Long-Range-Distance NMR Effects in a Protein Labeled with a Lanthanide–DOTA Chelate

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Abstract: A two-thiol reactive lanthanide–DOTA (1,4,7,10-tetraazacyclododecane-\(N^1,N^4,N^7,N^{10}\)-tetraacetic acid) chelate, CLaNP-3 (CLaNP = caged lanthanide NMR probe), was synthesized for the rigid attachment to cysteine groups on a protein surface, and used to obtain long-range-distance information from the \({}^{15}\text{N,}^1\text{H} \) HSQC spectra of the protein–lanthanide complex. The DOTA ring exhibits several isomers that are in exchange; however, single resonances were observed for most amide groups in the protein, allowing determination of a single, apparent magnetic-susceptibility tensor. Pseudocontact shifts caused by Yb-containing CLaNP-3 were observed for atoms at 15–35 Å from the metal. By using Gd-containing CLaNP-3, relaxation effects were observed, allowing distances up to 30 Å from the paramagnetic center to be determined accurately. Similar results were obtained with a Gd–DTPA (diethylene-triaminepentaacetic acid) chelate, CLaNP-1, bound in the same bidentate manner to the protein. This study demonstrates that bidentate attachment of a paramagnetic probe enables determination of long-range distances.

Keywords: cage compounds • lanthanides • NMR spectroscopy • paramagnetic relaxation • pseudocontact shifts

Introduction

Long-range-distance information is required for solution structure determination of flexible molecules, such as nucleic acids or multidomain proteins, as well as for complexes of macromolecules.

Paramagnetic NMR effects, such as pseudocontact shifts (PCS), paramagnetic relaxation enhancement (PRE), and paramagnetic cross-correlated relaxation, are caused by the presence of unpaired electrons and are felt by the nuclear spins situated up to tens of Ångströms away. These effects are well understood and provide useful tools in structure determination.

Paramagnetic effects can be used in protein NMR analysis by addition of transition metals,[1,2] lanthanide ions,[3,4] lanthanide chelates,[5–10] stable radicals, such as 2,2,6,6-tetramethylpiperidine-1-oxyl (TEMPO),[11–16] or dioxygen,[17] to decrease the signal overlap of the resonances observed, to probe the molecular surface of these large molecules, or to align macromolecules for residual dipolar-coupling measurements.

A wealth of structural information was obtained from the paramagnetic NMR on proteins that contain a paramagnetic metal, or in which a diamagnetic metal could be substituted for a paramagnetic one.[18–34] As was first demonstrated for the cytochrome \(f\)-plastocyanin complex,[35–38] paramagnetic effects can also be used for structure determination of protein complexes.[39] Metal substitution can also be used effectively for this purpose.[39] Although there are quite a few examples for which a paramagnetic metal exists or can be engineered inside the core of a protein, there are numerous others for which this is not possible. This disadvantage can be eliminated by the construction of paramagnetic probes...
for covalent attachment to the protein. One possibility is to use N- or C-terminal fusion peptides engineered to bind a paramagnetic metal for co-expression with the protein of interest, however, this limits the choice for the location of the metal.[41–45]

Another possibility is to attach site-specific probes onto the surface of the macromolecule by chemical methods. In this way, different paramagnetic centers, such as nitroxide spin labels,[46–54] coordinated divalent metals,[55–58] and lanthanide chelates,[59–64] were created and can be used for structure refinement. Lanthanides are strong paramagnetic agents and, therefore, particularly useful in providing a range of effects that can be fine-tuned over large distances. Most lanthanide chelates were synthesized and specifically attached to the proteins through one reactive arm. Depending on the number of bonds between the lanthanide and the protein backbone, such probes can have considerable flexibility. In our laboratory, a paramagnetic probe was designed[60] with two attachment points on the protein surface and, consequently, reduced mobility. A drawback of this probe is the presence of several isomeric forms of the lanthanide chelate that cause several resonances for a spin that experiences a PCS.

PCS values can provide long-range-distance information, of 5–40 Å from the paramagnetic center,[20] by exploiting the magnetic-anisotropy properties of different lanthanides with fast electron-spin relaxation. The magnitude of the PCS is proportional to the inverse cube of the distance between the paramagnetic metal ion and the nucleus observed. Among the lanthanides, Gd3+ is a useful PRE agent because of its long electronic relaxation time.

Results and Discussion

Synthesis and properties of the caged lanthanide NMR probe (CLaNP) molecule: The strategy to synthesize the modified DOTA derivative 3b, capable of ligation to two proximate engineered cysteine residues of a protein, is depicted in Scheme 1. Commercially available cyclen was transformed in three synthetic steps[65] to obtain the bis(tert-butyloxycarbonylmethylene) derivative, compound 1. Reaction of 1 with benzyl bromoacetate in the presence of potassium carbonate gave compound 2 in moderate yield after silica-gel column chromatography. Liberation of two carboxylic acid functionalities was accomplished by removal of the two benzyl groups of 2 by using a catalytic amount of palladium on carbon under a hydrogen atmosphere. The resulting dicarboxylic acid intermediate was coupled to 2-(aminoethyl)mercaptoethanesulfonate with the aid of 1-(3-dimethylaminopropyl)-3-ethylcarbodiimide-HCl (EDCI) to provide compound 3a in 26% yield over the two steps after purification by HPLC. Removal of the tert-butyl ester groups was accomplished with 33% trifluoroacetic acid in di-
chloromethane to give compound 3b in 36% yield after HPLC purification.

**Attachment of the CLaNP molecules to E51C/E54C-Paz:**

The CLaNP molecules (Figure 1A and B) were attached to the test protein in a bidentate fashion through two engineered cysteine residues (Figure 1C), limiting the rotational freedom of the lanthanide ion with respect to the protein. We found that attachment of the bidentate CLaNP molecules does not depend critically on the position or distance of the two Cys residues. Various proteins were tagged successfully with Cys–Cys CB distances within the range of 5–10 Å (not shown). The minimal distance between the sulfur atoms should be large enough to avoid intramolecular disulfide-bridge formation. Generally, exposed polar residues are selected for mutation to minimize effects on protein structure.

A variant of pseudoazurin (Paz) with mutations E51C and E54C was used to test the effects of the CLaNP molecules. Paz is a small copper-containing protein (14 kDa) and does not contain any exposed cysteine residues. The only cysteine in the wild-type protein provides one of the four ligands to the copper.

Attachment of the CLaNP-3 molecule to the protein was checked by mass spectrometry. Mass spectroscopy analysis (TOF, ES+) revealed a mass of 14316±2 Da for the ⁹⁰CLaNP-3 bound to ¹⁵N-Paz, consistent with binding of the probe to the protein through both reactive arms (expected mass 14317 Da for a 99% incorporation of ¹⁵N in ⁹⁰CLaNP-3–Paz). The attachment of the CLaNP-1 molecule to Paz in a bidentate manner was confirmed previously.[60]

The [¹⁵N,¹H] HSQC spectra of the ⁹⁰CLaNP-1, the diamagnetic control probe, bound to this mutant exhibited only small chemical-shift perturbations for several residues in the neighborhood of the attachment site, relative to the wild-type Paz spectra, however the remaining amide resonances remained unaffected. The overall similarity between these two structures was also confirmed by X-ray crystallography results.[60] The differences in chemical shift between ⁹⁰CLaNP-1 and ⁹⁰CLaNP-3 bound to this Paz mutant, for most of the amide protons, are less than 0.02 ppm (see Figure S1 in the Supporting Information). This indicates that CLaNP-3 binding also does not affect the overall three-dimensional structure of Paz. Notably, similar to the case of CLaNP-1 bound to this mutant Paz, the resonances for C51, C54, and the residues in their immediate vicinity are not observed in the HSQC spectra of the diamagnetic sample. The disappearance of these signals might be attributed to line broadening due to a slow exchange process in the residues on the loop to which the CLaNP is attached.

**Pseudocontact shifts caused by ⁹⁰CLaNP-3:** Lanthanide-substituted DOTA exhibits several isomers in solution, differing in conformation and water coordination.[66] The Ln–DOTA conformers represent two enantiomeric pairs of diastereoisomers. Two sets of peaks are generally observed in NMR spectra due to slow conversion of the DOTA isomers.[67,68] In the HSQC spectra of the diamagnetic ⁹⁰CLaNP-3–Paz, single peaks are observed for all residues, indicating that the protein amide groups are too far from the probe to sense possible exchange processes. In the case of the paramagnetic Paz–⁹⁰CLaNP-3, nearly all peaks exhibit changes in chemical shift that are of about the same size in the proton and nitrogen dimensions. This is a hallmark of PCS because the shifts are independent of the type of nucleus when expressed in ppm. Most residues show single peaks, some exhibit a small splitting in the ¹H dimension only, and some amides close to the probe (<18 Å) are not observed at all (Figure 2A). We attribute these observations to the DOTA isomers, as will be discussed below.

Due to the attachment of CLaNP-3 to the protein through its two reactive arms, the symmetry in the coordination complex is altered, resulting in four distinguishable isomers, giving rise to four possible χ tensors. The peaks that are split in the ¹H dimension represent the effects of two χ tensors, caused by exchange between two conformers of the CLaNP molecule.[69] The fact that the splittings are small and even absent for most peaks suggests that these two conformers have χ tensors with similar orientations.

Resonances due to the amide group at >10 Å from the metal should be observable, in agreement with the results obtained with ⁹⁰CLaNP-1–Paz.[69] The absence of resonances from amides close (<18 Å) to the metal cannot be caused

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Figure 1. Chemical structures of A) the bis(MTS) derivative of DTPA, CLaNP-1,[60] and B) the bis(MTS) derivative of DOTA, CLaNP-3. The leaving groups of the probe molecule are shown in gray. C) Schematic representation of the attachment reaction between the probe molecule and the cysteine residues of the protein.
by the paramagnetic relaxation effects. Neither can it be attributed to exchange effects in the protein due to probe attachment, because the resonances are clearly observed in the diamagnetic control, YCLaNP-3–Paz. We suggest that a second exchange process in the DOTA part of CLaNP-3 results in doubling of the resonances. If the exchange between these forms is on the order of several hundred per second, the resonances of amides with two very different PCS will be broadened; however, those with only slightly different PCS will show a single, average resonance. Thus, the signals of amides close to the metal will mostly broaden out, whereas those from more-distant amides will show a single resonance. The published data[70,71] are contradictory with regard to the sizes of the exchange rates of the various interconversions. Therefore, no attempt will be made here to assign the two proposed exchange processes to specific conformer isomerizations.

To establish whether the observed PCS could be used for structural calculations, the data were fitted to Equation (1) (see Experimental Section), yielding a good correlation for observed versus predicted PCS with a single apparent tensor, \( \chi^{pp} \) (Figure 2B). These results indicate that CLaNP-3 can provide useful PCS data, despite its exchange behavior.

**Paramagnetic relaxation enhancement (PRE) caused by \( ^{64}\text{Gd} \)ClaNp molecules:** Unlike other lanthanides, \( ^{64}\text{Gd} \) has a slow electronic relaxation rate, causing strong relaxation effects on surrounding nuclear spins. These effects are isotropic to a good approximation[72] and are, therefore, independent of the presence of more than one isomeric of the CLaNp molecules. The PRE falls off with the sixth power of distance from the \( ^{64}\text{Gd} \) ion [Eq. (3), Experimental Section], making it a very sensitive tool for distance measurements. The \( ^{64}\text{Gd} \) ion does not cause significant PCS.

To test the usefulness of PRE caused by \( ^{64}\text{Gd} \), the effects of \( ^{64}\text{Gd} \)ClaNp molecules on Paz amide protons were determined by acquiring \([^{15}\text{N},^{1}\text{H}] \) HSQC spectra of \( ^{64}\text{Gd} \)ClaNp-1 and \( ^{64}\text{Gd} \)ClaNp-3 bound to the \(^{15}\text{N}\)-labeled Paz. The intensities were compared with those in control spectra of \(^{6}\text{Cl} \)ClaNp–Paz. Large intensity differences were observed for numerous peaks. Figures 3A and B show the peak-intensity ratios of \(^{6}\text{Cl} \)ClaNp-1–Paz to \(^{64}\text{Gd} \)ClaNp-1–Paz and of \(^{6}\text{Cl} \)ClaNp-3–Paz to \(^{64}\text{Gd} \)ClaNp-3–Paz, respectively, for all Paz residues observed. The residues that appear unaffected by the \( ^{64}\text{Gd} \) ion show an intensity ratio of 1. For residues ob-

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**Figure 2.** Pseudocontact shifts of \( ^{6}\text{Cl} \)ClaNp-3 bound to Paz: A) a section of the overlaid \([^{15}\text{N},^{1}\text{H}] \) HSQC spectra of \(^{6}\text{Cl} \)ClaNp-3–Paz–Cu(I) (gray) and \(^{6}\text{Cl} \)ClaNp-3–Paz–Cu(I) (black). Some of the shifted amide resonances are connected by lines with slopes close to unity to indicate the pseudocontact nature of the shift; B) correlation between measured and predicted PCS for 70 amide protons of Paz. Predicted PCS were back-calculated with the optimized apparent \( \chi \) tensor. The axial and rhombic contributions of the \( Dc \) are given in units of \( 10^{-26} \text{m}^4 \).

**Figure 3.** Relaxation enhancements of \( ^{64}\text{Gd} \)ClaNp-1 and \( ^{64}\text{Gd} \)ClaNp-3 bound to Paz. Relative intensities of the diamagnetic versus paramagnetic sample for all observed residues in the NMR experiment: A) ratio of \(^{6}\text{Cl} \)ClaNp-1–Paz intensities to \(^{64}\text{Gd} \)ClaNp-1–Paz intensities; B) ratio of \(^{6}\text{Cl} \)ClaNp-3–Paz intensities to \(^{64}\text{Gd} \)ClaNp-3–Paz intensities. Error bars were calculated by using the background intensity. C) Map of the relaxation effects caused by the \( ^{64}\text{Gd} \) on Paz. A decrease in intensity of the amide resonances in \(^{6}\text{Cl} \)ClaNp-1–Paz versus \(^{6}\text{Cl} \)ClaNp-1–Paz of more than five times is shown in red, between five and two times in orange, between two and 1.2 times in yellow, and below 1.2 times in blue. The five- and two-fold borders are marked in panels A and B by dotted lines. The unassigned and Pro residues are gray. The CLaNp-1 molecule with the bound \( ^{3}\text{Y} \) is purple. The structure of the \(^{6}\text{Cl} \)ClaNp-1–Paz was taken from the PDB entry 1PY0[60]. D) Relative intensity \( (I_{\text{dia}}/I_{\text{para}}) \) for CLaNp-1–Paz residues versus the distance from the \( ^{64}\text{Gd} \) ion to the amide protons. The solid line represents a fit of the data by using the equation \( y = 1 + 4.61x^{-6} \).
served in the diamagnetic sample that had disappeared in the paramagnetic one, the noise-level intensity was used to calculate the minimal ratio of intensities in the two forms. The broadening effects of the Paz resonances caused by the Gd$^{3+}$ in the CLaNP-1 protein complex were then mapped onto the crystal structure of CLaNP-1–Paz (Figure 3C). A clear distance-dependent decrease in intensity of the Paz peaks with respect to the lanthanide is apparent (Figure 3D).

**Modeling the position of the lanthanide with respect to the protein by using NMR-derived distances as restraints:** To quantify the distances from the HSQC peak intensities, the $R_2^{para}$ values were determined by using Equation (2) (Experimental Section).[46] The use of intensity ratios yields $R_2^{para}$ values that are slightly less precise than those obtained through direct $R_2$ determination. However, it has been demonstrated that other approximations concerning PRE distance determination introduce larger errors.[49,51] Furthermore, this approach is simpler and is also applicable to larger protein complexes. According to Equation (3), at the magnetic field used, $R_2^{para}$ is proportional to the correlation time $\tau_c$. The latter depends on the electronic relaxation time of the Gd$^{3+}$ ion ($\tau_e$), the rotational correlation time of the metal–proton vector ($\tau_r$), and possible exchange processes ($\tau_{ex}$). The contribution of the $\tau_{ex}$ is likely to be relatively small. Both $\tau_e$ and $\tau_r$ are expected to be in the low-nanosecond range. To obtain $\tau_c$, the $R_2^{para}$ values were fitted to Equation (3) by using the crystal structure of Paz–$^{3}$CLaNP-1 as a reference for the distances, and with $\tau_c$ as the only variable. Values of 4.0 ± 0.9 and 5.9 ± 1.2 ns were obtained for $^{3}$CLaNP-1–Paz and $^{3}$CLaNP-3–Paz, respectively, indicating that within the error, the $\tau_c$ values are the same.

The NMR-derived distances obtained with these $\tau_c$ values were used to optimize the position of the Gd ion relative to Paz for both CLaNP molecules. Restrained energy minimization by using only the distance information placed the Gd$^{3+}$ ion at 2.4 Å and 4.8 Å from the lanthanide position observed in the crystal structure of Paz–CLaNP-1[160] for $^{3}$CLaNP-1–Paz and $^{3}$CLaNP-3–Paz, respectively (Figure 4A). Due to dependence on the sixth power of the distance, the predicted distances are not very sensitive to errors in $\tau_c$. A 50% higher or lower value for the $\tau_c$ led to an optimized position of the Gd ion that differed by 1.3 Å from the one obtained above.

These calculations suggest that the solution position of the lanthanide in CLaNP-1 is similar to that observed in the crystal. The lanthanide position differs slightly for CLaNP-3, which may be caused by the differences in the structure between the two lanthanide probes. The correlation plots for the observed versus predicted distances, based on this position for the Gd$^{3+}$ ion, show that distances of between 20 and 30 Å can be measured within an error margin of ±3 Å for both CLaNP-1–Paz and CLaNP-3–Paz (Figures 4B and C, respectively).

**Conclusion**

Paramagnetic effects can provide long-range-distance information that can be used in combination with other NMR-based restraints to determine the solution structure of proteins, global fold determination of larger proteins, and characterization of their different states. For applications to macromolecules that do not contain paramagnetic centers, paramagnetic surface tags can be employed. In this study, we present a new paramagnetic probe, covalently bound to a...
protein in a bidentate fashion. The good agreement between observed and calculated PCS and PRE data indicates that the bidentate attachment helps to limit the conformational freedom of the metal relative to the protein. Thus, this probe can be used as a relaxation agent and, with some approximations, as a pseudocontact-shift agent, to obtain long-distance restraints for solving the solution structures of large proteins and protein–protein complexes. Such studies are currently in progress.

### Experimental Section

Gadolinium(III) acetate (99.9%), yttrium(III) nitrate, and the HBr salt of S-(2-methylaminomethyl)-methanesulfonate (MTSO) were used as purchased without further purification. Reactions were monitored by TLC analysis using DC-fertigkoff (1720). Experimental Section

Protein production and purification: The expression plasmid containing the pseudoazurin gene of *Alcaligenes faecalis* S-6 and the mutagenesis of the residues 51 and 54 to cysteine, E51C/E54C, have been described previously.[60] The procedure for expression of the mutant gene in *Alcaligenes faecalis* was performed according to a published protocol[60] to yield two fractions of CLaNPyPaz molecules. In short, the Paz-E51C/E54C complex, from here on called CLaNPyPaz,[73] was monitored by LC-MS and H NMR spectroscopy. The pH was immediately brought to 7 and CLaNPyPaz was reacted with the protein (see below).

**Protein binding to 3b:** Formation of the CLaNPyPaz-3 molecule: Compound 3a (90 mg, 0.11 mmol) in CH3Cl (9 mL) was treated with TFA (4.5 mL) under an argon atmosphere at 0°C. The reaction was complete after 5 h, as monitored by LC-MS. The mixture was concentrated under reduced pressure and the residue was purified by RP-HPLC: flow rate 4.5 mL min⁻¹, step 0.5 CV; 7% buffer B followed by a linear gradient of 30 CV; 7–16% buffer B; R, 2.7 CV, yielding compound 3b (28 mg, 36%).

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**Attachment of the CLaNPyPaz molecule to the Cu-containing E51C/E54C-Paz:** Binding of CLaNPyPaz molecules to pseudoazurin through the two engineered Cys residues was performed as described,[60] but the step of replacement of Paz cofactor with Zn. In short, the Paz-E51C/E54C mutant was incubated with DT5 (5 mM) for 1 h at RT to dissociate the protein dimers. This was followed by removal of DT5 on a desalting PD10 column equilibrated with degassed sodium phosphate buffer (20 mM, pH 7.0). Five molar equivalents of the probe molecule were added slowly to the protein solution and the reaction was left to proceed overnight at 4°C under semiaerobic conditions. The protein-bound probe was then passed through a Superdex G75 column equilibrated with sodium phosphate buffer (20 mM, pH 7.0) containing NaCl (150 mM). Mass spectra analysis showed that almost all protein (>95%) treated in this way contained the bound probe.

**NMR spectroscopy:** The NMR samples of CLaNPyPaz (100–400μg) were prepared in sodium phosphate buffer (20 mM, pH 7.0) containing one equivalent of sodium ascorbate and 6% (v/v) D2O. [2H17]HSQC experiments were acquired at 303 K by using a Bruker Avance DMX600 spectrometer. The spectral widths used were 32 and 12.6 ppm for the 1H and 13C resonances, respectively.

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and $^1$H dimensions, respectively. The HSQC spectra were acquired with 2048 and 256 data points in $t_1$ and $t_1$, respectively. Data processing was performed by using the AZARA program (from http://www.bio.c.ucl.ac.uk/azara) and the spectra were analysed by using the program "Ansig for Windows."[39] The intensities of the crosspeaks were obtained by using parabolic peak fitting in the AZARA program. For resonances that were broadened beyond detection in the paramagnetic sample, an upper limit of the intensity was set by using the noise level. Assignments of the $^1$H and $^{13}$N amide nuclei of CuI(Paz) ESI1CE/S4C were taken from previous work.[39][40]

Determination of the magnetic-susceptibility tensor: The pseudocontact shifts ($\delta_0^r$) caused by bonding of Yb$^{3+}$ onto Paz were measured from the difference between the chemical shifts observed for the paramagnetic $^{19}$F-CLaNP-3-Paz and the diamagnetic $^{19}$F-CLaNP-3-Paz. The size of the pseudocontact shift experienced by a nucleus under the influence of a paramagnetic center is described by Equation (1):

$$\delta_0^r = \frac{1}{2\pi\gamma_0} \left[ \Delta\chi_{\text{ax}} \left[ \sin^2(\theta_0) - 1 \right] + \frac{3}{2} \Delta\chi_{\text{dy}} \sin(2\theta_0) \right]$$

(1)

in which $r$, $\theta$, and $\Omega$ are the polar coordinates of the nucleus with respect to the principal axes system of the magnetic-susceptibility tensor. $\Delta\chi_{\text{ax}}$ and $\Delta\chi_{\text{dy}}$ are the axial and rhombic anisotropy components of the magnetic-susceptibility tensor of the lanthanide ion, and are defined as $\Delta\chi_{\text{ax}} = \chi_{z} - \frac{1}{2}(\chi_{x} + \chi_{y})$ and $\Delta\chi_{\text{dy}} = \chi_{y} - \chi_{x}$, in which $\chi_{x}$, $\chi_{y}$, and $\chi_{z}$ are the orthogonal components of the magnetic-susceptibility tensor. The apparent $\chi$ tensor was obtained by Euler rotation of the Gd-centered molecular coordinate frame and by fitting of the $\delta_0^r$ values to Equation (1), according to the procedure described.[39] The data were then obtained by adding hydrogen atoms to the crystal structure of CLaNP-1-Paz (PDB entry 1PY0,[57] by using the program Xplor-NIH.[58]) The Gd$^{3+}$ position relative to the Paz structure was that obtained after optimization by using $^{19}$F-CLaNP-3-derived relaxation constraints (see below; the $x$, $y$, $z$-coordinates of the Gd were 5.083, 14.480, 2.680 in the frame of 1PY0, respectively).

Pseudocontact shifts for 70 amide resonances were used in the calculation. For eight shifted resonances clearly showing split peaks in the $^1$H dimension, an average $\delta_0^r$ was used, with the distance to the mean as the error value. The Euler angles obtained for rotation of the Gd-centered coordinate frame into the $\chi$ tensor frame are $\alpha = 140^\circ \pm 2^\circ$, $\beta = 112^\circ \pm 1^\circ$, $\gamma = 100^\circ \pm 6^\circ$, and $\Delta\chi_{\text{dy}} = 5(\pm 0.2) \times 10^{-3} \text{ m}^3$, $\Delta\chi_{\text{ax}} = 1(\pm 0.4) \times 10^{-3} \text{ m}^3$ (for $\Delta\chi_{\text{dy}} > \Delta\chi_{\text{ax}} > 0$).

PRE distances and optimization of the Gd positions: The intensity ratio of the amide resonances of the paramagnetic and diamagnetic form of the protein can be related to the paramagnetic contribution of the transverse $^1$H relaxation according to Equation (2),[30] assuming that the total INEPT relaxation time of the signal ($9\text{ms}$).

$$I_{\text{PRE}} = \frac{R_{\text{PRE}} \exp \left( -R_{\text{PRE}} \gamma_0 \right)}{R_{\text{PRE}} + R_{\text{g}} \gamma_0}$$

(2)

in which $I_{\text{PRE}}$ and $I_{\text{Av}}$ are the peak intensities for an amide resonance in the PRE and Av, respectively. For eight shifted resonances clearly showing split peaks in the $^1$H dimension, an average $\delta_0^r$ was used, with the distance to the mean as the error value. The Euler angles obtained for rotation of the Gd-centered coordinate frame into the $\chi$ tensor frame are $\alpha = 140^\circ \pm 2^\circ$, $\beta = 112^\circ \pm 1^\circ$, $\gamma = 100^\circ \pm 6^\circ$, and $\Delta\chi_{\text{dy}} = 5(\pm 0.2) \times 10^{-3} \text{ m}^3$, $\Delta\chi_{\text{ax}} = 1(\pm 0.4) \times 10^{-3} \text{ m}^3$ (for $\Delta\chi_{\text{dy}} > \Delta\chi_{\text{ax}} > 0$).

$$r = s = \frac{1}{15} \frac{\mu_0}{4\pi} \frac{1}{R_{\text{PRE}}} \sqrt{\gamma_0^2 R_{\text{g}}^2 (1 + J + 1)} \left( 4r_1 + 3r_2 \right) \left( 1 + \alpha p_0 r_1^2 \right)$$

(3)

in which $\mu_0$ is the magnetic permeability. $\gamma_0$ is the proton gyromagnetic ratio. $\mu_0$ is the Bohr magneton, $g$ is the electronic g factor, $\alpha p_0$ is the Larmor frequency of the proton spin, $J$ is the quantum number for the Gd (7/2),[31] $r$ is the lanthanide-to-amide proton distance in the protein, and $r_1$ is the total correlation time. The $r_2$ values were obtained from linear fits of $R_{\text{PRE}}$ versus $r_2$ from the crystal structure of CLaNP-1–Paz.[30] Values of 4.0 and 5.9 ns were obtained for $^{19}$F-CLaNP-1–Paz and $^{19}$F-CLaNP-3–Paz, respectively.

The distances were separated into three classes of restraints. The first class comprised the residues for which the intensity of the amide resonances was affected more than five-fold. These residues were considered to be too close to the paramagnetic center for an accurate distance to be measured. A restraint with only an upper boundary of 20 Å was defined for these residues, which means they can be found anywhere up to 20 Å from the lanthanide. Given that the distance from the lanthanide ion in a CLaNP molecule to the nearest amide in the protein is about 8–9 Å, a “blind region” on the protein can be estimated by using this probe to be 11–12 Å. The second class contained the residues for which the intensity of the amide resonances was affected less than five-fold, but more than 1.2-fold. For these residues, a restraint to the experimental distance was defined with an upper and lower boundary of 3 Å. The third class comprised the residues for which the peak intensity was decreased less than 1.2-fold and these residues were considered to be too far away to be affected by the paramagnetic probe. To these residues, a restraint with only a lower boundary of 30 Å from the metal was defined. The number of restraints in each class is listed in Table 1.

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</tbody>
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[a] Total number of assigned residues in the spectra, including the residues for which overlapping resonances could not be used for an accurate linewidth determination.

The Gd$^{3+}$ position with respect to the protein was determined through restrained energy minimization by using the program Xplor-NIH.[39] In these calculations, the protein was kept fixed (PDB, entry 1PY0),[57] and the Gd ion was allowed to move starting from ten random positions, solely under the influence of a distance-restraints energy term. This procedure yielded convergence to single positions for both CLaNP-1 and CLaNP-3 bound to Paz.

Acknowledgements

Dr. J. A. Peters is acknowledged for careful reading of the manuscript. H. van den Elst and N. Meeuwenoord are thanked for their excellent HPLC and LC/MS assistance. M.D.V., M.P., and M.U. were supported by the Netherlands Organization for Scientific Research, VIDI grant 700.52.425.

We use the name CLaNP (Caged Lanthanide NMR Probe) to be consistent with earlier work. CLaNP-2 is an EDTA-based molecule with two functional groups for protein attachment. This molecule bound to proteins did not show sufficient Ln coordination.


Received: June 27, 2006
Published online: November 20, 2006