confidently discuss a trend in the curves. However, it is clear that the fractal dimension increases with the simulation (or observation) time \( \tau \); on average \( \gamma \approx 1.67 \) for 10ps and rises to \( \gamma \approx 1.72 \) for 25ps. This behavior is seen in all our results ahead (see Figs. 6 and 7).

As is clear from the inset, the fractal exponent \( \gamma \) is essentially constant around room temperature, apart from the rather sharp drop at 330K.
The robustness of the results was tested by running the simulations at 300K and 400K with the heme group deleted. The removal of the heme group makes myoglobin considerably less compact, causing it to unfold rather easily. Hence we expected a significant change in the underlying dynamics. However, Fig. 4 shows (squares and circles) that the influence of heme deletion is very minor, and results in a small increase in $\gamma$.

C. BPTI

The results of our simulations of BPTI are shown in Fig. 5. The overall behavior is very similar to that of myoglobin. The differences are (1) overall the fractal dimensions are higher; (2) the minimum is more pronounced (beyond statistical error), visible also in the 25psec data, and has shifted to 50K.

The increase in $\gamma$ with increasing observation time for the longer simulation is unmistakable for BPTI as well.

We also performed a 10psec simulation including a water solvation shell (diamond). Clearly, this has no significant effect on the results.

As a consistency test, we calculated $\gamma$ for the complementary 10-25psec of the simulation (triangle). The result is a $\gamma$ value in between the 10psec and 25psec values.

Finally, the dip observed for myoglobin can be seen here as well as in the 25psec simulation, at T=320K.

D. PPT

Our PPT simulation results appear in Fig. 6. The general qualitative similarity to BPTI and myoglobin is noticeable, namely the increase in overall fractal dimension with simulation time, and the shallow minimum at around 50K. Several additional details are noteworthy: (1) We included a 50psec long simulation at 1K (same 0.5fsec time step), and the increase in $\gamma$ persists (full circle). (2) We also performed simulations in the “equilibration mode” (see Sec. II A). These are indicated as squares and can be seen most clearly in the inset. The results are virtually identical to the usual “no-equilibration mode”, demonstrating the agreement between the canonical and microcanonical ensembles for large (> 100 atoms) systems. The only exception is the point at 800K, where the “no-equilibration” value of $\gamma$ drops rather sharply. (3) A water solvation shell (diamonds, also in the equilibration mode) again does not produce statistically significant results.

E. Comparison Among Myoglobin, BPTI and PPT

Figs. 7 and 8 compare the results for the three proteins in the 10 and 25psec cases, respectively. These figures merely recapitulate the results from Figs. 1-6 and show no new data. The main points to notice are the striking similarity between PPT and BPTI, compared to the overall lower fractal dimension values for myoglobin. This holds in both the 10 and 25psec cases.

F. PPT Under a Variety of Conditions

We performed further tests of the robustness of our results by running 10psec simulations of PPT at 300K, under different conditions, as shown in Fig. 9. There are 11 data points in this figure, and the $\gamma$ values are arranged in increasing order. Let us briefly describe the results, noting that case 7 is the “standard” one (i.e., the result shown in Fig. 6).

The case “no $\alpha$-helix” refers to a simulation in which the entire PPT $\alpha$-helix was deleted, i.e., only the GLY PRO SER GLN PRO THR TYR PRO GLY ASP ASP groups remain. Under these conditions a significantly smaller $\gamma$ value is obtained. On the other hand data point 5 is the opposite case, where only the $\alpha$-helix remains. The corresponding $\gamma$ is much closer to the reference value of data point 7. This shows that the $\alpha$-helix is by and large responsible for the observed fractal dimension.

Data point 2 corresponds to a freezing of all bond vibrations. The remaining motions are therefore proper and improper rotations and center of mass translations. Since the deviation from the reference $\gamma$ value is small, it appears that the rotations are largely responsible for the observed fractal dimension.

Data point 3 is the result when a different integrator is used, which appears to have only a minor effect. This test was performed to check whether the results are a consequence of numerical instabilities.
Data point 4 is another deletion experiment, this time of just the SER residue. This has a very small effect which, curiously enough, is slightly larger than leaving only the α-helix, as indicated in case 5 (although due to the large error bars in fact this effect cannot be taken too seriously).

To calculate the γ corresponding to data point 6, the terminal hydrogens were treated as “extended”: unlike in all other cases, the hydrogens were not treated as atoms but were “absorbed” into the carbons. This modifies the carbon mass and radius to appropriate effective values. The effect, however is negligible.

Data points 8 and 10 relate to using the equilibration mode and adding a water solvation shell respectively, which (as shown also in Fig.6) do not have a significant effect either.

Data point 9 reports the result from a single 250psec long trajectory: the trajectory was divided up into 10 segments of 25psec each, and the γ’s were averaged. Data point 11 is due to a 100psec long trajectory which, however, employed a time step of 5fsec (10 times larger than the usual time step). In accordance with the observation of a fractal dimension which increases over time, the resulting average γ of data point 9, as well as that of data point 11, are larger than the reference value.

G. PPT vs Test Systems: Poly-alanine and NaCl

In order to estimate to what extent our results reflects properties unique to realistic proteins, we present calculations for two test systems: poly (16)-alanine and a 40-atom cluster of NaCl. Poly-alanine folds into an α-helix, so it has the crucial part of the protein topology (recall points 1 and 5 in Fig.9), but the underlying energy landscape is not expected to be rugged. This is because all the residues are identical so there is no frustration due to conflicting interactions between different segments of the polymer. (Nevertheless each residue has internal conflicting interactions, giving rise to some ruggedness). The ground state of the non-polymeric NaCl cluster is easily accessible, i.e., starting from a disordered state, it finds its crystalline ground state well before the end of our 10psec simulation. Thus it has neither the protein topology, nor its rugged energy landscape. The results of our simulations are shown in Fig.10. The 10 and 25psec runs of 16-alanine and NaCl at 250K, as well as 10psec at 300K, and 10psec for NaCl alone at 1K. For clarity of presentation the error bars have been removed. However, it should be remarked that when the error bars are taken into account the results for a given temperature overlap. Indeed, it is striking that the results for poly-alanine and NaCl do not differ significantly from those for PPT. In fact for 250K, NaCl and PPT agree almost exactly, and the agreement for 10psec at both 250K and 300K is quite close. Only at 1K does there seem to be a significant difference between NaCl and PPT. 16-alanine has lower γ values than PPT, but not very much so. Barring accidental agreement, these results clearly indicate that neither the protein topology, nor the rugged energy landscape are essential in order to obtain fractal \( V(t) \) curves, with fractal dimension close to that of realistic proteins.

H. The Role of Observation Time: Short Simulations of PPT

In order to systematically investigate the role of the total duration of the simulation and the correspondingly participating physical processes, we ran simulations of 20000 time steps of PPT at 300K, with progressively smaller time steps. This is different from the bulk of our simulations where a constant time step of 0.5fsec was employed, so care should be exercised in the comparison. The main point is that by employing a smaller time step, previously inaccessible fast motions are now observable. The results are shown in Fig.11. It is remarkable that the fractal dimension obeys an almost perfect logarithmic law over three orders of magnitudes:

\[
\gamma(\tau) = 1.60 + 0.060 \ln(\tau),
\]

for \( 0.1\text{psec} \leq \tau \leq 100\text{psec} \). The regression coefficient is \( R^2 = 0.999 \). Fig.11 thus demonstrates decisively that γ is indeed strongly dependent on the total observation time.

Below \( \tau = 100\text{fsec} \) (to the left of the arrows in Fig.11) the behavior of γ appears rather erratic and we believe this is due to numerical error. Indeed, the fastest motions in the protein are the OH or CH stretches, with a period of about 10fsec. Thus \( \tau = 10\text{fsec} \) would most likely not probe even a single period and can confidently be regarded as the lower physical limit of our simulations.

We turn next to an analysis and discussion of our results.

IV. DISCUSSION

The most striking observation presented in the previous section is the rise of the fractal dimension with time. It suggests (at least) two likely hypotheses for its explanation: As time evolves
1. there is increased access to local minima and “fine-structure” of the potential energy landscape;
2. low frequency vibrations and energy transfer between modes become increasingly activated.

The first of these attributes the growth in $\gamma$ over time to the increased roughness of the sampled potential energy landscape. In other words, the protein “visits” more and more local minima (or basins of attraction) as its samples its conformational space. This process is consistent with an increase in fractal dimension since the fractal potential energy landscape is not sampled all at once by the protein: instead the inner details are only progressively revealed. However, as shown by Czerminski and Elber, activated transitions between similar conformational structures confined to a small set of minima (separated by barriers on the order of $k_B T$) can occur at room temperature on a time scale of several picoseconds. A much larger time scale is required for such transitions at temperatures on the order of 1K. The insensitivity of $\gamma$ to temperature therefore appears to rule out the first hypothesis: at the low temperature end one would expect the protein to be confined to a single minimum within the time scale of our simulations, and the fractal dimension to be smaller as a consequence. Thus while increased access to local minima is consistent with the growth of $\gamma$ with time, it is inconsistent with the lack of temperature dependence. We conclude that the fractal dimension measured in our simulations is not a good probe of the global potential energy landscape topology.

This brings us to the second hypothesis which, as will be argued shortly, is consistent with both the observed time and temperature dependencies. According to this hypothesis, the primary factor determining the fractal dimension of the potential energy landscape is the self-generated dynamical noise, namely the very broad frequency spectrum appearing in the dynamics. This spectrum and the resulting self-generated noise is a consequence of the anharmonicity in the potential, as is demonstrated convincingly in Fig.12. Shown in this figure, along with the full 10psec simulation of PPT at 300K, are the results of the same simulations with all anharmonic interactions switched off: terms $V_{\text{vdW}}$, $V_{\text{electrostatic}}$ and $V_{\text{torsion}}$ are missing from the total potential energy $V$ of Eq.(6). The effect is dramatic: there is a strong decrease of $\gamma$ as the temperature is lowered. This shows that anharmonicity is of paramount importance in determining $\gamma$, at least up to 100K.

The mechanism we propose is thus as follows: $V(t)$ is a sum over many anharmonic terms, the number of which is proportional to the number of atoms in the protein. These anharmonic terms create overtones, which effectively fill the frequency spectrum and create a quasi-continuum. Now, on the one hand, as time evolves (at constant temperature) both very low and very high (previously inactive) frequency components become activated: the low frequencies are only sampled on long times, whereas the high frequencies correspond to highly energetic modes which must await energy transfer before equilibration sets in. On the other hand, as temperature increases (at constant $\tau$) intermode coupling causing energy transfer from rigid modes to soft modes is facilitated, and more frequency components are activated. Thus in both cases one expects an increase in dynamical noise, with a corresponding increase in the fractal dimension of the signal. This accounts for the behavior seen in Figs.11. Unfortunately we have not been able to study whether the fractal dimension converges on a limiting value upon full equilibration (at least as a function of temperature this does not seem to be the case – see Figs.6 and 7), since this requires prohibitively long simulations.

The behavior of the harmonic case observed in Fig.13 might seem curious in light of these arguments: after all in the purely harmonic case there is no intermode coupling, nor any dynamical noise. However, we recall that due to the use of non-Cartesian coordinates [see Eqs.(4), (6) and (7)], a certain residual degree of anharmonicity remains with respect to the Cartesian components. At very low temperatures vibrations and rotations occur very close to their equilibrium values and the anharmonic effects are barely noticeable. Only as the temperature rises do these vibrations and rotations start to deviate significantly from their equilibrium values and gain anharmonicity. This explains the rise of $\gamma$ in the “harmonic” case. In contrast, the other terms [Eqs.(2), (6) and (7)] make a strong anharmonic contribution at all temperatures, which accounts for the relative constancy of $\gamma$ in this case.

The different behavior of PPT and BPTI vs myoglobin can be attributed to the size of these proteins. Myoglobin has 156 residues, whereas PPT and BPTI only have 41 and 60 residues respectively. Therefore at any given temperature it takes longer to activate the overtones that give rise to the self-generated noise in myoglobin than in either PPT or BPTI, which explains the lower fractal dimension of myoglobin in Figs.7 and 8. The proximity in size between PPT and BPTI, on the other hand, is the reason for their quantitatively similar behavior. A related argument makes clear why deletion of the heavy heme group from myoglobin yields a higher $\gamma$ (Fig.9): without heme the energy redistribution among the modes occurs faster.

Also the curious similarity between NaCl and PPT can be attributed to anharmonic effects: NaCl, with its purely Coulombic interactions, is of course strongly anharmonic, even more so than PPT. On the other hand poly-alanine is more harmonic, so as expected it has lower $\gamma$ values.

It is thus seen that the “anharmonicity” hypothesis is capable of accounting for all of the salient features present in our results for the fractal dimension of the $V(t)$ curves. The role of multiple minima and ruggedness of the energy landscape seems to be more restricted: it is apparently not the global topology which is measured by $\gamma$. Nevertheless, it should be noted that different multiple minima will generally lead to a denser filling of the frequency spectrum or to the appearance of bands, which may overlap and thus induce additional intermode coupling.
An important remaining issue concerns the value of $\gamma$ observed in our simulations: definitely greater than 1.5 (except perhaps for the $\tau = 100$sec simulation, Fig.11). *A priori* it is not clear what the lower limit of $\gamma$ for polymer-like systems subject to the usual dynamics (like Newtonian or Langevin) should be. A plausible lower limit of 1.5 is derived in the appendix for the Rouse model. The Rouse model considers a purely harmonic set of beads under the influence of a Gaussian random force. In our simulations (in the "no-equilibration" – microcanonical mode) there is no randomness except for numerical truncation errors and "randomness" due to chaos caused by the anharmonic terms. However, we find $\gamma \approx 1.5$ as the result of our "harmonic" simulations at very low temperature (Fig.12). As argued above, this actually corresponds to the low-anharmonicity limit, so is in agreement with the result for the Rouse model, provided the truncation errors and/or the chaotic "randomness" can be modelled as a Gaussian random force acting on the protein (this is a speculation). Increasing the degree of anharmonicity can only increase the fractal dimension of the signal, by the mechanisms discussed before. These arguments then show that the Rouse model result could thus serve as a *lower bound* to the $\gamma$ values found in our simulations.

V. SUMMARY AND CONCLUSIONS

In this work we have studied the fractal properties of the total potential energy $V$ as a function of time $t$ for several realistic proteins. We performed simulations at a wide range of temperatures and a number of time scales. Our main conclusion is that the fractal behavior seems to be caused by the self-generated dynamical noise due to intermode coupling and equilibration promoted by anharmonic effects. There are universal aspects to this behavior, as exemplified by the similarity between totally different systems such as PPT and NaCl. Thus it cannot serve to characterize individual proteins and their energy landscape, but is rather a property common to all sufficiently large and anharmonic systems. We believe, however, that probing the fractal aspects of protein potential energy landscapes is a useful approach to the characterization of the ruggedness of these landscapes. To achieve this, it would be very interesting to examine $V$ as a function of appropriate spatial coordinates, rather than $t$. The question remains whether there are other ways leading to useful geometrical characterizations of the energy landscape, e.g., identification of multiple minima and maxima by real and imaginary frequencies.

*Note added:* While this paper was in the final stages of preparation we became aware of an article by García et al. The authors concluded that the time dependence of the mean-square displacement of crambin around the crystal structure is described by an effective fractal exponent $\gamma \approx 1.60$ (for $t$ long). This result was interpreted as indicative of multi-basin dynamics which would be typical in a rugged energy landscape.

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APPENDIX

AN ANALYTICALLY SOLVABLE EXAMPLE: THE ROUSE MODEL

It is useful to have a simple reference model in the background to compare the numerical results of Sec.III with. To this end, we consider the Rouse Model (RM), which is standard in polymer theory. It describes a polymer as a set of $N$ beads located at $(R_1, R_2, ..., R_N)$ connected along a chain, where each bead $(n)$ undergoes Brownian motion under the influence of random force $f_n(t)$. It should be remarked that this is essentially different from our simulations, in which no randomness (except for numerical truncation errors and chaotic "randomness" due to anharmonicity) is present. Nevertheless, it useful to study the RM in order to see analytically how a fractal $V(t)$ can come about.

The excluded volume and hydrodynamic interactions are disregarded in the RM. The dynamics is generally described by the Langevin equation:

$$\frac{\partial}{\partial t} R_n(t) = \sum_m H_{nm} \cdot \left( f_m(t) - \frac{\partial U}{\partial R_m} \right) + \frac{1}{2} k_B T \sum_m \frac{\partial}{\partial R_m} \cdot H_{nm}. \quad (A1)$$
In the RM the mobility tensor is assumed to be isotropic:

\[ H_{nm} = \frac{I}{\zeta} \delta_{nm}, \]  

(A2)

and the interaction potential is taken to be harmonic:

\[ U = \frac{k}{2} \sum_{n=2}^{N} (R_n - R_{n-1})^2, \]  

(A3)

so that Eq.(A1) reduces to

\[ \zeta \frac{dR_n}{dt} = -k(2R_n - R_{n+1} - R_{n-1}) + f_n \quad (n = 2, 3, ..., N - 1) \]  

(A4)

\[ \zeta \frac{dR_1}{dt} = -k(R_1 - R_2) + f_1 \]  

(A5)

\[ \zeta \frac{dR_N}{dt} = -k(R_N - R_{N-1}) + f_N. \]  

(A6)

The force and the potential define a time scale through

\[ \tau \equiv \frac{\zeta}{k}. \]  

(A7)

It is convenient to consider the last set of equations in the limit where \( n \) is taken as a continuous variable:

\[ \zeta \frac{\partial R_n}{\partial t} = k \frac{\partial^2 R_n}{\partial n^2} + f_N \]  

(A8)

\[ \frac{\partial R_{n=0}}{\partial n} = \frac{\partial R_{n=N}}{\partial n} = 0. \]

The random forces are assumed to be Gaussian, i.e., delta-function uncorrelated:

\[ \langle f_n(t) \rangle = 0 \]  

(A9)

\[ \langle f_n(t)f_m(t') \rangle = 2\zeta k_B T \delta(n - m) \delta(t - t'). \]

In the continuum limit the potential energy becomes:

\[ U = \frac{k}{2} \int_{0}^{N} dn \left( \frac{\partial R_n}{\partial n} \right)^2. \]  

(A10)

The last three sets of equations define the continuum RM.

1. **Normal Coordinates**

We are interested in calculating the fractal scaling of the potential energy \( U(t) \) defined in Eq.(A10). For this purpose it is convenient to transform to normal coordinates. It can be shown that in terms of the coordinates

\[ X_p \equiv \frac{1}{N} \int_{0}^{N} dn \cos(p\pi n/N)R_n(t) \quad p = 0, 1, 2, ... \]  

(A11)

the Langevin equation (A3) becomes

\[ \zeta_p \frac{dX_p}{dt} = -k_p X_p + f_p \]  

(A12)

where
\[ \zeta_0 = N\zeta \]  
\[ \zeta_p = 2N\zeta \quad \text{for } p = 1, 2, \ldots \]  
\[ k_p = \frac{2\pi^2 kp^2}{N} \quad \text{for } p = 0, 1, 2, \ldots \]  
and the forces satisfy

\[ \langle f_p(t) \rangle = 0 \]  
\[ \langle f_p(t)f_q(t') \rangle = 2\zeta_p k_B T \delta_{pq}\delta(t-t'). \]

In this representation the random forces as well as the motions of the \( X_p \)'s are independent, so that the motion of the polymer is decomposed into independent modes. Indeed, it is easily verified that the formal solution to Eq. (A12) is

\[ X_p(t) = \frac{1}{\zeta_p} \int_{-\infty}^{t} dt' e^{-\frac{(t-t')}{\tau_p}} f_p(t'), \]  
where

\[ \tau_p = \frac{\tau_1}{p^2}, \quad \tau_1 = \frac{\zeta_1}{k_1} = \frac{N^2}{\pi^2}\tau. \]

Finally, the inverse transform of Eq. (A11) is

\[ R_n = X_0 + 2\sum_{p=1}^{\infty} X_p \cos(p\pi n/N). \]  

### 2. Scaling of the Potential Energy

We will now present a calculation of the fractal scaling of the potential energy in the RM, as defined in Eq. (11): \[ C_U(t) = \langle (U(t) - U(0))^2 \rangle. \]  

Inserting the expression for \( R_n \) into Eq. (A10) and expanding yields:

\[ U = 2k \left( \frac{\pi}{N} \right)^2 \sum_{p,q=1}^{N} p^2 q^2 X_p(t) \cdot X_q(t) \int_0^{N} dn \sin(p\pi n/N)\sin(q\pi n/N). \]

The integral evaluates to \( \frac{N}{2}\delta_{pq} \), so that:

\[ U = k \frac{\pi^2}{N} \sum_{p=1}^{N} p^2 X_p^2(t). \]

Thus the fluctuation can be written as:

\[ C_U(t) = \left( k \frac{\pi^2}{N} \right)^2 \left\langle \left( \sum_{n=1}^{N} p^2 (X_p^2(t) - X_p^2(0)) \right)^2 \right\rangle \]

\[ = \left( k \frac{\pi^2}{N} \right)^2 \sum_{p,q=1}^{N} p^2 q^2 \left[ (X_p^2(t)X_q^2(t)) - 2(X_p^2(t)X_q^2(0)) + (X_p^2(0)X_q^2(0)) \right]. \]

Inserting the formal solution for the normal coordinates, Eq. (A18), the averages can be calculated from:
\[
\langle X_p^2(t)X_q^2(t') \rangle = \frac{1}{\zeta_p \zeta_q} \int_{-\infty}^{t} dt_1 \int_{-\infty}^{t'} dt_2 \int_{-\infty}^{t} dt_3 \int_{-\infty}^{t'} dt_4 \times \\
e^{-\frac{(2t-t_1-t_2)}{\tau_p} e^{-\frac{(2t'-t_3-t_4)}{\tau_q}}} \langle f_p(t_1) \cdot f_p(t_2) \cdot f_q(t_3) \cdot f_q(t_4) \rangle
\] (A25)

To proceed the averages over the forces must be evaluated. This can be done with the help of Wick’s theorem:

\[
\langle x_{n_1}x_{n_2} \cdots x_{n_{2p}} \rangle = \sum_{\text{all pairings}} \langle x_{m_1}x_{m_2} \cdots \langle x_{m_{2p-1}}x_{m_{2p}} \rangle \rangle
\] (A26)

where the \(x_n\)’s are Gaussian random variables. Employing the orthogonality relations from Eq. (A17) thus yields:

\[
(2k_B T)^2 \left[ \zeta_p \zeta_q \delta(t_1-t_2)\delta(t_3-t_4) + \zeta_p^2 \delta_{pq} \delta(t_1-t_3)\delta(t_2-t_4) + \zeta_q^2 \delta_{pq} \delta(t_1-t_4)\delta(t_2-t_3) \right]
\] (A27)

In addition we need the rule

\[
\int_{-\infty}^{t} dt_2 f(t-t_2)\delta(t_2-t_1) = f(t-t_1)\Theta(t-t_1)
\] (A28)

where \(\Theta(x)\) is the Heavyside step-function. Combining these results, the delta functions simplify the expression in Eq. (A26) as follows:

\[
(2k_B T)^2 \left[ \frac{1}{\zeta_p \zeta_q} \int_{-\infty}^{t} dt_1 e^{-\frac{(t-t_1)}{\tau_p+1/\tau_1}} \int_{-\infty}^{t'} dt_3 e^{-\frac{(t'-t_3)}{\tau_1+1/\tau_1}} + \right.
\]
\[
\frac{2 \delta_{pq}}{\zeta_p^2} \left( \int_{-\infty}^{\min(t,t')} dt_1 e^{-\frac{(t+t'-2t_1)}{\tau_p}} \right)^2
\]
\[
(2k_B T)^2 \left[ \left( \frac{\tau_p \tau_q}{\tau_p + \tau_q} \right)^2 \frac{1}{\zeta_p \zeta_q} + 2 \delta_{pq} \frac{\tau_p^2}{\zeta_p^2} 2(2\min(t,t')-t')/\tau_p \right].
\] (A29)

Inserting this into Eq. (A25) yields:

\[
C_U(t) = \left( \frac{k_B T}{N} \right)^2 \sum_{p,q=1}^{N} p^2 q^2 \left[ \left( \frac{\tau_p \tau_q}{\tau_p + \tau_q} \right)^2 \frac{1}{\zeta_p \zeta_q} + 4 \frac{\tau_p^2}{\zeta_p^2} \left( 1 - e^{2t/\tau_p} \right) \delta_{pq} \right].
\] (A30)

Using the definitions of the various parameters appearing here the final result becomes:

\[
C_U(t) = (k_B T)^2 \left[ \sum_{p,q=1}^{N} \frac{1}{p^2 + q^2} + 4 \sum_{p=1}^{N} \left( 1 - e^{-2\pi^2 + \frac{q^2}{N^2}} \right) \right].
\] (A31)

The time dependence of \(C_U(t)\) for large but finite \(N\) arises from the term \(\sum_{p=1}^{N} \exp(-2\pi^2 tp^2/\tau N^2)\). To derive an expression for \(\gamma\), it is useful to approximate the above sums by integrals. Defining \(x = p/N\), \(y = q/N\), we have:

\[
C_U(t) \approx (k_B T)^2 \left[ 4N + \frac{1}{N^2} \int_{1/N}^{1} dx \int_{1/N}^{1} dy \frac{1}{x^2 + y^2} - 4 \int_{1/N}^{1} dx e^{-2\pi^2 x^2} \right].
\] (A32)

The first integral is easily evaluated after a transformation to polar coordinates, and the second corresponds to the error-function (in the second integral the error in taking \(1/N = 0\) is negligible):

\[
C_U(t) \approx (k_B T)^2 \left[ 4N + 2\pi \ln N \frac{N^2}{N^2} - 2\sqrt{\pi} \frac{\text{erf}(\sqrt{\theta})}{\sqrt{\theta}} \right], \quad \theta = 2\pi^2 t/\tau.
\] (A33)

In accordance with the preceding comments we next take the \(t \ll \tau\) limit. Then, using \(\text{erf}(x) \approx 2/\sqrt{\pi}(x - x^3/3! + x^5/5! - ...)\) to the first non-trivial (third) order, we can write:
\[ C_U(t \ll \tau) \approx (k_B T)^2 \left[ 4N + 2\pi \frac{\ln N}{N^2} - 4(1 - \frac{1}{3} \theta) \right] \]
\[ \sim (k_B T)^2 \frac{8\pi^2 t}{3} \sim t^{2(2 - \gamma)}, \]  
(A34)

where in the last line the time-independent terms were ignored, and ultimately \( \gamma = 3/2 \). Thus the Rouse model has a potential energy fluctuations which have a fractal dimension of 3/2, just like a classical, fully correlated random walk.

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FIG. 1. Total potential energy as a function of time for a single run of BPTI at 310K. Annealing from 1K occurs during the first 0.4 psec, which are not shown. The highly irregular fluctuations are reminiscent of fractional Brownian noise.

FIG. 2. $\epsilon$-variation result for the curve shown in Fig. 1. A straight line can be observed for about 1.5 decades. The bending at high $\epsilon$ values is due to finite size effects.
FIG. 3. All 25 fractal dimensions calculated for BPTI at 310K. Individual error bars are due to linear regression. The dashed line is the average fractal dimension.

FIG. 4. Fractal dimension as a function of temperature for myoglobin. Lines connecting data points are guides to the eye only.

FIG. 5. Fractal dimension as a function of temperature for BPTI.
FIG. 6. Fractal dimension as a function of temperature for PPT.

FIG. 7. Comparison among myoglobin, BPTI and PPT for 10psec simulations.
FIG. 8. Comparison among myoglobin, BPTI and PPT for 25psec simulations.

![Graph showing comparison among myoglobin, BPTI and PPT for 25psec simulations.]

FIG. 9. A comparison of results for PPT at 300K, 10psec, under a variety of conditions.

![Graph showing a comparison of results for PPT at 300K, 10psec, under various conditions.]

FIG. 10. A comparison between PPT, 16-alanine and NaCl (40 atoms). Error bars have been omitted for clarity and are replaced by small dots.

![Graph comparing PPT, 16-alanine and NaCl (40 atoms). Error bars are omitted for clarity and replaced by small dots.]
FIG. 11. 20000 time step simulations of PPT at 300K with varying total duration (x-axis). Results to the left of the arrows are in the unphysical range.

FIG. 12. 10psec simulation results of PPT with the full potential (solid line) and just the harmonic part (dashed line).