Conformational gel analysis and graphics: Measurement of side chain rotational isomer populations by NMR and molecular mechanics

Christopher Haydock
Applied New Science LLC, Rochester, Minnesota 55901, USA

Conformational gel analysis and graphics systematically identifies and evaluates plausible alternatives to the side chain conformations found by conventional peptide or protein structure determination methods. The proposed analysis determines the populations of side chain rotational isomers and the probability distribution of these populations. The following steps are repeated for each side chain of a peptide or protein: first, extract the local molecular mechanics of side chain rotational isomerization from a single representative global conformation; second, expand the predominant set of rotational isomers to include all probable rotational isomers down to those that constitute just a small percentage of the population; and third, evaluate the constraints vicinal coupling constants and NOESY cross relaxation rates place on rotational isomer populations. In this article we apply conformational gel analysis to the cobalt glycyl-leucine dipeptide and detail the steps necessary to generalize the analysis to other amino acid side chains in other peptides and proteins. For a side chain buried within a protein interior, it is noteworthy that the set of probable rotational isomers may contain one or more rotational isomers that are not identified by conventional NMR structure determination methods. In cases such as this the conformational gel graphics fully accounts for the interplay of molecular mechanics and NMR data constraints on the population estimates. The analysis is particularly suited to identifying side chain rotational isomers that constitute a small percentage of the population, but nevertheless might be structurally and functionally very significant.

I. INTRODUCTION

Simple changes in functional groups can often make multiple order-of-magnitude changes in biological activity [1]. This suggests that protein conformations populated at the 10%, 1.0%, or even 0.1% level may play a very significant role in function.

The most common NMR protein structure determination methods generate perhaps a few dozen complete protein structures [2]. The structures in this ensemble are independently fit to the data and each final structure should fit the data about equally well. The optimization of a multiconformer model differs sharply from these standard fitting methods because the measure of goodness-of-fit is not defined for any single structure in the ensemble, but depends simultaneously on the entire ensemble of structures. In contrast to the standard methods, no single structure in the final multiconformer ensemble need be a particularly good fit to the data [3]. In favorable cases multiconformer models can give some indication of the conformational variability of proteins in solution [4]. For example, particular secondary structure elements, loops, or even side chains of the multiconformer model might have a larger RMS deviation indicating variability of these parts in solution. Information about local variability thus comes from a global fitting procedure. Whether conformational variability is assessed by conventional methods or multiconformer models, the essential idea is to narrow down the vast global conformational space of a protein by applying the constraints of real data.

This point also marks one of the key differences of conformational gel analysis because conformational gel analysis identifies local conformations based solely on molecular mechanics and a single representative global conformation previously determined from current or preexisting NMR data or even crystallographic data. Instead of applying NMR data to pick out new global conformations, the NMR data is analyzed to determine the extent to which it constrains the populations of the local conformations identified by molecular mechanics.

Detailed information about local conformations is often available from molecular mechanics. In the case of protein side chain rotational isomerization, good estimates of the position and shape of potential energy wells are known and approximate depths of these wells are also known [5]. Even in cases where such information might be exploited to reduce or eliminate the conformational search problem, its potential use is overlooked or even dismissed, apparently because the experimental data is judged more reliable than the molecular mechanics results [6], the molecular mechanics geometries and energies may be judged globally coupled to such an extent that local conformational information can not be separated out [7, 8, 9], or perhaps because the molecular mechanics models are not readily available [10, 11, 12].

The cobalt glycyl-leucine dipeptide [22] NMR data analyzed in this article was previously analyzed in a more comprehensive and perhaps less accessible manner [13]. The present analysis of this data is not designed to extend the knowledge of the cobalt dipeptide system, but rather to be readily generalizable to proteins. The cobalt dipeptide analysis is particularly suited to generalization to proteins because the cobalt chelate ring system fixes the dipeptide backbone in a definite conformation and

*Electronic address: haydock@appliednewscience.com
FIG. 1: Stereo views of the predominant rotational isomers of the leucine side chain of the cobalt glycyl-leucine dipeptide: top, trans gauche$^+$; middle, gauche$^-$ trans; and bottom, gauche$^+$ gauche$^-$. The atom gray scale tones are: white, hydrogen; light gray, carbon; medium gray, nitrogen; dark gray, oxygen; black, cobalt. The leucine side chain projects outward towards the viewer and the three chelated nitro groups are visible below, behind, and above the cobalt dipeptide ring system.

FIG. 2: Molecular mechanics energy schematic for the three predominant leucine side chain rotational isomers. Isomerization energies are plotted as a function of the $\chi^1$ torsion angle for rotation about the leucine side chain $\alpha$ to $\beta$-carbon bond. The horizontal axis labels refer to only the $\chi^1$ torsion angle of each rotational isomer. Solid, molecular mechanics energy function; dashed, NMR experimental data energy function; dotted, energy function distribution from estimated molecular mechanics errors. This energy function distribution displays essentially the same information as a gel graphic.

the simplifying assumption of a single backbone conformation also applies to the analysis of proteins.

II. CONFORMATIONAL GEL ANALYSIS AND GRAPHICS

Conformational gel analysis and graphics provides detailed molecular mechanics population estimates and assesses the constraints that NMR data places on these population estimates. Much of this information is expressed in the form of gel graphics. Though gels are widely applied to the separation, characterization, and identification of all types of biomolecules and their developed images are widely seen in the biochemical literature, the gel graphics employed here are entirely computer generated and their interpretation is in many ways novel. The basic facts about their interpretation are perhaps most easily explained by showing their connection to molecular mechanics energy plots. We will make this connection in the opening paragraphs of this section in a highly simplified analysis of the three predominant rotational isomers of the leucine side chain of the cobalt dipeptide (Figure 1) and then move on to a detailed explanation of conformational gel analysis and graphics in the three numbered subsections of this section.

A gel graphic can convey the uncertainty of rotational isomer populations obtained either by molecular mechanics calculations or by fitting NMR data. The distinction between calculated or fit populations and the uncertainties of these populations parallels that between a simple energy function and an energy function distribution (Figure 2). In this example all the energy functions give energies for rotation about the leucine side chain $\alpha$ to $\beta$-carbon bond. The three troughs of each sinusoidal function are the energy wells of the three predominant rotational isomers and the three crests are the energy barriers to interconversion. Note that the molecular mechanics energy function (Figure 2 solid) can be calculated from the full $\chi^1 \times \chi^2$ energy map (Figure 3) by rotating the $\chi^1$ torsion angle and as appropriate adjusting the $\chi^2$ torsion angle in such a way as to pass over the energy barriers separating the three predominant rotational isomers, see methods.

The molecular mechanics energy function is calculated from an empirical energy function, which has many parameters, such as bond length and bond angle equilib-
Figure 3: Molecular mechanics energy map for rotational isomerization of cobalt dipeptide leucine side chain. Contour levels are dashed, 1, 3, 5, 7, 9; solid, 2, 4, 6, 8, 10; dotted, 15, 20 kcal/mol. Zero corresponds to −39.4 kcal/mol. The nine rotational isomers are labeled at the position of their energy well minima.

| Rotational Isomer | Population |
|------------------|------------|
| g−g−             | 0.0        |
| g+g−             | 0.2        |
| tg−              | 0.4        |
| g−g+             | 0.6        |
| g+g+             | 0.8        |
| tg+              | 1.0        |
| g−t              | 0.0        |
| g+t              | 0.2        |
| tt               | 0.0        |

The Boltzmann factor establishes a relation between the energy and the population of a state. The connection between rotational isomer population uncertainties and an energy function distribution follows from this relation. For example the energies at the trans, gauche−, and gauche+ well minima of the molecular mechanics energy function are 0.5, 0.4, and 2.4 kcal/mol (Figure 2, solid) and at 300 degrees K the Boltzmann factors for these energies give populations of 45, 53, and 2% for the three predominant rotational isomers. It is also important to note that though it is meaningful to talk about the energy of rotational isomerization of a single molecule, the population must refer to an ensemble average over many molecules or at least to a time average for a single molecule. Even though there is an ensemble of molecules there is no uncertainty in the populations predicted by the simple energy function. The population probability distributions shown in the gel graphic are calculated for an uncertainty of the map energy well depths of ±1.0 kcal/mol and a temperature of 300 K. Each gray scale step of the stepwedge bar corresponds to a two-fold change in probability density.
FIG. 5: Conformational gel graphic for the entire probable set of rotational isomers of the cobalt dipeptide leucine side chain. The gel graphic visually portrays the extent to which the NMR data constrains the populations of the probable set of isomers. The population probability distributions shown in the gel graphic are constructed by repeated fitting of the rotational isomer populations to Monte Carlo NMR data. Each gray scale step of the stepwedge bar corresponds to a two-fold change in probability density.

FIG. 6: Conformational gel graphic for the predominant set of rotational isomers of the cobalt dipeptide leucine side chain. The restriction of the probable set of rotational isomers to those that molecular mechanics suggest are predominant clearly reduces the uncertainty in the populations. For a protein side chain such a restriction is not always desirable because less conspicuous rotational isomers might be structurally and functionally very significant.

gives explicit visual representation to the errors inherent in molecular mechanics.

The interpretation of conformational gel graphics (Figures 5 and 6) is in many respects similar to that of a molecular mechanics gel graphic as described above. Again we will focus on the populations estimates for the three predominant rotational isomers (Figure 6, lanes 5–7) and leave detailed consideration of the entire probable set of rotational isomers to a later subsection. If a model with only the trans, gauche\(^-\), and gauche\(^+\) predominant rotational isomers is fit to the NMR data for the cobalt dipeptide, the predicted populations of these rotational isomers are 21, 54, and 25%. Though it is natural to fit the NMR data by adjusting rotational isomer populations rather than well energies, these fit populations may be converted to energies by inverting the Boltzmann factor procedure described in the previous paragraph. At 300 degrees K the fit populations are equivalent to a simple energy function with trans, gauche\(^-\), and gauche\(^+\) energy well minima at 0.9, 0.4, and 0.8 kcal/mol (Figure 2, dotted). The standard Monte Carlo method for estimating the distribution of the rotational isomer population estimates assumes that the estimated populations are the true populations, generates a large number of simulated NMR data sets, and fits these simulated NMR data sets to generate the population distribution, see methods. The gel graphic (Figure 6, lanes 5–7) displays this population distribution. By the inverted Boltzmann factor procedure the population distribution could be converted into an energy function distribution and plotted in an energy schematic. In the same way that the molecular mechanics gel graphic (Figure 4, lanes 5–7) has an equivalent energy function distribution (Figure 2, dotted) so also does a conformational gel graphic (Figure 6, lanes 5–7) have an equivalent energy function distribution (not shown). Only the origins of the distributions differ. The molecular mechanics energy function distribution is generated directly by applying Monte Carlo energy errors to a simple molecular mechanics energy function, but the energy function distribution equivalent to a conformational gel is generated from the underlying rotational isomer population distribution, which is in turn indirectly generated from Monte Carlo NMR data errors as outlined above. In short, the conformational gel graphic gives explicit visual representation to the errors inherent in the NMR data.

We now turn to a detailed explanation of the conformational gel analysis and graphics of the cobalt dipeptide and along the way indicate how this analysis can be generalized to proteins.
1. Extract local molecular mechanics from a single representative global conformation

In this work we confine our attention to the rotational isomerization of peptide and protein side chains as examples of local molecular mechanics. The cobalt dipeptide energy map for the rotational isomerization of the leucine side chain (Figure 3) is a very simple example of local molecular mechanics. To calculate this map the leucine side chain torsion angles are constrained to values on a 5 degree grid over the $\chi^1 \times \chi^2$ torsion space and the entire dipeptide structure is energy minimized [13]. To extract local molecular mechanics of proteins local backbone flexibility and local side chain interactions must be carefully controlled [6]. A single representative global conformation can be routinely extracted from an ensemble of NMR protein structures by averaging and constrained minimization [2].

Though only the side chain conformation at an energy minimum of a well is required for the gel analysis, the entire map is useful for automated identification of the energy minima and is absolutely essential for correctly controlling the local backbone flexibility and neighbor side chain interactions. To control the backbone flexibility the entire protein backbone is fixed except for backbone segments of two or at most three amino acids. Essentially, the number and length of these free atom segments must be sufficient to accurately determine the position of the energy well minima, but this number and these lengths must not be so unduly generous as to make energy minimizations unnecessarily expensive or as to obscure the energy map with transitions of nonlocal backbone degrees of freedom. The energy map is very effective tool for eliminating these nonlocal transitions because they show up as discontinuities of the energy surface. If a map has these discontinuities then it must be recomputed with reduced backbone flexibility. The accuracy of the energy map can be judged by comparing the position, shape and depth of energy wells of maps computed at two or three different levels of backbone flexibility.

The effects of neighbor side chain interactions are assessed by truncating neighbor side chains at the $\beta$-carbon. By comparing the shape and position of energy wells of maps calculated with and without neighbor side chain truncation it is possible to gauge the extent to which particular interactions influence particular side chain rotamers. In rare cases a neighbor side chain interaction may be judged to preclude a particular energy well at any potentially interesting level of population. More commonly a neighbor side chain interaction will simply increase the uncertainty of a target well’s energy depth because the uncertainty of the interaction strengths of the target well with all of the neighbor side chain wells and the uncertainty in the energy depth of all the neighbor wells must be folded into the uncertainty of the target well’s depth.

The energy map for the leucine side chain of the cobalt dipeptide (Figure 3) is similar to maps computed for side chains in a variety of protein environments. One important similarity is that during calculation of the side chain energy map the cobalt dipeptide backbone is fixed in a definite conformation by the cobalt chelate ring system. This parallels the approach to calculating a protein side chain energy map described above, where the protein backbone is fixed in the same conformation as found in the single representative global conformation. The cobalt dipeptide and protein maps both give the local molecular mechanics of side chain rotational isomerization. The leucine map of the cobalt dipeptide is also similar to protein side chain maps in that the total number of wells present in the map is the same or close to the number of ideally possible rotational isomers, for example, nine in the case of a leucine side chain. This is true even for a side chain buried within a protein because neighbor side chains are usually truncated at the $\beta$-carbon in order not to exclude target side chain conformations that interact unfavorably with the neighbor side chain conformation that happens to exist in the single representative global conformation of the protein. As already mentioned the uncertainty of the energy depth of such target side chain wells is substantially increased. The positions of the well energy minima on the energy map for the leucine side chain of the cobalt dipeptide differ from their ideal positions by from about 10 degrees (trans gauche$^+$ rotational isomer) to almost 60 degrees (trans trans rotational isomer). The average departure from ideal positions tends to increase for side chains buried within proteins.

2. Expand the predominant set to include all probable rotational isomers

The single representative global conformation determined by conventional methods already gives a preferred conformation for each side chain of a protein. For each side chain it is usually possible to adjoint the preferred rotational isomer with one or two others and form a predominant set of rotational isomers that account for say 90% of the population of the side chains rotational isomers. The side chain rotational isomers included in this predominant set might be present in an ensemble of conformations determined by conventional NMR, would probably have low energy wells in the molecular mechanics energy map, and might possibly be suggested by rotamer preference libraries compiled from the protein data bank. There are several different scenarios that may arise and these are best illustrated by referring to the cobalt dipeptide molecular mechanics gel graphic (Figure 1).

The molecular mechanics gel graphic plays an important role both in identifying the predominant set of rotational isomers and in expanding this set to make the set of all probable rotational isomers. As discussed in the opening paragraphs of this section this gel graphic differs from the energy map in that it takes into account not only the energy depth of each rotational isomer’s well, but also the uncertainty of the energy well depths. A comparison of
cobalt dipeptide gel graphic and gel graphics computed for side chains in a variety of protein environments follows along much the same lines as the comparisons between energy maps in the previous section. Like a protein energy map, the resulting protein molecular mechanics gel reflects the local molecular mechanics of the target side chain. Just as the protein energy map usually has energy wells corresponding to each of the ideally possible rotational isomers, the resulting protein molecular mechanics gel has the same corresponding lanes. Unlike the well depth uncertainties of the cobalt dipeptide molecular mechanics gel, which are all equal because there are no neighbor side chains, the uncertainties of a protein gel would be significantly larger for rotational isomers that interact strongly with neighbor side chains. It is difficult to draw reliable conclusions from the molecular mechanics energy map without taking these uncertainties into account.

From the cobalt dipeptide molecular mechanics gel graphic (Figure 4) the predominant rotational isomers of the leucine side chain are trans gauche$^+$ and gauche$^-$ trans and possibly gauche$^+$ gauche$^-$. Analysis of NMR data suggests that these first two rotational isomers are the most populated and that the third may also make a significant contribution [13]. This agreement between experiment and molecular mechanics comes about because the experimental analysis is designed to reproduce the rotational isomer preferences observed in the protein data bank [12], which not coincidently are much the same as the relative rotational isomer stabilities predicted by molecular mechanics. We expect that for many protein side chains the same set of predominant rotational isomers will be given by the molecular mechanics gel graphic and the ensemble of conformations determined by conventional NMR, but for slightly different reasons than those just mentioned for the cobalt dipeptide leucine side chain. Perhaps the most important reason that the molecular mechanics energy map is likely to predict the same predominant rotational isomers is that it is computed with the backbone fixed in the same conformation as found in the single representative global conformation. The side chain energy map is thus likely to favor the rotational isomers populated in the ensemble of conventional NMR conformations because the map is computed with the backbone fixed in a conformation that is representative of this very same ensemble.

The set of all probable rotational isomers includes all the predominant rotational isomers as well as those that make smaller contributions to the population down to contributions as small as perhaps a few tenths of a percent. The molecular mechanics gel graphic for the cobalt dipeptide leucine side chain suggests that all rotational isomers except gauche$^+$ gauche$^-$ are in the probable set. This is shown visually by a single band at zero population and a complete lack of any upwards extension of this band. As mentioned earlier in this section the molecular mechanics gel graphics of protein side chains have lanes corresponding to most of the ideally possible rotational isomers. Even for a side chain buried in a protein interior a good fraction of these will at least be in the probable set of rotational isomers. Compared to the leucine side chain of the cobalt dipeptide the distinction between predominant and probable sets of rotational isomers may not be as clear cut because of the increased energy uncertainty of rotational isomers that strongly interact with a neighbor side chain. A large uncertainty shows up on the molecular mechanics gel graphic as a lane with bands at population zero and one and a much fainter extensions stretching between the these two extremes. Rotational isomers with such a large molecular mechanics energy uncertainty should certainly be included in the probable set and may even be sufficiently populated to be included in the predominant set [2].

3. Evaluate the constraints vicinal coupling constants and NOESY cross relaxation rates place on rotational isomer populations

Thus far we have described the molecular mechanics of the leucine side chain of the cobalt dipeptide, the selection of predominant and probable rotational isomers, and how this can be generalized to side chains of proteins. The key point about the generalization to proteins is that the molecular mechanics remains local. By local it is meant that the molecular mechanics model depends on the single representative global conformation, which is always readily available from the preexisting NMR structure. The side chain molecular mechanics does not depend on a multiconformer model, which would in some way require the simultaneous solution of all the side chain rotational isomer populations. In this section the molecular mechanics model is fit to the NMR data to evaluate the populations of the predominant and the probable rotational isomers. This can be generalized to proteins by exploiting the locality of both the molecular mechanics model and NMR data to carry out the evaluation independently for each side chain.

The conformational gel graphic for the probable set of rotational isomers of the cobalt dipeptide leucine side chain (Figure 5) shows that the experimental data, which consists of eight vicinal coupling constants and ten NOESY cross relaxation rates [13], places little constraint on the populations of these eight isomers. This is indicated by the intense bands extending from zero population up to thirty to fifty percent population for each of the rotational isomers in the probable set. A comparison of the molecular mechanics and conformational gel graphics (Figures 4 and 5) gives a striking graphical portrayal of the dramatic variation in the usefulness of molecular mechanics and NMR data for determining rotational isomer populations. Apparently, the populations of most of the cobalt dipeptide side chain rotational isomers are best determined either by the NMR data alone or by the molecular mechanics calculations alone. Except for the three more extended lanes (gauche$^+$ gauche$^+$, trans gauche$^+$,
and gauche$^-$ trans) near the middle of the molecular mechanics gel graphic (Figure 1) all the rest of the rotational isomers have bands at zero population that at most have a relatively small upwards extension. These rotational isomer populations are better determined by molecular mechanics. In contrast the trans gauche$^+$ and gauche$^-$ trans rotational isomers have bands stretching all the way from population zero up to one in the molecular mechanics gel graphic (Figure 1) as compared to much less extended bands in the conformational gel graphic (Figure 5). These rotational isomer populations are better determined by fitting the NMR data. The similarity of the gauche$^+$ gauche$^-$ rotational isomer bands in both figures suggests both molecular mechanics and NMR data must be taken into account to determine the population of this rotational isomer. All these conclusions are born out by more detailed analysis [13].

Clearly it is desirable to reduce the size of the probable set of rotational isomers to the point that the NMR data does place significant constraints on the populations. In the case of the cobalt dipeptide this point is reached when the probable set is reduced to the three rotational isomers of the predominant set defined in the previous section. The conformational gel graphic for the predominant set (Figure 6) displays population errors that are somewhat to considerably smaller than the predicted populations of the rotational isomers. This is displayed visually by the modest to large distance from the zero population horizontal grid line and to the beginning of the high density region of the bands. Note that the bands extend nearly three standard deviations above and below the mean, but the high density region extends only about two standard deviations in each direction. Comparing the conformational gel graphics for the probable and the predominant sets of rotational isomers (Figures 5 and 6) a particularly striking improvement is seen in the significance of the population estimate of the gauche$^-$ trans rotational isomer.

For any protein side chain it is in principle possible to define a predominant set of rotational isomers that is just small enough to yield significant population estimates. The practical difficulty with this is that the molecular mechanics does not always give reliable ordering of the energy well depths because of the sources of error discussed in the previous section. As the size of the probable set is reduced it will not always be clear which rotational isomers to include or exclude. All three gel graphics (Figures 1, 5, and 6) must be considered together to obtain a complete picture of the rotational isomer populations that fully accounts for the interplay of molecular mechanics and NMR data. A still more detailed analysis should also consider measurability and over-fitting of rotational isomer populations [13], but an easily accessible description of the application of these concepts is beyond the scope of this article.

III. CONCLUSIONS

This work makes new theoretical predictions of interest to a broad range of chemists studying the structure and function of proteins or other complex molecules. Particularly important is the prediction that the local molecular mechanics of protein side chains can be extracted from a single representative global conformation determined by conventional methods. The local molecular mechanics can identify low population though potentially functional rotational isomers of buried protein side chains. Conformational gel analysis and graphics is an important new tool for display and understanding of conformational population estimates and of the sources and level of errors in these estimates. By helping us see more clearly the extent of both our knowledge and our ignorance we hope to fuel the demand and inspire and guide the development of more powerful NMR instruments and analysis methods.

IV. METHODS

Detailed descriptions as well as working computer input files for calculating molecular mechanics energy maps, fitting NMR data, and generating gel graphics, have been previously published [5, 13]. Briefly, custom topology and parameter input files were created and $\chi^1 \times \chi^2$ energy maps for the leucine side chain of the cobalt dipeptide were calculated with the CHARMM molecular mechanics program [10]. Based on the positions of the energy well minima nine energy minimized rotational isomers were prepared and interatomic distances and torsion angles for modeling cross relaxation rates and vicinal coupling constants were output with the CHARMM correlation and time series analysis command. The optimization design matrix was obtained by a MATLAB function file that input a list of NOESY cross relaxing protons and vicinally coupled spins, read in the appropriate distance and angle data files output by the analysis command, calculated the cross relaxation rates and the vicinal coupling constants for each rotational isomer, and normalized each of these observables by a composite experimental and cross relaxation rate or Karplus coefficient error. Note that the original version of this MATLAB function file [13] was somewhat more complex than described here because it was designed to examine the effects of intramolecular motions by averaging over the molecular mechanics energy map. The rotational isomer populations were fit by minimizing the differences between the experimentally measured and predicted observables subject to the constraints that the populations were nonnegative and that their sum was one. This linear least-squares with linear constraints problem was solved as the equivalent quadratic programming problem [17]. The probability density functions of the fit rotational isomer populations were computed by the standard Monte Carlo recipe [18]: the experimental NMR observables
were fit to yield fit rotational isomer populations and fit NMR observables, random errors were added to the fit NMR observables and these simulated NMR observables were fit to give simulated fit populations, these last steps were repeated many times to make the Monte Carlo probability density functions of the populations.

The energy functions in the energy schematic were obtained by cubic interpolation from the positions of the energy minima and maxima. The derivatives of the interpolating functions were constrained to zero at the positions of these minima and maxima. Each energy function was shifted by an energy constant so that at 300 degrees K the sum of the Boltzmann factors of the energy minima equaled one. The molecular mechanics energy function was obtained from the molecular mechanics energy map by matching the energies of the predominant rotational isomer minima and the energies of the lowest energy barriers between the predominant rotational isomers. The horizontal positions of the energy function minima and maxima were adjusted somewhat to make the maximum slopes of the energy barriers about equal while still approximately matching the rotation of the $\chi^1$ torsion angle. The horizontal axis of the energy schematic is labeled steric rotation angle rather than $\chi^1$ torsion angle to reflect this approximation and to emphasize the steric relationship between the side chain atoms rather than the exact rotation angle.

The molecular mechanics gel graphic was generated from the energy map by a simple Monte Carlo procedure. Random energy errors were added to the rotational isomer well depths and Boltzmann weighted populations were calculated and normalized, these steps were repeated many times, and the resulting large set of simulated rotational isomer populations was histogramed to make Monte Carlo probability density functions of the populations.

Monte Carlo probability density functions were displayed as gel graphics, which were designed to visually indicate both the discrete probability fraction at zero population and shape of the continuous probability density over the range of population from zero and one. This was accomplished by a simulated photographic process where the degree of film overexposure indicates the probability fraction at zero population and continuous gray tones represent the continuous part of the probability density. To simulate film overexposure at zero population and smooth the lane edges along the continuous part of the probability distribution a Gaussian blur filter was applied to the image so that the typical probability density at zero population was still considerably greater than that along the continuous part of the probability distribution. The pixel values of the blurred image were treated like scene luminances [19] and converted into photographic print densities with a characteristic curve [20] that had a maximum point-gamma of 1.5. The print densities were linearly mapped into gray scale values so that maximum printable density was somewhere along the continuous part of the probability distribution. A stepwedge bar of the 11 zones in the Zone System [21] was added to the gel graphic as an aid to calibrating the probability densities.

Vector PostScript molecular graphics were generated with the RasMol program [23].

V. ACKNOWLEDGMENT

This work was financially supported by Franklyn G. Prendergast in the Department of Biochemistry and Molecular Biology at the Mayo Clinic in Rochester, Minnesota, USA.

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