Direct NMR observation of a substrate protein bound to the chaperonin GroEL

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The reaction cycle and the major structural states of the molecular chaperone GroEL and its cochaperone, GroES, are well characterized. In contrast, very little is known about the nonnative states of the substrate polypeptide acted on by the chaperonin machinery. In this study, we investigated the substrate protein human dihydrofolate reductase (hDHFR) while bound to GroEL or to a single-ring analog, SR1, by NMR spectroscopy in solution under conditions where hDHFR was efficiently recovered as a folded, enzymatically active protein from the stable complexes upon addition of ATP and GroES. By using the NMR techniques of transverse relaxation-optimized spectroscopy (TROSY), cross-correlated relaxation-induced polarization transfer (CRIPT), and cross-correlated relaxation-enhanced polarization transfer (CRINEPT), bound hDHFR could be observed directly. Measurements of the buildup of hDHFR NMR signals by different magnetic environments intermediate in polarity between native and fully unfolded states, suggesting an unstable tertiary structure (21). Subsequent hydrogen–deuterium exchange experiments revealed that there are only low degrees of exchange protection, suggesting the absence of stable regular secondary structures (17, 22–25). Crystallographic observations on substrates bound to GroEL have been made in two cases but involved only short peptides associating with single apical helices (26, 27). The peptides were bound in an extended state in the hydrophobic groove between two apical α-helices, but it is unclear whether such well-ordered structures inform about the behavior of full-length polypeptide substrates. Indeed, it has been suggested that these structures may more closely mimic the binding to the apical surface of the so-called “mobile loop” of the cochaperonin GroES (28). Thus, it would be desirable to directly inspect an intact nonnative polypeptide while bound to GroEL. This inspection has become possible through the development of NMR techniques that can observe very large molecules in solution (29, 30). In particular, such techniques have recently been applied to the GroEL system, examining binding of the C15N-labeled and perdeuterated 70-kDa cochaperonin, GroES, to the unlabeled 800-kDa chaperonin, GroEL. Nearly all of the 94 amide protons of the GroES subunits were observed both in the free cochaperonin and in the 870-kDa complex, and chemical shift changes were detected upon association of GroES with GroEL, localizing to the GroES mobile loop region (31). We now report NMR observations of an isotope-labeled substrate polypeptide, human dihydrofolate reductase (hDHFR), while bound to unlabeled or C15N-depleted GroEL.

Materials and Methods

hDHFR Expression. For uniform C15N-labeling, the E. coli strain BL21(DE3) harboring the T7 expression plasmid was grown on hydrogen-exchange studies (17), but how is the unfolded state obtained? Is there active unfolding mediated by multivalent binding (18)? Or is the apparent unfolding action a passive result of thermodynamic partitioning, in which GroEL binds less-folded conformers with greater affinity, shifting the ensemble toward a less-folded state (19, 20)? As a basis for distinguishing between different possible mechanisms, at least the stable GroEL-bound “end-state” of a nonnative substrate bound in an open GroEL ring must be characterized.

Early studies revealed that GroEL-bound polypeptides were exquisitely protease susceptible, with their tryptophans in environments intermediate in polarity between native and fully unfolded states, suggesting an unstable tertiary structure (21).

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C haperone proteins are involved in nearly every cellular process in which proteins must become unfolded or refolded, including transport across membranes, assembly and disassembly of macromolecular complexes, and de novo folding of newly synthesized cytosolic proteins. Chaperones of the Hsp60 class, also named chaperonins, are of special interest in this regard, because folding takes place within a central, closed cavity. The chaperonin architecture consists of two homo-oligomeric rings stacked back to back. The polypeptide substrate binds in the open end of one of these rings and is incorporated upon subsequent binding of the cochaperonin partner protein.

The essential cellular action of chaperonin-mediated folding involves cycles of polypeptide binding and ATP/GroES-driven release, with only a small percentage of input molecules reaching the native state in any given cycle (1–5). With each round of binding, it appears that the same ensemble of chaperonin-associated nonnative conformers becomes stably associated with the chaperonin, giving the bound molecules essentially the same chance at reaching the native state with each round of release (1, 6). In the case of the Escherichia coli chaperonin GroEL, binding is mediated through a hydrophobic lining of its central cavity (7), which forms multivalent contacts (8) with exposed hydrophobic surfaces of the nonnative substrate protein (9–14). Such interaction serves to prevent substrate protein from irreversible misfolding and aggregation (15, 16).

Although the overall action of polypeptide binding by GroEL has been well described, possible correlations between the binding mode and the substrate conformation have not yet been rationalized on the molecular level. GroEL can catalyze complete unfolding of a small protein, a result obtained from
D₂O-based M9 medium by using [¹⁵N]ammonium chloride as the sole nitrogen source. Samples containing an overall deuteration level of ~85% were prepared by using unlabeled glucose as the sole carbon source, whereas samples containing >98% deuteration were prepared by using deuterated acetate as the sole carbon source (31–35), requiring preconditioning of cells, first to medium containing D₂O with unlabeled glucose, followed by the full labeling medium.

Specific labeling of the leucine residues with [¹⁵N] was carried out on a transaminase-deficient strain (35). Cells were grown to mid-log in LB, then in M9 with either H₂O (~80% final deuteration) or D₂O (~90% final deuteration), then supplemented with 1 g/liter deuterated algae extract (CELTOX-d powder, Spectra Stable Isotopes, Columbia, MD), 75 mg/liter [¹³C,¹⁵N]-leucine, and 0.5 mM isopropl β-D-thiogalactoside. Cells were harvested after 3–5 h. hDHFR was purified by using methotrexate-agarose affinity chromatography (24).

hDHFR Sample Preparation. Reference samples of folded hDHFR were diaalyzed against binding buffer containing 250 μM dihydrofolate and 5 mM ATP, and 5% D₂O was added before the NMR measurements. Denatured hDHFR was prepared by diluting the denatured hDHFR ~10-fold into 6.6 M guanidine hydrochloride and 50 mM Tris (pH 7.5), followed by PD10 chromatography. Samples were concentrated to ~10 mg/ml, supplemented with 1–5 mM DTT, and frozen at ~80°C until use for complex formation with GroEL or single ring variant 1 of GroES (SR1). Denatured hDHFR reference NMR samples were prepared by dilution of the stock protein solutions into the same buffer.

Chaperonins, hDHFR–GroEL Complexes, and Refolding. GroES and SR1 were expressed as described in refs. 31 and 36. Complexes of either GroEL or SR1 with hDHFR were prepared by diluting 10 μl of 0.5 mM GuHCl-unfolded hDHFR 50-fold into binding buffer containing 15 μM chaperonin protein at 23°C. To prepare a sample, a number of such mixtures were pooled, centrifuged at 18,000 × g for 5 min to remove aggregated unbound hDHFR, then exchanged to remove residual GuHCl and concentrated to 0.3 ml and supplemented with 5% D₂O. The final protein concentration was typically 60–80 mg/ml. Refolding was accomplished by addition of dihydrofolate (500 μM final), GroES (1.5:1 GroES to SR1 or GroEL), and ATP (5 mM final). After 10 min at 16°C, the sample was centrifuged, and NMR was performed.

Assay for hDHFR Enzymatic Activity. hDHFR subjected to refolding exactly as just described also was assayed for recovery of enzymatic activity (24), reflecting production of the native state. Recovery of activity was observed with single-exponential kinetics (k = 0.31 min⁻¹), reaching nearly complete reactivation by 10 min.

Detection of Internal Dynamics of hDHFR Bound to SR1 by Using NMR. Many different experiments have been proposed to measure internal dynamics in proteins (37–41). However, the sensitivity of these experiments is too low for large macromolecular structures of several hundred kilodaltons in size, so we used a previously undescribed approach based on comparison of the efficiencies of [¹⁴N] magnetization transfer in backbone [¹⁴N]-H moieties to generate [¹H] magnetization in antiphase to [¹⁵N] when using either cross-correlated relaxation-induced polarization transfer (CRIPT) or insensitive nuclei enhanced by polarization transfer (INEPT) coherent transfer (42).

The transfer of [¹H] magnetization into [¹H] magnetization in antiphase to [¹⁵N], Sᵣᵣ, by cross-correlated relaxation between dipole–dipole (DD) coupling and chemical shift anisotropy (CSA) interactions by means of CRIPT or by scalar coupling with INEPT is described by (42, 43)

\[ S_r(T) = A_s(T) \exp(-R_cT), \quad v \in \{\text{INEPT, CRIPT}\}, \]  \[ A_{\text{INEPT}}(T) = \sin(\pi T), \quad A_{\text{CRIP}}(T) = \sinh(R_c T). \]

\[ R_I \] is the transverse relaxation rate of [¹H], T is the transfer time, and \( R_C \) is the relaxation rate due to cross-correlation between [¹H] CSA and [¹⁵N] DD coupling.

\( R_I \) and \( R_C \) are given by Eqs. 3 and 4 for isotropic rotational tumbling of rigid, high-molecular-weight systems at high magnetic field (44)

\[ R_I = \frac{2}{5} \left( \frac{\beta \gamma_H \gamma_N}{r_{SH}} \right)^2 + \frac{2}{9} \left( \gamma_H \Delta \sigma_B B_0 \right)^2 T_1 + \frac{1}{2 T_2(N)} \]  \[ R_C = \frac{4}{15} \left( \frac{\beta \gamma_H \gamma_N}{r_{SH}} \right) \left( \gamma_H \Delta \sigma_B B_0 P_2(\theta_{\text{DD-CSA}}) \right) T_2. \]

\( \tau \) is the isotropic rotational correlation time of an equivalent sphere representing the molecule of interest (41), \( r_{SH} \) is the distance between the two nuclei, \( \Delta \sigma_B \) is the CSA of [¹⁴N], \( B_0 \) is the static magnetic field, and \( \gamma_H \) and \( \gamma_N \) are the gyromagnetic ratios of [¹H] and [¹⁵N], respectively. The second-order Legendre polynomial, \( P_2(\theta_{\text{DD-CSA}}) \), accounts for deviations of the angle \( \theta_{\text{DD-CSA}} \) between the DD vector and the principal axis of the CSA tensor from 0°. The term \( 1/T_2(N) \) is the longitudinal relaxation rate of [¹⁵N], and \( 1/T_2(S) \) is the contribution to the transverse relaxation rate of [¹⁴N] due to DD coupling with remote protons H at distances \( r_{HH} \). Assuming for large structures that \( 1/T_2(S) = 2(\gamma_H^2/2 I_{HH})^2 \tau_H \gg 1/T_2(N) \), the optimal CRIPT transfer period, \( T_C \), becomes

\[ T_C = (1/R_C) \arctan(h R_C/R_I), \]

and is inversely proportional to \( \tau \) (Eq. 4). Measurement of CRIPT build-up curves for the determination of \( T_C \) therefore enables an estimate of the size of the molecular particle studied (42), assuming that internal motions are sufficiently limited so as to have at most small effects on \( T_C \).

Usually, three distinct regimes are distinguished when assessing the influence of internal motions (characterized by the correlation time \( \tau_m \)) on the overall relaxation rates (45). For slow internal motions with \( \tau_m \gg \tau_c \) (\( \tau_m \) in the microsecond to millisecond time range), the most evident effect is resonance line broadening (39, 41, 46). For high-molecular-weight systems, such additional line broadening may be difficult to quantitate, because it is added to the inherently large line width because of transverse relaxation (30, 47, 48). Internal motions with correlation times \( \tau_c \) comparable with \( \tau_c \), with \( 1 \leq \tau_m/\tau_c \ll 10 \) (\( \tau_m \) in the nanosecond to microsecond time range) affect directly the effective correlation time for transverse relaxation (41). Finally, very rapid internal motions, with \( \tau_m \ll \tau_c \) (\( \tau_m \) in the picosecond to nanosecond time range) reduce the efficiency of relaxation pathways, but the respective correlation times cannot be measured directly by NMR methods (41, 49, 50). Internal motions with \( \tau_m/\tau_c \leq 10 \) that are not correlated with the overall rotational tumbling always reduce \( R_C \) and \( R_I \) (41). Slow internal motions (\( \tau_m \gg \tau_c \)), conversely, do not influence \( R_C \) but can increase the \( R_I \) value due to chemical shift modulation (39, 41).

Here, we make use of this situation to assess the internal dynamics of substrate proteins bound to GroEL/GroES by...
estimating the effective $R_C$ and $R_I$ values by using a combination of INEPT and CRIP'T build-up curves.

For the determination of $R_C$ and $R_I$ from build-up curves, the optimal INEPT transfer delay $T_1$ and the ratio between the optimal CRIP'T and INEPT transfer efficiencies, $A_{CIP'T}$, are needed. These parameters are defined by

$$T_1 = (1/\pi J)\arctan(\pi J/R_I),$$

and

$$A_{CIP'T} = S_{CRIP'T}(T_C)/S_{INEPT}(T_1),$$

where $J = 93$ Hz is the $\text{^1H}J_{\text{^15N},\text{^1H}}$ coupling constant in the amide moiety. $R_I$ can be estimated from experimental $T_1$ values ($T_1^{\text{exp}}$) by using Eq. 6

$$R_I = \pi J/\tan(\pi J/T_1^{\text{exp}}).$$

With the experimental value for $A_{CIP'T}$, one can then estimate $R_C$ by numerical evaluation of the roots of the function $F(R_C)$

$$F(R_C) = A_{CIP'T}(R_C, T_1^{\text{exp}}) - A_{CIP'T}^{\text{exp}},$$

where $A_{CIP'T}(R_C, T_1^{\text{exp}})$ is given by

$$A_{CIP'T}(R_C, T_1^{\text{exp}}) = \sinh(R_C T_1^{\text{exp}})/(\sinh(R_C T_1^{\text{exp}}) + \sin(\pi J T_1^{\text{exp}}).$$

$T_C$ and $R_I$ are functions of $R_C$ and $T_1^{\text{exp}}$ (Eqs. 5 and 8). The roots of Eq. 9 were determined numerically by applying the Newton–Raphson method (51), using the MAPLE 9 software package (Maplesoft, Waterloo, ON, Canada).

**NMR Spectroscopy.** All NMR spectra were recorded on a DRX 750-MHz spectrometer (Bruker, Fällanden, Switzerland) equipped with a triple-resonance probehead and a shielded z-gradient coil. NMR spectra were collected at 25°C. Two-dimensional $[^{15}\text{N},^{1}\text{H}]$ transverse relaxation-optimized spectroscopy (TROSY), 2D $[^{15}\text{N},^{1}\text{H}]$-cross-correlated relaxation-enhanced polarization transfer (CRINEPT)–heteronuclear multiple-quantum correlation (HMQC)–[^{1}\text{H}]TROSY, and 2D $[^{15}\text{N},^{1}\text{H}]$-CRIP'T–TROSY experiments were recorded as described in refs. 31 and 44. Details of the parameter settings are given in the figure legends. The spectra were processed with the program PROSA (52) and analyzed with the program XEASY (53).

**Results**

**NMR Observation of Substrate Binding to Chaperonins.** Our experimental setup (Fig. 1) includes that hDHFR was assessed in the folded state by using NMR spectroscopy. The 2D $[^{15}\text{N},^{1}\text{H}]$-TROSY spectrum of [u-$^{15}\text{N},\text{u}~80\%$ $^{2}\text{H}]$hDHFR (Fig. 2a) contains ~190 resonances from backbone amide groups, which show large chemical shift dispersion in the proton and nitrogen dimensions, as is characteristic of a folded protein. There is evidence for two conformations of the protein, which differ in the occupation of the NADPH binding site of hDHFR (54–56). The $^{15}\text{N}$-labeled, deuterated protein was then denatured in 6.6 M guanidinium hydrochloride solution and then diluted 50-fold into buffer containing either of the chaperonins SR1 or GroEL. Unbound aggregated protein was removed by centrifugation. The substrate–chaperonin complex used for the NMR studies was collected from the supernatant. After the NMR measurements, hDHFR was refolded from the complex by addition of a mixture of GroES (green), ATP, and dihydrofolate (DHF). Two sets of NMR spectra that were recorded, respectively, with uniform or residue-specific $^{15}\text{N}$-labeling of hDHFR, are shown in Fig. 2, as indicated.

**NMR Characterization of Bound hDHFR.** The distribution of resonances in 2D $[^{15}\text{N},^{1}\text{H}]$-CRIP'T–HMQC–[^{1}\text{H}]TROSY spectra of the stable binary complex between unlabeled SR1 and [u-$^{15}\text{N},\text{u}~85\%$ $^{2}\text{H}]$hDHFR (Fig. 2b) has a small chemical shift dispersion, indicating that the observed species does not adopt isotope distribution or uniformly $^{14}\text{N},^{2}\text{H}$-labeled, such that they do not give rise to $^{15}\text{N}$ NMR signals, and the observed spectra arise solely from [u-$^{15}\text{N},\text{u}~80\%$ $^{2}\text{H}]$hDHFR. We obtained the highest sensitivity with the 2D $[^{15}\text{N},^{1}\text{H}]$-CRIP'T–HMQC–[^{1}\text{H}]TROSY scheme (42). The peak patterns (data not shown), with higher signal intensities in the SR1 complexes of hDHFR with GroEL and SR1 were very similar (data not shown), with higher signal intensities in the SR1 complexes. The spectra of hDHFR bound to unlabeled or $^{14}\text{N},^{2}\text{H}$-labeled SR1 were identical. Therefore, we used unlabeled SR1 for further studies. A sample of [u-$^{15}\text{N},\text{u}~85\%$ $^{2}\text{H}]$hDHFR diluted from denaturant in the absence of chaperonin yielded no NMR signal after centrifugation at 18,000 × g to remove aggregated hDHFR, ruling out the possibility that the observed signals arise from unbound hDHFR.
The same experiments as presented in either uniformly labeled with $^{15}$N or with specific $^{15}$N-labeling of the 19 Leu residues. The transfer period was 1 s.

For an assessment of whether or not the bound state observed by NMR is a true folding intermediate, we added GroES, ATP, and dihydrofolate to the hDHFR/SR1 complex (24) (Fig. 1). Such incubation produced efficient re-folding of hDHFR to an enzymatically active form within 10 min. In the 2D $[^{15}$N,$^1$H]-TROSY spectrum of the recovered substrate (Fig. 2c), the broad peaks in the random coil region of the spectrum in Fig. 2b have completely disappeared, and a widely dispersed spectrum characteristic of a globular protein is present.

To gain further insight into the dynamic properties of hDHFR bound to SR1, we recorded $^{15}$N-selected 1D $^1$H NMR spectra of $[^{15}$N,$^1$H]hDHFR bound to unlabeled SR1, after a CRIPT or an INEPT magnetization transfer step (Fig. 3a). The signal of the 1D spectrum using the INEPT element shows nearly three times higher signal intensity than that using the CRIPT element, and the resonances in both spectra have antiphase character. For a substrate rigidly bound to the 400-kDa SR1, one would expect that the CRIPT and INEPT transfer have similar transfer efficiencies at 750 MHz and that one component of the INEPT buildup curve fitted to the experimental data by using Eqs. 1 and 2. The spectra were recorded on a Bruker DRX 750 spectrometer. The number of transients per $T$ value in the CRIPT and INEPT experiments was 512.
uniformly 15N-labeled hDHFR (Fig. 2). We do not presently know why only part of the hDHFR spectrum is seen after dissociation from SR1; quite likely this result is because of the lower concentration of the refolded protein. Nonetheless, recovering a globular protein with hDHFR enzymatic activity and hDHFR-like NMR chemical shifts supports the hypothesis that the NMR-obscorable GroEL-bound hDHFR represents a bona fide folding intermediate in the chaperonin cycle. More direct evidence for this conclusion comes from the experiments with residue-selective labeling in the following section.

Studies with [15N,2H]Leu-hDHFR. The large line widths and the severe overlap of the resonances in the spectrum of the binary complex make it difficult to estimate the number of resonances observed in Fig. 2b. Assuming that all of the side-chain NH groups of Asn and Gln contribute to the peak clusters indicated by I and II in Fig. 2b, we estimate that the intensities of the other peaks in Fig. 2b correspond to ~25% of all backbone 15N-1H moieties of hDHFR. This apparent low intensity could be due either to more extensive line broadening of the backbone resonances or to their representing only a subset of signals, corresponding to discrete parts of the molecule.

For a further investigation of this issue, we chose to simplify the spectrum by labeling only the Leu residues with 15N. A 2D [15N,1H]-HSQC spectrum of the denatured [15N,1H]-Leu-u–80% 2H]hDHFR contains the expected 19 resonances (see Fig. 5, which is published as supporting information on the PNAS web site), and folded hDHFR shows the typical wide distribution of the Leu resonances (Fig. 2d). The spectrum of the complex with SR1 prepared with this material, however, contains only one continuous area of signal intensity (Fig. 2e), and a precise count of the number of cross-peaks is not possible. The NMR spectrum of the enzymatically active hDHFR recovered from the binary complex as described in the preceding section (Fig. 2f) shows the same chemical shift dispersion and closely similar peak patterns as native hDHFR (Fig. 2d), but as with uniformly 15N-labeled hDHFR (Fig. 2e), only an incomplete set of the resonance lines is seen (see the preceding section).

Discussion
The qualitative evaluation in the preceding section of solution NMR data on the complexes between hDHFR and either SR1 or GroEL indicates that the NMR-obscorable parts of hDHFR do not adopt a defined 3D structure and have high internal mobility. This conclusion is in line with previously reported small amide proton protection factors for hDHFR (6, 24) and other substrate proteins (17) in complex with GroEL. Here, we elaborate further on the analysis of the signal build-up data of Fig. 3 and on the biological implications of the present observations.

Protein Dynamics from NMR Signal Build-Up Measurements. To test the significance of the data in Fig. 3, we performed model calculations of buildup curves over a wide range of different relaxation parameters. Fig. 4 shows pairs of T1,AC/I values calculated for different Rс values by applying Eq. 10, where the thick solid lines indicate trajectories with constant Rс in the T1,AC/I plane. T1,AC/I combinations for an isolated 15N-1H two-spin system lie on the dotted curve, and those for 15N-1H moieties in perdeuterated regular antiparallel β-sheets and α-helices, respectively, lie on the broken curves. Experimental T1,AC/I combinations for the complexes of GroES with SR1 and of hDHFR with SR1 are indicated as black diamonds. The GroES/SR1 value is compatible with a β-sheet in a rigid protein structure with the correlation time for GroES/SRI of τc = 175 ns (open diamond in Fig. 4); the respective estimated Rс values (Eq. 9) are 200 and 220 Hz. In contrast, the hDHFR/SR1 complex has a much lower Rс value of ~100 Hz, which is incompatible with a rigid structure of the size of SR1. From the T1 value of 2.5 ms for hDHFR/SR1, Eq. 10 shows that the Rс value is ~350 Hz, as compared with the Rс value for the GroES/SR1 complex of ~250 Hz. The increased Rс value is an indication for slow internal motions, which also would explain the line broadening observed in the 2D [15N,1H]-HSQC spectrum.

Overall, the analysis of the CRIT and INEPT build-up curves shows that hDHFR is subject to internal mobility in the SR1 complex, which includes components with correlation times in the microsecond to millisecond time range, as well as components with frequencies in the picosecond to nanosecond time range. Such wide distribution of the frequencies of internal motions has previously been suggested for “collapsed conformations” of polypeptide chains, which would not have well-defined tertiary structure packing (57, 58). For the hDHFR/GroEL system, the interpretation of this analysis is limited, because we have not yet succeeded in determining whether or not the entire polypeptide chain of the bound hDHFR is NMR-obscorable or in identifying which parts of the bound hDHFR are observed (see Results). Quite generally, however, the approach used here opens an avenue for assessing the dynamics of distinct polypeptide segments in very large structures.

Implications for the Mechanism of GroEL Action. The NMR data presented in this work support a general model of GroEL action in which the chaperonin maintains proteins in an unfolded state while they are bound. The initial act of binding of nonnative proteins in an open ring may be associated with an unfolding action on the part of the chaperonin, but that is not resolvable
by the present analysis. In particular, in the present study, hDHFR was diluted from guanidine denaturant into an aqueous mixture containing GroEL, where early folding intermediates are formed in <20 ms and are efficiently recognized by GroEL (ref. 24 and M. Goldberg, J. Beechem, and A.L.H., unpublished data). Thereby it is difficult to assess whether partially structured states become less structured upon binding to GroEL. However, GroEL also efficiently binds later-folding intermediates of hDHFR. As yet with certainty whether apical-associated stretches of states become less structured upon binding to GroEL. However, GroEL also efficiently binds later-folding intermediates of hDHFR, which are formed many seconds or even several minutes after dilution from denaturant (24). It thus may become informative, as further understanding of the NMR-observable features of GroEL-bound hDHFR is attained, to assess whether these more structured states will produce similar or different spectroscopic behavior when they become associated with GroEL.

The apparent dynamic nature of GroEL-bound hDHFR also invites the question of whether there is ongoing binding and release of distinct segments of hDHFR from the apical domains of GroEL within the binary complex. Although we do not know as yet with certainty whether apical-associated stretches of polypeptide are NMR-observable (see Results), it is noteworthy that the observed hDHFR resonances were not visibly affected by deuteration of SR1.

In sum, here we have provided direct observations of a substrate protein while bound to the chaperonin GroEL in solution by using advanced NMR methods. The present work makes clear that we are confronting the problem of one smaller, relatively unstructured substrate polypeptide, moving with a distinct set of motional regimes inside the far larger chaperonin “container,” which presents a previously undescribed set of physical problems whose resolution will provide important biological insights. Further techniques of both biology and physics may be needed to resolve this system.

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