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Two well-defined motifs in the cAMP-dependent protein kinase inhibitor (PKIα) correlate with inhibitory and nuclear export function

JENNIFER A. HAUER, 1 PHILIPPE BARTHE, 2 SUSAN S. TAYLOR, 1 JOSEPH PARELLO, 3 AND ANDRÉ PADILLA 2

1 Department of Chemistry and Biochemistry, University of California, San Diego, 9500 Gilman Drive, La Jolla, California 92093-0654
2 Centre de Biochimie Structurale CNRS UMR 9955, INSERM U414, University of Montpellier I, 15 Av. Charles Flahault, 34060 Montpellier Cedex 1, France
3 UPRESA, 5-74 CNRS, University of Montpellier I, France, and Cancer Research Center, The Burnham Institute, 10901 N. Torrey Pines Rd., La Jolla, California 92037

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Abstract

The heat stable inhibitor of cAMP-dependent protein kinase (PKIα) contains both a nuclear export signal (NES) and a high affinity inhibitory region that is essential for inhibition of the catalytic subunit of the kinase. These functions are sequentially independent. Two-dimensional NMR spectroscopy was performed on uniformly [15N]-labeled PKIα to examine its structure free in solution. Seventy out of 75 residues were identified, and examination of the Cα chemical shifts revealed two regions of upfield chemical shifts characteristic of α-helices. When PKIα was fragmented into two functionally distinct peptides for study at higher concentrations, no significant alterations in chemical shifts or secondary structure were observed. The first ordered region, identified in PKIα (1–25), contains an α-helix from residues 1–13. This helix extends by one turn the helix observed in the crystal structure of a PKIα (5–24) peptide bound to the catalytic subunit. The second region of well-defined secondary structure, residues 35–47, overlaps with the nuclear export signal in the PKIα (26–75) fragment. This secondary structure consists of a helix with a hydrophobic face comprised of Leu37, Leu41, and Leu44, followed by a flexible turn containing Ile46. These four residues are critical for nuclear export function. The remainder of the protein in solution appears relatively unstructured, and this lack of structure surrounding a few essential and well-defined signaling elements may be characteristic of a growing family of small regulatory proteins that interact with protein kinases.

Keywords: 2D-NMR; cAPK; conformational disorder; kinase inhibitor; nuclear export signal; PKIα

Signaling through cAMP-dependent protein kinase (cAPK) is a common pathway for many cellular processes. Regulation of cAPK is achieved by both inhibition and subcellular localization. The best understood control mechanism for cAPK kinase activity is achieved through the regulatory (R) subunit. Two catalytic (C) subunits bind to a dimer of R subunits to yield an inactive holoenzyme. Cooperative binding of two molecules of cAMP to each R subunit causes dissociation of the holoenzyme complex and release of two active C subunits (Taylor et al., 1990). In addition to being inhibitors of C, the R subunits also contribute directly to subcellular localization of cAPK by interacting with AKAPs. These proteins anchor the cAPK holoenzyme complex to a wide variety of cellular locations (Dell’Acqua & Scott, 1997).

Another mode by which cAPK is simultaneously localized and inhibited is by PKI, the heat stable protein kinase inhibitor identified by Walsh et al. (1971). While PKIα has only 75 residues, it has two sequentially distinct functional elements. The first third of PKIα contains the PKI pseudo-substrate sequence preceded by a high affinity binding region. This segment contains nearly all of the residues critical for the subnanomolar inhibitory constant of PKIα.
addition to inhibiting the kinase, PKIα residues 5 to 22 of PKI~a was also shown in solution to contain an α-helix at its N-terminus by 1D NMR (Reed et al., 1989). In addition to inhibiting the kinase, PKIα also traffics the catalytic subunit from the nucleus to the cytoplasm through its Nuclear Export Signal (NES). The NES (residues 37–46) consists of a regular repeat of large hydrophobic amino acids, and acts to rapidly and actively export from the nucleus to the cytoplasm any protein to which it is bound, including the catalytic subunit in a C:PKIα complex (Fantozzi et al., 1994; Wen et al., 1995).

Based on CD and FTIR studies, PKIα does not appear to be highly structured in solution (Thomas et al., 1991). CD measurements indicated that PKIα has 21% α-helix, 15% antiparallel β-sheet, 34% turn, and 30% random coil. This suggested the presence of another helix in PKIα, in addition to the N-terminal helix described previously.

Determining the structural features of PKIα is critical for understanding its bifunctionality. How does the structure of such a small protein allow for both high affinity binding to the C subunit and export of the enzyme complex from the nucleus? A structural study of the inhibitor free in solution was undertaken by 2D NMR to define structural features within the full length protein. Due to severe overlapping of proton resonances, uniformly 15N enriched PKIα was used to resolve most of the proton resonances as well as the proton–proton Nuclear Overhauser effect (NOE) connectivities. Two regions of α-helix were identified by Chemical Shift Index analysis, but overall there was a lack of medium- and long-range NOEs. This indicated that PKIα has a rather flexible structure in solution, and the presence of different shape “conformers” of PKI was apparent from early studies as well (Whitehouse & Walsh, 1982). At this point we adopted a strategy to analyze, by homonuclear 2D NMR, two functionally separate fragments, allowing the secondary structure of each functional domain to be more precisely determined. We identified two α-helices, which overlap with the two functional elements of PKIα. The remainder of the molecule appears to be highly disordered.

Results

Sequential assignments of [15N]PKIα

Figure 1 shows the HSQC spectrum of uniformly 15N labeled [Gly0]PKIα. As reported in Figure 2 and Table 1 of the Electronic Supplementary Material, 70 of 75 spin systems expected were sequentially assigned through a combination of HMQC-TOCSY and HMQC-NOESY spectra. Side-chain CαH3 resonances of Leu residues were assigned using NOE connectivities in the HMQC-NOESY spectra. One spin system has been tentatively assigned to Asp24 according to standard chemical shift values for aspartic acid (Wüthrich, 1986), and the most high-field shifted 15N-NH cross peak has been assigned to Arg15 by comparison with the [Gly0]PKIα (1–25) peptide spectra (see below).

No medium- or long-range NOEs were observed in the HMQC-NOESY spectra. This may be because the protein appears to be flexible in solution, with a relatively low content of secondary structure elements. This would reduce the NOE intensities through an equilibrium between relatively structured and less structured states of the protein. Alternatively, analysis of PKIα by dynamic light scattering at the concentration used for the NMR experiments showed that the inhibitor aggregates in solution (data not shown). The presence of such polymeric species could affect the NOE intensities through dynamic processes involving a variety of environments to give proton nuclei.

Identification of secondary structure elements in PKIα

To identify any regions of secondary structure within full length PKIα, the CαH chemical shift values in ppm of the 70 assigned residues were compared with the random coil chemical shift values for residues in a GGXGG peptide (Merutka et al., 1995), revealing two regions of upfield shifted CαH chemical shifts (Fig. 3). Chemical shift index analysis of these CαH chemical shift values identified these upfield shifted regions as helical. The first α-helical region extends from residue 7 to 13, and corresponds to the high affinity binding helix seen in the crystal structure. Residues 2 and 3 also have indices of −1, suggesting an earlier start to the helix. The second helix identified is from residue 34 to 43 of PKIα, and overlaps with the Nuclear Export Signal of PKIα. The overall distribution of −1 indices in this region is less than the previous helix, but two of the residues with “0” index values (35 and 42) differ from the tabulated CαH chemical shifts by −0.09 ppm, indicating a consistent trend in upfield chemical shifts and these residues are indicated in Figure 3 by asterisks. These residues were included in defining the helix. Residue 44 has an index value of +1, terminating the helix. One other region of −1 chemical shift indices is seen: a stretch of three −1 values from 54 to 56. Three −1 values are not enough to define an α-helix, but suggest some secondary structure in this region. The remainder of PKIα is not structured.
Sequential assignments of PKI fragments

Based on the chemical shift index identification of the two helices but the lack of corresponding medium- and long-range NOEs to substantiate these helices in full length PKI, we decided to investigate the secondary structure of two functionally distinct PKI fragments, with the expectation that polydispersity in solution could be reduced, if not totally avoided.

Proton resonances of the two fragments of $^{15}$N-labeled PKI were assigned by the standard 2D-NMR strategy (Fig. 2, Table 1) (Wüthrich, 1986). For the N-terminal fragment, all sequential assignments were made, with the exception of the sequential connectivity between Ser13 and Gly14. Due to the low solubility of the N-terminal fragment near neutral pH, all studies were performed at pH values close to 4. In the C-terminal fragment, all but five sequential $d_{AN}$ connectivities from Gly32 to Glu65 were assigned. This fragment has a high content of serine and glutamic acid residues, and several of their NH-$C^a$H cross peaks in the fingerprint region are superimposed. Three Ser residues, Ser28, Ser29, and Ser31, among nine in total, could not be identified. The residues Leu26 and Ala30 were assigned at the latest stage to the remaining spin systems. Despite strong overlap, several $J_{AN}$ coupling constants were measured.

Comparison of chemical shifts

We compared the chemical shifts of full length PKI$\alpha$ and the PKI$\alpha$(26–75) fragment to look for any conformational changes that may have occurred upon fragmentation. Five chemical shift differences could not be calculated due to the absence of proton resonance assignments in either full length or fragmented PKI$\alpha$. As can be seen in Figure 4, chemical shift differences of both NH and CaH are less than 0.05 ppm for most residues in PKI$\alpha$(26–75). This indicates that no dramatic change occurs in the magnetic environment of the NH and CaH protons, and therefore in the secondary and tertiary structure, upon fragmentation. No large changes are evident in the N-terminal fragment either, but this is less meaningful since the experiments were performed at quite different pH values. Any differences observed are likely to correspond to local conformational changes, since none of them involves several adjacent residues in the sequence.
Secondary structure of PKI fragments

α-Helices are characterized by sets of \((i, i+3)\) NOE connectivities, slow NH exchange rates, and \(J_{\alpha N}\) coupling constants less than or equal to 6 Hz. By such criteria, two helices were identified in the fragments of PKI\(\alpha\), corresponding to the regions of α-helix identified by the Chemical Shift Index. Helix I extends from Asp2 to Ser13 of [Gly0]PKI\(\alpha\) (1–25) and Helix II from Asn35 to Leu44 of PKI\(\alpha\) (26–75). Molecular models of each of the two α-helical regions were constructed from the distance constraints imposed by the NOEs. Superimposition of the family of structures generated for the backbone stretch of Thr1–Ser13 gave an average rmsd value of 1.11 Å (Fig. 5) and ensemble-averaged refinement was applied to the average structure.

The N-terminus of PKI\(\alpha\) (Thr-Asp-Val-Glu) appears to act as an N-capping motif. In such a motif, the first and fourth residues of a helix make two additional intrahelical hydrogen bonds, but depart from the helical values for Φ and Ψ angles (Presta & Rose, 1988; Richardson & Richardson, 1988). Slow NH exchange rates were observed for Val3 and Glu4, and we found a NOE between CgH3 Thr1 and NH Val3, in agreement with the geometrical requirements of an N-capping sequence (Zhou et al., 1994). Some variability does occur in the N-terminal stretch as shown by ensemble-averaged refinement and the B-factors for these first four residues are higher than for the rest of the helix (Fig. 6). We conclude overall that the first helix starts with the N-capping sequence from Thr1 to Glu4, a highly conserved sequence in all PKI isoforms identified. A small \(J_{\alpha N}\) value and a slow NH exchange rate for Ser13, in contrast to a fast NH exchange for Gly14, indicates that Ser13 is the last residue in Helix I.

A second helix (Helix II) was found within the C-terminal fragment, from Asn35 to Leu44, defined by a set of \(da\alpha\)(\(i, i+3\)) and \(da\beta\)(\(i, i+3\)) NOE connectivities (Fig. 2). This helix, following ensemble-averaged refinement applied to the average structure, is

Fig. 5. A: Space filling model of PKI\(\alpha\) (1–13). The hydrophobic residues critical for high affinity binding are labeled and shown in red. B: Superimposition of 23 structures for PKI\(\alpha\) (1–25) from the NMR data in purple and PKI\(\alpha\) (5–24) from the crystal structure in red (1APM) (Knighton et al., 1991b). The location of the three Arg residues is highlighted in each structure and the position of the residues is indicated in the PKI\(\alpha\) (5–24) structure. C: A 90° rotation of the view in B with the three Arg residues highlighted in each peptide.
displayed in Figure 7. The average RMSD value for the superimposition of this backbone stretch was 1.30 Å. In contrast to Helix I, this helix displays several \( J_{\text{NN}} \) constants that are slightly greater than 6 Hz, ranging from 6.3 to 6.6 Hz. Two large \( J_{\text{NN}} \) coupling constant values, 7.4 and 6.3 Hz, were found at the N- and C-terminal residues of the helix, Asn35 and Leu44. This is not surprising since the ends of a helix are expected to “fray” due to weak individual H-bonds. Ensemble-averaged refinement indicates relatively low \( B \)-factors (between 5–10 Å\(^2\)) for the stretch of residues between 35 and 40, whereas higher \( B \)-factors are found for the C-terminal half of this helix (Fig. 6). Several NOE connectivities of type \((i,i+2)\) are found at the C-terminus of Helix II, between Gly43 and Asp45, Leu44 and Ile46, and Asp45 and Asn47. This is consistent with Helix II having a well-defined N-terminal half whereas the C-terminal half is much more flexible. The last observed NOE connectivity of type \((i,i+3)\) indicates the break in Helix II occurs at Leu44. Interestingly, Asp45, Ile46, Asn47, and Lys48 have slow NH exchange rates, which indicates protection from solvent accessibility. Residues 44 to 47, while not participating in Helix II, certainly do not adopt a random conformation, but rather can be better described as a flexible turn structure.

Both helices have amphipathic character with one face of the helix formed by hydrophobic residues: Val3, Tyr7, Phe10, and Ile11 for Helix I (Fig. 5A), and Leu37, Ala38, Leu41, Ala42, and Leu44 for Helix II (Fig. 7). Helix I starts with an N-cap motif, which is less constrained than the core of the helix. In Helix II, the essential leucine residues cluster along the hydrophobic face of the helix, including Leu37, Leu41, and Leu44. Leu39, which does not play a role in the NES, lies on the opposite face. Two polar residues, Glu36 and Lys40, are found on the polar face of Helix II where they may form an ionic interaction.

**Discussion**

Free PKI\(\alpha\) in solution has two \( \alpha \)-helical regions surrounded by stretches of unstructured residues, and each helix overlaps with
a distinct biological function. We show here that Helix I starts with the N-terminal capping box motif, Thr-Asp-Val-Glu, at the N-terminus of the protein, extends to Ser13, and is the helix responsible for high-affinity binding to the C subunit. Residues critical for this interaction have been previously determined by mutational studies (Walsh et al., 1990), and the crystal structure of the C subunit-PKIα (5–24) complex showed that hydrophobic residues on one face of this helix interact with a hydrophobic pocket on the surface of the C-subunit (Knighton et al., 1991b). The N-terminus of PKIα, which extends the helix by one full turn, is a highly conserved region among PKI isoforms, including murine PKIβ2 in which this region is preceded by a 21 amino acid sequence. This N-terminal capping box acts to anchor the formation of this high affinity binding helix of PKI. This extension may create an additional interaction between Asp2 of PKIα and Arg256 of the C subunit since previous work with inhibitory peptides showed a tighter C-subunit affinity for the PKIα (1–24) peptide than a PKIα (5–24) peptide (Scott et al., 1985).

In the crystal structure, Gly14 of PKI is part of a turn from Gly14 to Gly17 followed by an extended structure of the pseudo-substrate site in a groove between the small and large lobes of the kinase (Knighton et al., 1991b). In the protein free in solution, no such turn was observed and no structural information was obtained for the five-residue consensus recognition sequence, encompassing Arg18 to Ile22. This region is much more flexible than the binding helix. A parallel NMR study of PKI peptides from the N-terminal region up to Asp24 showed that bent structures are present in solution, with the consensus recognition sequence running antiparallel to Helix I. However, such structures are highly dependent on the solution conditions, especially the pH (Padilla et al., 1997). The binding of PKIα to the C-subunit is bipartite with the basic consensus site docking with low affinity to the active site cleft and the N-terminal helix required for high-affinity binding. The existence of the helix free in solution followed by a flexible consensus sequence can permit high-affinity docking of PKI to the hydrophobic pocket of the C-subunit subsequent to the long-range electrostatic attraction of the positively charged consensus sequence to the negatively charged active site region of the kinase.

Helix II, from residue 35 to 44, is well defined by medium range NOE connectivities but has fraying N- and C-termini with a C-terminal half less resolved than the N-terminal half. Many of the hydrophobic residues in the 37–46 sequence are critical for the nuclear export function of PKI. As shown in Figure 7, three leucines, Leu37, Leu41, and Leu44 form a hydrophobic face to Helix II. Mutation of Leu41 and Leu44 to alanine in NES consensus II, at least one small

Another region of flexible structure occurs for residues Asp54 to Arg57. In this case, however, a small JaN coupling constant is observed for Ala55, and the daN (54, 56) has a higher intensity than that observed in the Leu44–Asn47 segment. Interestingly, slow NH exchange rates are observed for residues encompassing this region, indicating that it is not fully exposed to solvent. This further defines this region as another flexible turn structure with, as yet, no known function.

In closing, it can be concluded that PKIα contains two relatively stable helices, which are essential for the two functional properties of PKIα: high affinity binding to the C subunit and nuclear export. According to the NMR data, 22 residues are found in these two helical domains within residues 2–13 and 35–44. The presence of 22 helical residues, as inferred from our NMR data, is slightly greater than the α-helical content of 21% previously reported by CD (Thomas et al., 1991). The β-sheet structure determined by CD was not found by NMR in the fragments of PKIα. This may be due to aggregation of full length PKI in solution, where β-sheet structures may form in the aggregates. The remainder of PKIα is thus remarkable for its lack of well-defined or stable secondary structure. p21 and p16, two regulatory proteins that bind to CDK2, display similar properties and are largely unstructured when they are free in solution (Kriswacki et al., 1996; Russo et al., 1996). This property may be characteristic of many of the small proteins that bind to protein kinases and regulate their function.

Materials and methods

Preparation of NMR samples

The [Gly0]PKIα cDNA corresponding to rabbit skeletal muscle with a glycine residue between the start methionine and the first residue was the gift of Dr. R.A. Maurer (Volum Institute, Portland, Oregon). This was transformed into Escherichia coli BL21 (DE3) and grown in minimal media with either (15NH4)2SO4 or 15NH4Cl as the sole nitrogen source. The protein was purified as described previously (Thomas et al., 1991). The numbering of residues throughout this work refers to the rabbit skeletal muscle protein kinase inhibitor (PKIα).

The [I35M]PKIα vector, in a derivative of pET3C with tetracycline resistance, was graciously provided by Dr. R.A. Maurer. The protein was expressed and purified as above. The purified protein was digested with cyanogen bromide in 70% formic acid and the resulting peptides were separated by HPLC chromatography and identified with a gas phase amino acid sequencer with an on-line PHT analyzer (Applied Biosystems, Inc. models 470A and 120). Vydac C18 analytical or semi-prep columns were used. Solvent A was 0.1% TFA in MQ water, and B was 0.85% TFA in ACN/MQ water (95:5). The 15N enriched PKI sample was prepared by mixing 8 mg protein in 450 μL (2.2 mM) of solvent (D2O or a mixture of 95% H2O/5% D2O), and the pH adjusted to 6.6. Samples of [Gly0]PKIα (1–25) were prepared by dissolving 6.2 mg of peptide

Figure 7; Lee et al., 1997.
in 450 μL (5 mM) of solvent (D₂O or a mixture of 95% H₂O/5% D₂O), and the pH adjusted to between 3.8 and 4.1. The PKIα(26–75) sample was prepared by dissolving 5.9 mg of peptide in 450 μL (2.5 mM) of solvent (D₂O or a mixture of 95% H₂O/5% D₂O), and adjusted to a pH of 7.1. All samples were sealed under argon.

Choice of experimental conditions

Preliminary homonuclear NMR studies of PKIα at 32 °C by DQF-COSY and jump-return NOESY techniques indicated low NOE intensities and strong signal overlap (data not shown). Chemical shift dispersion and linewidths were unaffected by increased ionic strength, up to 100 mM NaCl or 50 mM MgSO₄, or decreased protein concentration to 0.5 mM. However, decreasing the temperature had two effects: (1) the NOE intensities increased, and (2) NH exchange was slow enough to allow the use of water presaturation. Even though NH intensities were not affected by temperature changes between 2 and 12 °C, the NOEs involving NH or side-chain protons were stronger at the lower temperature.

NMR techniques

Spectra of ¹⁵N labeled-[Gly0]PKIα and unlabeled [Gly0]PKIα(1–25) and PKIα(26–75) were recorded at 600 MHz on an AMX 600 Bruker spectrometer. Heteronuclear experiments were carried out at 2 and 12 °C. The ¹⁵N-¹H HSQC, ¹⁵N-¹H HMBC-TOCSY, and ¹⁵N-¹H HMQC-NOESY spectra were run with the published sequences (Bodenhausen & Ruben, 1980; Bax et al., 1990; Norwood et al., 1990), respectively. The HSQC was recorded with spectral widths of 1,824 and 6,613 Hz while the HMQC-TOCSY and HMQC-NOESY spectra and were recorded with spectral widths of 1,946 and 6,024 Hz in the ¹⁵N and ¹H dimensions, respectively. For each 2D experiment, 2,048 points by 512 FIDs (128 for the HSQC) were recorded. Water suppression was achieved with 1s presaturation. Eighty scans were collected for the HMBC-NOESY with 150 and 200 ms mixing times, and 32 scans were collected with a mixing time of 48 ms for the HMQC-TOCSY experiments. TIPPI was used in t1 for all experiments (Marion & Wüthrich, 1983).

Homonuclear experiments on the PKI fragments were carried out at 2 °C. The NOESY, TOCSY, and DQF-COSY spectra were recorded with a spectral width of 6,024 Hz, with 1s water presaturation. For each 2D experiment, 4,096 data points by 512 FIDs were recorded. Eighty scans were run for both the NOESY (with a 200 ms mixing time) and the TOCSY (with 40 ms mixing time) experiments, and 128 scans for the DQF-COSY.

Slow exchanging amide protons were determined for each fragment by recording two 30 min COSY experiments at 2 °C just after dissolving the peptides in D₂O. The data collected were 2,048 points by 512 FIDs.

After Fourier transform, data sets of 2,048 by 512 real points were further processed by polynomial baseline correction in both dimensions. Coupling constants between COH and NH were measured from the DQF-COSY experiments after reprocessing the data with 0.3 Hz per point resolution in the F2 dimension. The ¹H-NMR chemical shifts at 2 °C were referenced by taking water resonance at 5.14 ppm referenced against tspd4. All ¹⁵N-NMR chemical shifts were referenced according to external 1.85 M ¹⁵NH₄Cl in 1 M HCl at 24.93 ppm.

Chemical Shift Index analysis

The CaH chemical shifts of [¹⁵N(Gly0)]PKIα were analyzed by the Chemical Shift Index (Wishart et al., 1992). Briefly, the measured CaH chemical shift values in ppm were subtracted from the tabulated values for each residue. If the difference was greater than 0.1 ppm, a value of +1 was assigned to the residue. If the difference was more negative than −0.1 ppm, a value of −1 was assigned. A value of 0 was assigned to all other residues. A local density of three or more ±1 indices over 70% of the residues in a stretch is indicative of a β-strand, and a local density of 4 or more −1 indices over 70% of the residues in a stretch is indicative of an α-helix. Termination of secondary structures can be identified by a chemical shift index of the opposite sign or by two consecutive zeros.

Experimental constraints

NOE constraints were derived from the NOESY spectra at 200 ms mixing time and applied as 3.0, 4.0, and 4.7 Å upper limits for strong, medium, and weak NOEs, respectively. Pseudo-atom corrections were added (Wüthrich et al., 1983). Distance constraints for hydrogen bonds were not introduced. Values of JaN coupling constants listed in Figure 2 were directly used in the final refinement stage.

Computer modeling methods

Modeling of [Gly0]PKIα(1–25) and the peptide stretch Ser34–Lys48 of PKIα(26–75) was performed by distance geometry and simulated annealing using the X-PLOR program Version 3.8 (Brünger, 1992) on an HP 735 work station and is based on the original work of Clore and Gronenborn and others (Nilges et al., 1988a, 1988b). Distance geometry was applied to a subset of atoms (Ca, Ha, N, NH, C, CB, and Cy) to generate 30 substructures. Simulated annealing was done by 3 ps molecular dynamics at 2,000 K (where the force constants were increased in small steps), an annealing step consisting of 5 ps dynamics during which the temperature was decreased from 2,000 to 100 K in 50 K steps, and 200 cycles of Powell minimization with constraints. The refinement protocol consisted of 10 ps dynamics during which the temperature was decreased from 1,000 to 100 K in 50 K steps and 200 cycles of Powell minimization with constraints. The structures were further refined by applying 200 cycles of Powell minimization and 20 ps dynamics at 300 K, where an electrostatic energy term was introduced. Finally, direct J-coupling refinement was done by applying 5,000 cycles of Powell minimization introducing a JaN-coupling error term (Kuszewski et al., 1995). Structures were selected using the following criteria: (1) no violation of distance constraints greater than 0.5 Å, and (2) good geometry (ca. RMSD from ideality for bonds or angles less than 0.01 Å or 2°, respectively). An average structure was obtained by fitting backbone atoms from residue 1 to 13 for [Gly0]PKIα(1–25) and from residue 35 to 44 for PKIα(26–75). The average RMSD value for the superimposition of these backbone stretches was 1.11 and 1.30 Å, respectively. Both average structures were further minimized by applying 1,000 cycles of Powell minimization.

Ensemble-averaged refinement

The protocol used is similar to the one described by Bonvin and Brünger (1995), starting from the averaged structures of
Supplementary material in Electronic Appendix

Table I. Chemical shift assignments for PKIα and the two PKIα peptides at 2°C (file name NMR assignments). Nitrogen chemical shifts were referenced to external 15NH4Cl (1.85 M) in 1 M HCl at 29.43 ppm relative to liquid ammonia and proton chemical shifts were referenced to the 1H2O signal at 5.14 ppm. (A) Chemical shift assignments for 15N labeled PKIα at pH 6.6. (B) 1H chemical shift assignments for [Gly0]PKIα(1–25) at pH 4.1. (C) 1H chemical shift assignments for PKIα(26–75) at pH 7.1.

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