Determination of Multiple ϕ -Torsion Angles in Proteins by Selective and Extensive ¹³C Labeling and Two-Dimensional Solid-State NMR

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We describe an approach to efficiently determine the backbone conformation of solid proteins that utilizes selective and extensive ¹³C labeling in conjunction with two-dimensional magic-anglespinning NMR. The selective ¹³C labeling approach aims to reduce line broadening and other multispin complications encountered in solid-state NMR of uniformly labeled proteins while still enhancing the sensitivity of NMR spectra. It is achieved by using specifically labeled glucose or glycerol as the sole carbon source in the protein expression medium. For amino acids synthesized in the linear part of the biosynthetic pathways, [1-13C]glucose preferentially labels the ends of the side chains, while [2-13C]glycerol labels the C^{\alpha} of these residues. Amino acids produced from the citricacid cycle are labeled in a more complex manner. Information on the secondary structure of such a labeled protein was obtained by measuring multiple backbone torsion angles ϕ simultaneously, using an isotropic-anisotropic 2D correlation technique, the HNCH experiment. Initial experiments for resonance assignment of a selectively ¹³C labeled protein were performed using ¹⁵N-¹³C 2D correlation spectroscopy. From the time dependence of the ¹⁵N-¹³C dipolar coherence transfer, both intraresidue and interresidue connectivities can be observed, thus yielding partial sequential assignment. We demonstrate the selective ¹³C labeling and these 2D NMR experiments on a 8.5-kDa model protein, ubiquitin. This isotope-edited NMR approach is expected to facilitate the structure determination of proteins in the solid state. © 1999 Academic Press

Key Words: ¹³C labeling; proteins; torsion angle; resonance assignment; solid-state NMR.

INTRODUCTION

Recently solid-state NMR has been increasingly applied to the structure determination of noncrystalline and disordered biological molecules. Techniques that yield both distance and orientational constraints to high precision are now available. The sensitivity of these NMR experiments is greatly enhanced by the enrichment of NMR-sensitive isotopes such as ¹³C and ¹⁵N. So far, most NMR structural studies of solid peptides and proteins have adopted a specific-labeling approach whereby a small number of sites in the molecule is labeled in each synthesis and then one or two distances or torsion angle constraints are measured in each experiment (*1*–*7*). To determine protein structures more efficiently and comprehensively, it is

desirable to employ more extensive isotopic labeling approaches and to extract multiple structural constraints from each experiment. The second condition may be fulfilled by performing two-dimensional (2D) NMR experiments where one dimension separates and identifies the chemically distinct sites through their isotropic chemical shifts, while the other dimension yields the structural parameters (8). The first condition can be fulfilled by uniform 13C and 15N labeling of the proteins, which is used routinely now in solution NMR to determine the structure of globular proteins (9, 10). In solidstate NMR, uniform ¹⁵N labeling has also been utilized to obtain information such as the orientation of protein segments in the membrane bilayer (11). However, uniform ¹³C labeling of proteins leads to several spectroscopic difficulties for solidstate NMR. First, the one-bond ¹³C-¹³C *J*-couplings and the multispin dipolar couplings that are effective in fully ¹³Clabeled proteins cause significant line broadening. The J-coupling-induced line broadening cannot be removed by magicangle spinning (MAS) alone (12, 13) while the dipolar-induced line broadening is only removed completely at very high spinning speeds (>25 kHz) (14). Thus the spectral resolution of fully ¹³C-labeled proteins is limited under normal MAS speeds (3–15 kHz). Second, the weak ¹³C–¹³C dipolar couplings that reflect long-range ¹³C-¹³C distances are obscured by the strong couplings between directly bonded 13C spins. This occurs by spin diffusion within the dense ¹³C spin network and by the dipolar truncation mechanism (15, 16). These spectroscopic challenges, combined with the high cost of ¹³C labeling, have hindered the routine measurement of multiple structural constraints in solid proteins from a single experiment and on a single sample.

In this paper we explore selective and extensive 13 C labeling as a means to facilitate the NMR investigation of solid proteins and demonstrate the application of ϕ torsion measurement on such a labeled model protein, ubiquitin. By exploiting the biosynthetic pathways of amino acids in bacteria, we can incorporate 13 C labels into some sites in the proteins while leaving other sites unlabeled. Easily feasible with the standard protocol for protein expression, the selective and extensive (S&E) labeling approach reduces the number of directly-bonded 13 C spin pairs or larger clusters in the molecule, either by not labeling one of the carbons in the chain, or by reducing



the labeling levels of the sites involved. The principal advantage of this S&E labeling approach for solid-state NMR lies in the resolution enhancement that it achieves. This is demonstrated by the 13 C spectra of several differently labeled ubiquitin ($M_r = 8565$, 76 residues). The abundant X-ray crystal diffraction (17) and solution-state NMR (18–21) data that are available for ubiquitin allow us to verify the 13 C labeling protocol and to demonstrate the solid-state NMR techniques. The modest molecular weight of ubiquitin also makes it possible to assess the realistic sensitivities of the NMR experiments. Similar selective 13 C labeling approaches have been applied in solution NMR for measuring precise relaxation parameters (22, 23); however, to our best knowledge this is the first demonstration of selective 13 C labeling for solid-state NMR investigation of protein structure.

Using a 13 C selectively and 15 N uniformly labeled ubiquitin sample, we demonstrate the residue-specific measurement of backbone torsion angles ϕ by correlating the N–H and C^{α} –H $^{\alpha}$ bond orientations (24). The ϕ -angle experiment yielded site-specific information on the backbone conformation with an accuracy of ± 5 to $\pm 30^{\circ}$, by comparison with the X-ray crystal structure. Since structure determination requires the assignment of spectral resonances to the residues in the protein sequence, we applied a 15 N– 13 C 2D correlation technique and demonstrate that it is possible to achieve partial sequential assignment directly in the solid state.

RESULTS AND DISCUSSION

1. Selective and Extensive ¹³C Labeling

The selective and extensive labeling method relies on the specificity of the amino acid biosynthetic pathways in bacteria. Under aerobic conditions, bacteria can utilize glucose or glycerol as the sole carbon source to synthesize the 20 amino acids. This is achieved via three major enzymatic pathways: glycolysis, the pentose phosphate pathway, and the citric-acid cycle (25, 26). As an example, we outline the metabolic fates of the glucose C1 carbon in Fig. 1. During glycolysis, the six-carbon glucose is oxidized to the three-carbon pyruvate, converting glucose C1 into the pyruvate methyl carbon, C3. Since glucose C1 and C6 contribute equally and indistinguishably to pyruvate C3, only 50% of pyruvate results from glucose C1. Pyruvate is next oxidized to acetyl-CoA, which enters the citric-acid cycle, where two key intermediates, α -ketoglutarate and oxaloacetate, serve as the precursors for 10 amino acids. Oxaloacetate is also produced directly from phosphoenolpyruvate and pyruvate through several anaplerotic reactions. Both α -ketoglutarate and oxaloacetate are reused in the cyclical reactions, thus labeling multiple carbon sites, particularly C^{α} , C^{β} , and C^{γ} , of these 10 amino acids. In contrast, amino acids synthesized by the linear enzymatic reactions have fewer glucose C1-derived carbons. From pyruvate C3, the side chain ends of Ala, Val, and Leu are labeled. Gly loses the glucose C1 label in its synthesis from 3-phosphoglycerate, which also generates Ser and Cys with

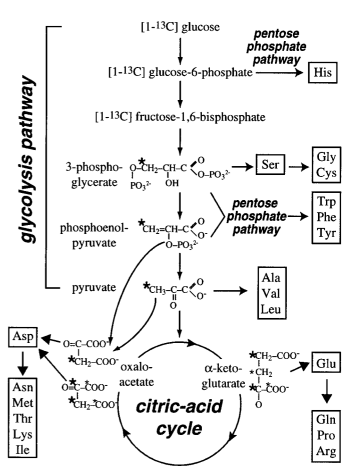


FIG. 1. Amino acid biosynthetic pathways in bacteria, illustrating selective and extensive 13 C labeling of proteins. Most compounds in the pathways are precursors for the 20 amino acids. The fate of the glucose C1 is traced. The resulting labeled carbons in the amino acid precursors are designated by asterisks. The smaller asterisks in oxaloacetate and α -ketoglutarate indicate lower labeling levels.

labeled side chain ends. The 3 aromatic amino acids are produced from phosphoenolpyruvate via the pentose phosphate pathway and are labeled at C^{β} and several ring positions.

Therefore, we can obtain a protein that is selectively labeled in ¹³C by supplementing the bacterial growth media with ¹³C specifically labeled glucose such as [1-¹³C]glucose. The absence of labeling for certain sites will alleviate the line broadening, spin diffusion, and dipolar truncation problems mentioned above. The chemical symmetry of the glucose molecule can be exploited to control the maximum labeling level to 50%. Thus, the probability of a ¹³C spin pair is at most 25% even for two directly bonded labeled sites. This further reduces the influence of the undesirable J-couplings. For amino acids produced from the citric-acid cycle, the labeling levels of many sites are reduced to \sim 25% due to the loss of some ¹³C labels as CO₂ and the distribution of labels between chemically indistinguishable sites by two symmetric intermediates (succinate and fumarate) of the cycle. The lower labeling levels compensate for the extensiveness of labeling in these amino acids. When a labeling

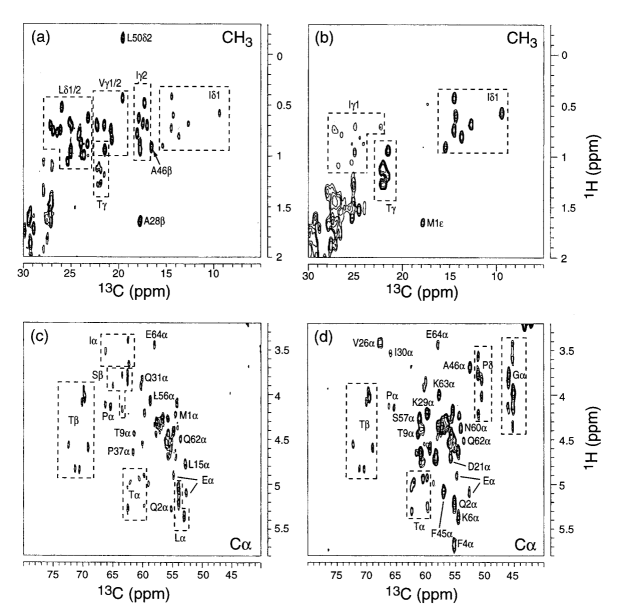


FIG. 2. Solution NMR $^{13}\text{C}-^{1}\text{H}$ heteronuclear correlation spectra of $^{13}\text{C1}$ -ubiquitin and $^{13}\text{C2}$ -ubiquitin, verifying the selective and extensive ^{13}C labeling patterns. (a) CH₃ region of $^{13}\text{C1}$ -ubiquitin. (b) CH₃ region of $^{13}\text{C2}$ -ubiquitin. (c) C $^{\alpha}$ -H $^{\alpha}$ region of $^{13}\text{C1}$ -ubiquitin. (d) C $^{\alpha}$ -H $^{\alpha}$ region of $^{13}\text{C2}$ -ubiquitin. Spectral assignment is based on the known chemical shifts of this protein (27) and is indicated for some peaks to illustrate the selectivity of the labeling and the partial complementary nature of the $^{13}\text{C1}$ and $^{13}\text{C2}$ labeling schemes. Areas of the same type of carbons are enclosed by dashed lines.

level of 100% is desired, [1, 6-¹³C]glucose may be used. This selective labeling approach can also be extended to specifically labeled glycerol and pyruvate, which undergo similar enzymatic reactions and have been used in several solution NMR studies of protein dynamics (22, 23).

To demonstrate this selective and extensive ¹³C labeling approach, we obtained two ubiquitin samples that were expressed from bacterial media supplemented with [1-¹³C]glucose and [2-¹³C]glycerol as the sole carbon source. To verify and compare the distribution of ¹³C-labeled sites, we employed 2D ¹³C-¹H correlation experiments on these samples dissolved in aqueous solution (Fig. 2). The labeled carbon sites were identified by using the solution NMR chemical shifts known

for this protein (27) and are compiled in Tables 1 and 2. As expected, the two labeling schemes exhibit partial complementarity, in that many sites labeled by $[1^{-13}C]$ glucose are not labeled by $[2^{-13}C]$ glycerol, and vice versa. For example, in the $[1^{-13}C]$ glucose-derived ubiquitin (heretofore referred to as $^{13}C1$ -ubiquitin), most side chain methyl carbons are labeled without directly bonded ^{13}C neighbors (Fig. 2a). In comparison, the $[2^{-13}C]$ glycerol-derived ubiquitin (heretofore referred to as $^{13}C2$ -ubiquitin) exhibits poor methyl carbon labeling except for Ile $C^{\delta}1$ and Thr C^{α} (Fig. 2b). The isolated methyl carbon labels are potentially useful for measuring long-range $^{13}C^{-15}N$ and $^{13}C^{-13}C$ distances between residues to constrain the protein tertiary structure. The $[2^{-13}C]$ glycerol scheme pref-

TABLE 1

13C Labeling Pattern of [1-13C]Glucose

	¹³ C labels			¹³ C labels	
Residue	Theory	Expt.	Residue	Theory	Expt.
Gly	None	None	Glu	$[\mathrm{CO}^a],lpha,eta,\gamma,-$	α, β, γ
Cys^b	β	N.A.	Gln	$[CO^a]$, α , β , γ , $-$	α, β, γ
Ser	β	β	Pro	$[CO^a]$, α , β , γ , $-$	α , β , γ , $-$
Ala	β	β	Arg	$[CO^a]$, α , β , γ , $-$, $-$	α , $-$, γ , $-$, $-$
Leu	α , δ 1, δ 2	α , δ 1, δ 2	Asp	$[\mathrm{CO}^a],lpha,eta,[\gamma^a]$	α , β
Val	$\gamma 1, \gamma 2$	$\gamma 1, \gamma 2$	Asn	$[CO^a], \alpha, \beta, [\gamma^a]$	α , β
His	CO^1	δ2	Met	$[CO^a]$, α , β , $[\gamma]$	$\alpha, \beta, -$
Phe	β , δ 1, δ 2	β , γ , δ 1, δ 2	Thr	$[CO^a]$, α , β , $[\gamma]$	$\alpha, \beta, [\gamma]$
Tyr	β , δ 1, δ 2	β , δ 1, δ 2, ζ	Lys	$-, \beta, [\gamma], \delta, \epsilon$	$-$, β , δ , ϵ
Trp^b	β, δ1, δ2	N.A.	Ile	[CO 1], α , $-$, γ 1 γ 2, [δ 1]	α , $-$, γ 1, γ 2, $[\delta 1]$

Note. The theoretically predicted labeled sites are compared with the experimentally observed ones obtained from the solution NMR data in Fig. 2a. For amino acids produced from the citric-acid cycle (right half of the table), two types of labeling levels are predicted: the normal level of \sim 50% (no brackets), and a lower labeling level of \sim 25% (square brackets). The experimental peak intensities, classified as strong (no brackets) and weak (square brackets), agree well with the theoretical predictions. For amino acids produced from the citric-acid cycle, dashes are used to indicate the unlabeled sites.

erentially labels the 13 C $^{\alpha}$ of the linear-chain amino acids, while the [1- 13 C]glucose scheme does not. Isolated 13 C $^{\alpha}$ sites are useful for measuring ϕ torsion angles to directly probe the protein secondary structure. Those amino acids with both 13 C $^{\alpha}$ and 13 CO labels are suitable for ψ torsion angle measurements using the NCCN technique (28, 29) and techniques that correlate the 13 CO chemical shift anisotropy with the 13 C $^{\alpha}$ - 1 H $^{\alpha}$ dipolar coupling (30, 31).

Overall, the predicted labeling patterns are borne out well by the solution NMR spectra, as shown in Tables 1 and 2. No comparison is made for the two missing amino acid types in ubiquitin and for the unprotonated carbons, which are not detectable by the experiment. The good agreement between the experiment and the prediction indicates that label scrambling is negligible. Based on residual signals of glycine C^{α} in the $^{15}N-^{13}C$ correlation spectra of $^{13}C1$ -ubiquitin (Fig. 6), we estimate that scrambling contributes $\sim 10\%$ of the labeling. Low scrambling is crucial for securing the selectivity of labeling, especially for amino acids produced from the linear part of the biosynthetic pathways. If desired, the more extensive labeling

TABLE 2

13C Labeling Pattern of [2-13C]Glycerol

Residue	¹³ C labels			¹³ C labels	
	Theory	Expt.	Residue	Theory	Expt.
Gly	α	α	Glu	$[\mathrm{CO}^a], (\alpha), [eta], -, \delta^a$	$[\alpha], [\beta], -,$
Cys^b	α	N. A.	Gln	$[CO^a]$, (α) , $[\beta]$, $-$, δ^a	$[\alpha], [\beta], -$
Ser	α	α	Pro	$[CO^a]$, (α) , $[\beta]$, $-$, δ	$[\alpha], [\beta], -, \delta$
Ala	α	α	Arg	$[CO^a]$, (α) , $[\beta]$, $-$, δ	$[\alpha], [\beta], -, \delta$
Leu	CO^a , β , γ	β, γ	Asp	$[\mathrm{CO}^a], [lpha], (eta), [\gamma^a]$	$[\alpha], [\beta]$
Val	α, β	α, β	Asn	$[\mathrm{CO}^a], [lpha], (eta), [\gamma^a]$	$[\alpha], [\beta]$
His	α	δ2	Met	$[\mathrm{CO}^a], [lpha], (eta), [\gamma]$	$[\alpha], -, [\gamma], \epsilon$
Phe	$\alpha, \gamma, \epsilon 1$	$\alpha, \gamma, \epsilon 1$	Thr	$[\mathrm{CO}^a], [\alpha], (eta), [\gamma]$	$[\alpha], [\beta], \gamma$
Tyr	$\alpha, \gamma, \epsilon 1$	$\alpha, \gamma, \epsilon 1, \zeta$	Lys	α , $-$, $[\gamma]$, (δ) , $[\epsilon]$	α , $-$, γ , $[\delta]$, ϵ
Trp^b	α , $\delta 2$, $\zeta 3$	N. A.	Ile	$[CO^a], [\alpha], \beta, (\gamma 1), [\delta 1]$	$[\alpha], [\beta], [\gamma 1], \delta 1$

Note. The theoretically predicted labeled sites are compared with the experimentally observed ones obtained from the solution NMR data in Fig. 2b. For amino acids produced from the citric-acid cycle, three types of labeling levels are predicted: the normal labeling level is \sim 100% sites in square brackets are labeled at \sim 50%, and sites in round brackets are labeled at <25%. The experimental peak intensities, classified as strong (no brackets) and weak (square brackets), show general agreement with the theoretical predictions. For amino acids produced from the citric-acid cycle, dashes are used to indicate the unlabeled sites.

^a Carbon sites predicted to be labeled but cannot be confirmed by the ¹³C-¹H correlation experiment due to the lack of directly bonded protons.

^b Amino acids not present in ubiquitin. The experimental data are not available (N. A.).

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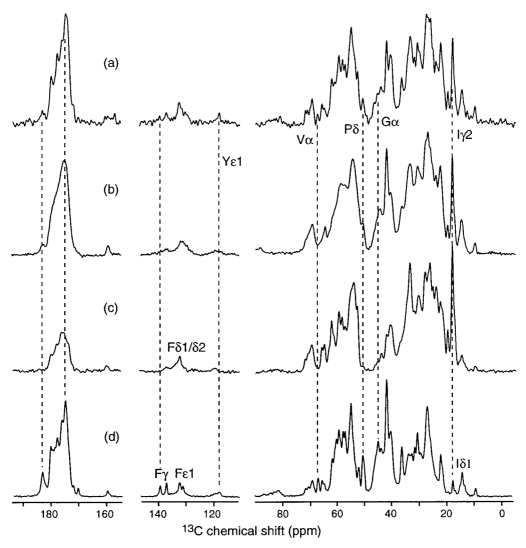


FIG. 3. Solid-state ¹³C CPMAS spectra of differently labeled ubiquitin samples. (a) Unlabeled ubiquitin, 35 mg, 4096 scans. (b) [U-¹³C₆]glucose, ¹⁵N-labeled ubiquitin, 10 mg, 64 scans. (c) [1-¹³C]glucose, ¹⁵N-labeled ubiquitin, 15 mg, 128 scans. (d) [2-¹³C]glycerol, ¹⁵N-labeled, 20 mg, 128 scans. The different intensity distributions of spectra (c) and (d) compared to (a) and (b) reflect the selectivity of labeling. Assignments for several resolved signals are indicated to illustrate the effects of selective labeling. Spectrum (b), of the uniformly labeled sample, has the lowest resolution due to the ¹³C-¹³C scalar couplings and the residual dipolar interaction.

of the amino acids from the citric-acid cycle can be avoided by supplementing the minimal medium with unlabeled intermediates of the citric-acid cycle or the unlabeled amino acid products themselves (32).

The selective and extensive ¹³C labeling approach (22, 23) differs from other selective labeling methods that require bacterial auxotrophs (33–35) and from random fractional labeling (36, 37). It is easy to implement with the standard protein expression protocol and is very versatile, since specifically ¹³C-labeled glucose, glycerol, and pyruvate are commercially available in many forms. The feasibility and versatility of this labeling approach, together with the resulting potential efficiency in the NMR structure determination, compensate for the reduced control in the placement of the isotopic labels compared to site-specific labeling methods.

2. One-Dimensional ¹³C NMR Spectra of Selectively Labeled Ubiquitin

To demonstrate the effects of S&E ¹³C labeling on the resolution of the solid-state NMR spectra and to estimate the labeling levels, we acquired 1D ¹³C CPMAS spectra of four different ubiquitin samples. The spectrum of the unlabeled ubiquitin (Fig. 3a) serves as a reference to those of the labeled samples in three ways. First, its intensity distribution is a fingerprint of the complete carbon skeleton of the protein, since all carbons in the molecule contribute equally at the natural abundance (1.1%) level. Deviations from this total intensity distribution would thus reflect the preferential labeling of certain carbon sites. Second, the homogeneous linewidth of the spectrum is minimal at the natural abundance, since the effects

of the ¹³C-¹³C scalar couplings and residual dipolar couplings are absent. Third, the signal-to-noise ratio of the spectrum provides a benchmark against which the labeling levels of the [1-¹³C]glucose and [2-¹³C]glycerol schemes can be estimated.

For U-¹³C, ¹⁵N-labeled ubiquitin, the spectrum (Fig. 3b) exhibits significant line broadening, as expected when residual ¹³C-¹³C dipolar and scalar couplings are present. This dipolar-and scalar-coupling induced line broadening was also reported recently for crystalline organic compounds (*14*, *38*). It was found that to remove the effects of these couplings, very high spinning speeds (>25 kHz) and proton decoupling fields (~200 kHz) needed to be employed. These experimental conditions are currently only feasible on the smallest MAS rotors, which have limited sample volumes and thus reduced experimental sensitivity. In addition, such high spinning speeds make rotor-synchronized NMR pulse sequences difficult to implement.

Under [1-13C]glucose labeling, the ubiquitin spectrum (Fig. 3c) exhibits reduced intensities than the reference spectrum in the C α region between 50 and 65 ppm and at ~40 ppm. In comparison, the [2-13C]glycerol labeling scheme yields low intensities in the side chain region between 15 and 35 ppm (Fig. 3d). These different intensity distributions reflect the selectivity of labeling. For example, the Gly and Val 13 C $^{\alpha}$ peaks are conspicuously missing in the ¹³C1-ubiquitin spectrum, while the Ile 13C² carbons are mostly unlabeled in ¹³C2-ubiquitin. (These resonances are assigned based on the solution NMR data (27) and are confirmed by the 2D ¹⁵N–¹³C spectra shown later.) In the carbonyl region, 13C2-ubiquitin displays substantially higher intensities, suggesting that the carbonyl carbons are labeled at higher levels than in ¹³C1ubiquitin. In general, based on the relative intensities of the ¹³C1- and ¹³C2-ubiquitin spectra to the reference spectrum (Fig. 3a), and taking into account the sample amounts and signal-averaging times, we find approximately a factor of 2 enhancement in the labeling levels of the [2-13C]glycerol scheme (50–90%) over the $[1-^{13}C]$ glucose scheme (20–45%). This is consistent with the predictions based on the amino acid biosynthetic pathways.

Both selectively labeled ubiquitin samples exhibit sharper resonances than the uniformly labeled sample. The linewidths in spectra (c) and (d) range from 0.8 ppm for the unprotonated carbonyl carbons and the mobile methyl carbons, to $\sim\!1.3$ ppm for the CH and CH $_2$ groups. In contrast, the fully labeled sample under the same experimental conditions exhibits linewidths of $\sim\!1$ ppm for CH $_3$ groups and $>\!2$ ppm for CH $_2$ carbons. These linewidth differences confirm the importance of reducing directly bonded $^{13}{\rm C}$ spin pairs for obtaining highly resolved solid-state NMR spectra.

3. Site-Specific Determination of Multiple φ Torsion Angles by HNCH Correlation

The high spectral sensitivity and resolution achieved with S&E ¹³C labeling makes possible a variety of solid-state 2D

correlation techniques for measuring structural constraints such as torsion angles (24, 28, 30, 31, 39, 40) and for sequential resonance assignment (41–43). We first demonstrate the feasibility of residue-specific torsion-angle determination on such a selectively labeled protein by measuring backbone ϕ angles in ¹³C2-ubiquitin. The recently introduced HNCH technique correlates the N–H and the C^{α} –H^{α} dipolar coupling tensors (24) in order to extract the mutual orientation of the two bonds, which reflects the ϕ torsion angle about the N–C^{α} bond. The ϕ -dependent dipolar patterns are manifested in the ω_1 dimension and separated in the ω_2 dimension according to the C^{α} isotropic chemical shifts of the residues. Due to this isotropic–anisotropic correlation, the HNCH technique is capable of extracting multiple ϕ torsion angles simultaneously, provided that the ¹³C^{α} chemical shift dispersion is sufficient.

The HNCH spectrum of ¹³C2-ubiquitin is displayed in Fig. 4. The projection onto the ω_2 dimension and the ω_1 cross sections of four selected sites are shown. In the ω_2 dimension, only carbons directly bonded to nitrogen spins, i.e., C^{α} and CO, are observed, since a short ¹⁵N-¹³C double- and zero-quantum excitation time was used (0.47 ms). In the ω_1 dimension, only the intensity envelopes connecting the tips of the sidebands are plotted in order to emphasize the ϕ -angle-induced differences in the intensity distribution. The spectra consist of 14 sidebands and one center band, as determined by the number of t_1 intervals measured. The dipolar cross section of Val26 C^{α} (67 ppm) displays a dip in the central intensity and is best-fit with $\phi = -60^{\circ}$ (Fig. 4b). This corresponds to the α -helix conformation, which is in agreement with the crystal structure (17). The dipolar cross section of Thr12 and Thr14 C^{α} (62 ppm) is dominated by a sharp intensity maximum at the center, which is the signature for the β -sheet conformation (Fig. 4c). The Ala46 (53 ppm) cross section exhibits a broad high plateau in the center and corresponds to $\phi = +20^{\circ}$ or $+100^{\circ}$, both suggesting a β -turn structure (Fig. 4d). This is also corroborated by the crystal structure (17). Even in the absence of the crystal structure, this result, combined with the fact that Gly47 is the neighboring residue, would strongly suggest a type I'-III' β -turn (44).

The experimental spectra are each best fit (Figs. 4b–4d, right) with two ϕ -torsion angles because the HNCH technique has an intrinsic double degeneracy (24). However, one of these two angular values can often be ruled out based on its rare occurrence in nature (45). The best-fit simulations are found at the minimal root-mean-square deviations (rmsd) between the normalized experimental spectrum $S_{i, \exp}$ and the normalized simulated ones $S_i(\phi)$. The rmsd curve is calculated according to

$$rmsd(\phi) = \left\{ \frac{1}{N} \sum_{i=1}^{N} [S_{i,exp} - S_i(\phi)]^2 \right\}^{1/2},$$
 [1]

where the summation i is carried out over the central six

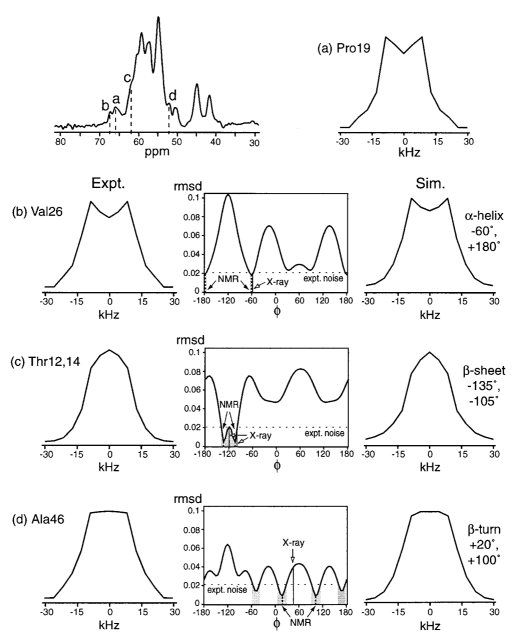


FIG. 4. HNCH spectrum of 13 C2-ubiquitin for determining the ϕ torsion angles. The experimental dipolar cross sections (ω_1 dimension) are the C^{α} of (a) Pro19, (b) Val26, (c) Thr12 and Thr14, and (d) Ala46. The assignment is obtained from solution NMR data (27) and is confirmed by the solid-state 15 N- 13 C 2D correlation spectra. The root-mean-squared deviations (rmsd) between the experiment and simulation are displayed in the middle column of (b-d). The best-fit torsion angles (dashed lines) are obtained at the minimum rmsd value. The experimental uncertainties are indicated by the shaded areas. The crystal structure results are indicated by the solid lines. For cross section (c), two solid lines are shown that correspond to Thr12 ($\phi = -120^{\circ}$) and Thr14 ($\phi = -101^{\circ}$). The best-fit simulations are shown in the right column of (b-d). The effective C-H dipolar coupling can be extracted from the Pro19 slice (a) due to its lack of the amide proton. The sample was spun at 4209 Hz. A 15 N- 13 C coherence transfer time of 475 μ s was used to select carbons directly bonded to the amide 15 N. The spectrum was acquired in 14 h.

spinning sidebands and the center band in each spectrum (46). The outer eight side bands were excluded in the rmsd calculation since their intensities are low and relatively similar for various ϕ angles. The global minima of each rmsd curve correspond to the best-fit torsion angles. A pair of minima is present for each rmsd curve, which reflects the double degeneracy of the HNCH technique. For the α -helix (b) and β -turn

(d) conformations, an additional pair of angles exhibit nearminimum rmsd values. This results from an accidental, near fourfold degeneracy in these angular regions, which has been observed in 2D (ϕ, ϕ) rmsd plots previously (46).

To assess the angular resolution of the technique, we measured the experimental noise and used its intersection with the rmsd curve to define an angular uncertainty (shaded areas).

Based on this criterion, the Val26 spectrum has the highest angular resolution, while the Ala46 result is the least precise. The accuracy of the NMR measurement can also be assessed, by comparing the NMR-derived torsion angle values (dashed lines) with the crystal structure (solid lines). We find relatively high accuracy for the α -helical and β -sheet conformations, but low accuracy for the positively valued, β -turn conformation. This may partly result from the fact that the C^{α} signal of Ala46 is not sufficiently resolved from other sites. It is also possible that the flexibility of the β -loop makes the NMR or the X-ray crystal structure an average of a number of conformations.

An internal check for the precision of the HNCH experiment is provided by the dipolar cross sections of the chemically unique amino acid, proline. Due to the lack of the amide proton in proline, its HNCH cross section exhibits a narrow envelope (Fig. 4a) that reflects the effective C^α -H $^\alpha$ dipolar coupling scaled by the homonuclear decoupling sequence. Knowledge of this effective C-H coupling makes the HNCH simulation independent of any adjustable parameter other than the torsion angle ϕ .

The advantage of the selective and extensive ¹³C labeling scheme compared to uniform ¹³C labeling lies in the fact that the uniformly labeled sample has practically no single-site resolution in the C^α region to yield any useful HNCH spectrum. Although higher spinning speeds could improve the chemical shift resolution, they are incompatible with the HNCH sequence, which requires a sufficient number of t_1 points as defined by the MREV-8 cycle time to be measured within each rotor period. This slow-spinning requirement is the main limitation of the torsion-angle technique, because even with the enhanced resolution afforded by ¹³C selective labeling, the ω_2 dimension still does not resolve all C^{α} resonances. To extract more torsion angles, it is necessary to extend the 2D HNCH technique into a third dimension, where ¹⁵N chemical shift may be employed to further disperse the resonances. This 3D experiment should not entail prohibitively long signalaveraging time, since the dipolar dimension of the experiment, incremented in a constant-time fashion (24), requires only a small number of points to be measured.

4. Resonance Assignment by ¹⁵N-¹³C Correlation

The structure elucidation of extensively labeled proteins by NMR requires the assignment of the measured structural constraints to the protein sequence (47). So far, solid-state NMR techniques for resonance assignment have been demonstrated mainly on small molecules such as amino acids and oligopeptides (42, 48–51), and few solid-state NMR studies of resonance assignment for proteins have been shown (41, 43, 52). The S&E ¹³C labeling approach facilitates resonance assignment by enhancing the spectral resolution. As a first demonstration of this feasibility, we carried out a 2D ¹⁵N–¹³C correlation experiment (42) where the coherence is transferred between ¹³C and ¹⁵N via the recoupled dipolar interaction (53). The spectra yield sequence-specific assignment based on $-N_i$ —

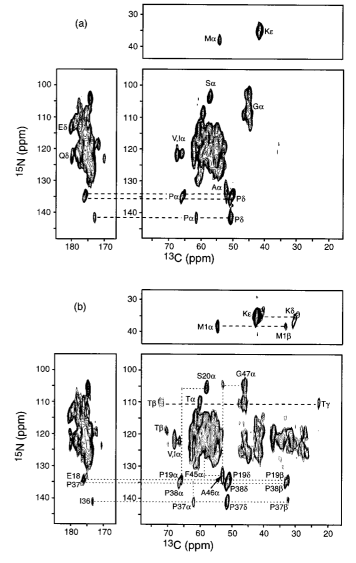


FIG. 5. ¹⁵N-¹³C 2D correlation spectrum of ¹³C2-ubiquitin with a ¹⁵N-¹³C mixing time of (a) 0.6 ms and (b) 2.3 ms. Several dozens of resonances are either partially or completely resolved in each spectrum. Amino-acid type assignment and partial sequential assignment were achieved based on the connectivity patterns, the characteristic chemical shifts, and the labeling patterns derived from Fig. 2 and Tables 1 and 2. The spectra were each acquired in 13 h.

 $C^{\alpha}_{i}-N_{i+1}-C^{\alpha}_{i+1}-$ and $-N_{i}-CO_{i}-N_{i+1}-CO_{i+1}-$ connectivities. Due to the distance-dependent nature of this coherence transfer mechanism, one can first identify the N-C^{α} and $CO_{i-1}-N_{i}$ cross peaks by using a short $^{15}N-^{13}C$ mixing time, then identify two- and three-bond $^{15}N-^{13}C$ spin pairs such as the intraresidue N-C^{α} and the interresidue $C^{\alpha}_{i-1}-N_{i}$ peaks using a longer mixing time (42).

We applied this ¹⁵N-¹³C correlation technique to ¹³C2-ubiquitin. Two ¹⁵N-¹³C spectra, acquired with a ¹⁵N-¹³C mixing time of 0.6 and 2.3 ms, are displayed (Fig. 5). Each spectrum yields several dozens of resonances that are either partially or completely resolved. Based on the chemical shifts, the connec-

tivity patterns, the labeling pattern specific to this sample, and the time-dependence of the resonance signals, we can identify several sequential connectivities. For example, in the spectrum acquired with the short mixing time (a), the six most downfield ¹⁵N resonances can be assigned to prolines due to their characteristic chemical shifts. Among these peaks, the signals at 13 C chemical shifts of ~ 50 ppm can be assigned to proline C^{δ} , while those at \sim 62 and \sim 67 ppm are proline C^{α} . Both C^{δ} and C^{α} are labeled by the [2- 13 C]glycerol scheme and are directly bonded to the amide ¹⁵N, thus they are observed at the short mixing time. At this point, only type assignment is possible. When the longer ¹⁵N-¹³C mixing time is used, we observe a weak but clear cross peak between two of the three proline C^{α} resonances. This unambiguously assigns the two peaks to the two consecutive prolines in the ubiquitin sequence, Pro37 and Pro38. Thus the remaining proline signals must belong to Pro19. One can continue along the connectivity pattern to identify the cross peak between Pro37 and Ile36, as well as the Pro19-Ser20 cross peak. Another observed sequential connectivity is Gly47-Ala46-Phe45 (43). In addition to interresidue assignment, the long mixing-time experiment also allows intraresidue assignment to be made to some extent, since the sidechain resonances appear in the same ¹⁵N cross section as the primary C^{α} signal but have more upfield ¹³C chemical shifts. One example of the intraresidue assignment is provided by the prolines, whose C^{β} site, two bonds from the amide ¹⁵N, is clearly shown in Fig. 5b. Similarly, the Thr C^{β} 's and C^{γ} 's can also be assigned, since Thr C^{β} 's have uniquely downfield chemical shifts. These assignments were made based on the solid-state spectra alone, without utilizing solution NMR chemical shift information. The latter, however, is useful as an eventual measure of the uniqueness of the solid-state assignment given the achievable linewidths (41). In these spectra, the limiting values of the full width at half maximum are ~ 0.8 ppm for ¹³C and ~2.5 ppm for ¹⁵N. This resolution is quite reasonable for an unoriented powder sample and is reflected by the partial separation of the six glycine resonances in the

Although the selective ¹³C labeling makes complete assignment from a single sample not possible, with two samples labeled in a complementary fashion, one can retrieve the missing information to some extent. For example, the ¹³C1-ubiquitin spectrum acquired at the long ¹⁵N–¹³C mixing time (Fig. 6b) exhibits the C^{β} signals of Ala, Ser, and Pro. Among these, Ser20 C^{β} can be assigned based on its cross peak with Pro19 C^{α} , and Ala46 C^{β} is identified by its cross peak with Gly47 C^{α} . Thr9 can be assigned by its cross peak with Leu8 C^{α} . Similarly, the sequential connectivities Thr66–Ser65 and Pro19–Glu18 can be identified.

To assign the 13 C and 15 N spectra of a solid protein more completely, additional correlation experiments that fully exploit the labeling patterns are required. For example, 3D experiments that correlate the C^{α} and sidechain carbon chemical shifts with the backbone 15 N are useful for more complete amino acid type assignment of the resonances. Homonuclear

¹³C correlation schemes will also be useful for type assignment. Experiments along these lines will be detailed in a future paper (54). On the other hand, variations of the selective and extensive labeling approach can be exploited to label carbons that are either unlabeled or labeled at low levels by the [1-¹³C]glucose and [2-¹³C]glycerol labeling schemes. This will be important to achieve complete resonance assignment of the proteins.

CONCLUSIONS

We have shown that selective and extensive ¹³C labeling in conjunction with high-resolution solid-state NMR is a useful approach to the efficient structure investigation of solid proteins. The selective ¹³C labeling primarily refers to the fact that the statistical probability of directly bonded ¹³C–¹³C spin pairs is close to 0% while the probability of one of the two carbons being ¹³C is significant. This is achieved by using specifically labeled glucose or glycerol as the sole carbon source in the bacterial growth media. We found, both theoretically and experimentally, that the selectivity of labeling is most prominent for the 10 amino acids produced from the linear part of the biosynthetic pathways. In these amino acids, many carbon pairs with 45-90% and 1.1% (i.e., natural abundance) labeling levels are found. In comparison, in amino acids derived from the more complex citric-acid cycle, many adjacent carbons have 50% or lower labeling levels for both sites, thus the spin-pair probabilities are below 25%.

The selective and extensive ¹³C labeling approach provides a general method of spectral narrowing and editing, which facilitates resonance assignment in solid proteins. The enhanced spectral resolution makes many solid-state NMR techniques practical for measuring multiple structural constraints per experiment and per sample. We showed here that constraints on the backbone conformation of such a labeled protein can be obtained by measuring the ϕ torsion angles using the HNCH technique. The angular resolution and accuracy of the measurement were analyzed. We also showed that the sequence-specific assignment of these structural constraints is partially feasible by 2D ¹⁵N-¹³C correlation. The extent of resonance assignment is still limited at this point by the spectral resolution due to the lack of ultrafast spinning and ultrahigh magnetic field. Combining the advance of these technologies, various heteronuclear and homonuclear correlation techniques, and 3D spectroscopy, more complete resonance assignment of proteins in the solid state is well foreseeable.

MATERIALS AND METHODS

The three ¹³C, ¹⁵N-labeled ubiquitin samples used in this work were custom-synthesized by VLI-Research (Malvern, PA). The human ubiquitin gene was expressed in *Escherichia coli*. Cells were grown in M9 minimal medium, with the sole nitrogen source being ¹⁵N-labeled ammonium sulfate, and the sole carbon source being one of the following three com-

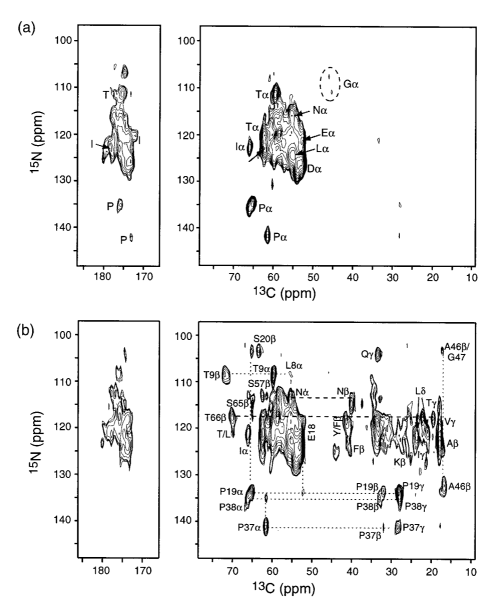


FIG. 6. 15 N $^{-13}$ C 2D correlation spectrum of 13 C1-ubiquitin with a 15 N $^{-13}$ C mixing time of (a) 0.6 ms and (b) 2.3 ms. Dashed circle indicates the weak glycine C^a signals, whose presence indicates low-level scrambling in the 13 C labeling scheme.

pounds: $[1^{-13}C]$ glucose, $[2^{-13}C]$ glycerol, and $[U^{-13}C_6]$ glucose (Cambridge Isotope Labs, Inc.). For a typical ubiquitin expression experiment, a 5-liter culture was induced and the solid paste was purified by column chromatography to homogeneity. The purity of the final products was analyzed by gel electrophoresis and amino acid composition analysis to be >99%. The unlabeled ubiquitin sample was purchased from Sigma and used directly without purification. All samples were hydrated to $\sim 30\%$ (w/w) prior to the NMR experiments. Hydration was found to be important for narrowing the spectral lines, mainly by decreasing the conformational heterogeneity present in lyophilized samples (55-57).

The solution-state ¹³C-¹H 2D correlation experiments for verifying the labeling schemes were carried out on a Bruker DPX-300 spectrometer using a 5-mm inverse probe. A refo-

cused INEPT pulse sequence was used (58). The samples consist of 0.7 mM ubiquitin dissolved in a 90% $\rm H_2O/10\%~D_2O$ sodium acetate buffer at pH 4.5.

All solid-state NMR experiments were performed on a Bruker DSX-300 spectrometer using a 4-mm triple-resonance MAS probe. Typical radiofrequency field strengths were 120 kHz for 1 H decoupling, 65 kHz for 13 C, and 50 kHz for 15 N. Cross polarization from 1 H to 13 C was achieved with a linearly ramped (59, 60) RF field on the 13 C channel and with a contact time of 0.5 ms. The FIDs were detected under 1 H TPPM decoupling (61). The evolution period of the ϕ torsion experiment employed a MREV-8 pulse train for homonuclear decoupling. Otherwise, continuous-wave decoupling was applied on the 1 H channel. Typical recycle delays were 3 s.

The 1D ¹³C CPMAS experiments and the 2D ¹⁵N-¹³C cor-

relation experiments were performed under 7 kHz of magicangle spinning. The spinning speed was controlled to ±2 Hz by a Bruker MAS controller. The ¹⁵N-¹³C correlation experiments were carried out using the pulse sequence in reference (42), except that the ¹⁵N-¹³C REDOR mixing period consisted of a single ¹³C 180° pulse and multiple ¹⁵N 180° pulses (62, 63). A total of 4 and 16 rotor periods were used to excite the double- and zero-quantum ¹⁵N-¹³C coherence of one-bond and multiple-bond ¹⁵N-¹³C pairs, respectively.

The ϕ torsion angle experiment was performed using the basic pulse sequence in Ref. (24), except that the apparent N-H and C^{α} -H $^{\alpha}$ dipolar couplings were doubled relative to the spinning speed to achieve high angular resolution. The doubling was achieved by moving the 180° pulses on the ¹³C and ^{15}N channels as a function of t_1 during the constant-time evolution period (46). Meanwhile, MREV-8 homonuclear decoupling (64) was applied for the entire rotor period on the ¹H channel. The MREV-8 90° pulse length was 2.2 µs and the short-window delay was 0.9 μ s. They were optimized by maximizing the sideband intensities of the leucine 13 C $^{\alpha}$ signal detected under MREV-8 decoupling. At a spinning speed of 4209 Hz, 16 t_1 points were collected at an increment of 14.85 μs, so that the maximum evolution time corresponded to one rotor period. Before the t_2 period, a TOSS sequence (65) was applied to remove the carbonyl sidebands in the ω_2 dimension, since they overlap with some of the aliphatic resonances.

To eliminate uncertainties in the MREV-8 scaling factor, the magnitude of the one-bond C–H dipolar coupling was directly measured using a DIPSHIFT pulse sequence (66-68). The result was further corroborated by the proline C^{α} cross section in the ϕ -angle spectra. The one-bond C–H couplings were thus found to be 10 ± 1 kHz.

The ϕ -angle spectra were simulated using a FORTRAN program described previously (24). A long-range C^{α} -H(N) dipolar coupling of 1.2 kHz was taken into account in the simulation. The theoretical ϕ -angle spectra were calculated at 5° increments over the entire 360° range. The root-mean-square deviations between the experiment and the simulations were calculated in MATLAB as a function of the ϕ -angle.

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