A Selective NMR Probe to Monitor the Conformational Transition from Inactive to Active Kinase

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Abstract
Kinases control many aspects of cellular signaling and are therefore therapeutic targets for numerous disease states. Monitoring the conformational changes that drive activation and inactivation of the catalytic kinase core is a challenging experimental problem due to the dynamic nature of these enzymes. We apply [13C] reductive methylation to chemically introduce NMR-active nuclei into unlabeled protein kinases. The results demonstrate that solution NMR spectroscopy can be used to monitor specific changes in the chemical environment of structurally important lysines in a [13C]-methylated kinase as it shifts from the inactive to active state. This approach provides a solution based method to complement X-ray crystallographic data and can be applied to nearly any kinase, regardless of size or method of production.

Disciplines
Biochemistry, Biophysics, and Structural Biology | Molecular Biology

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A Selective NMR Probe to Monitor the Conformational Transition from Inactive to Active Kinase

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Supporting Information

ABSTRACT: Kinases control many aspects of cellular signaling and are therefore therapeutic targets for numerous disease states. Monitoring the conformational changes that drive activation and inactivation of the catalytic kinase core is a challenging experimental problem due to the dynamic nature of these enzymes. We apply [13C] reductive methylation to chemically introduce NMR-active nuclei into unlabeled protein kinases. The results demonstrate that solution NMR spectroscopy can be used to monitor specific changes in the chemical environment of structurally important lysines in a [13C]-methylated kinase as it shifts from the inactive to active state. This approach provides a solution based method to complement X-ray crystallographic data and can be applied to nearly any kinase, regardless of size or method of production.

Kinase inhibitor development has driven significant investment in understanding the precise structural features that control kinase activity. Most of our knowledge regarding kinase structure comes from X-ray crystallography, a technique that provides spectacular views of the molecular determinants that control the catalytic activity of these enzymes. Despite its power to resolve atomic level details, X-ray diffraction captures static structural snapshots; the intermediates along the trajectory of a conformational transition are often lost. Moreover, use of X-ray crystallography to solve numerous drug or ligand bound structures can be laborious or not feasible. To complement and enhance available X-ray derived structural information, solution-based techniques must continue to be developed to adequately interrogate kinase structure–function relationships.

Here we report a simple and rapid NMR approach to assess the conformational preferences of any kinase in solution. Using [13C]-labeled formaldehyde, we have used reductive methylation chemistry to introduce a spectroscopic probe into the active site at the β3 strand lysine of a model kinase. The chemical environment surrounding the β3 lysine changes on transition from inactive to active kinase, providing a direct reporter of the activation trajectory. The method can be applied to any kinase regardless of expression system and creates a novel platform to study kinase regulatory mechanisms under a range of solution conditions. In addition to providing insight into how exogenous proteins and/or second messengers affect kinase regulation, the method can be used to directly measure the effect of disease-causing mutations or small molecule modulators on the conformational preferences of the kinase active site.

Src tyrosine kinase was chosen as a model system for NMR method development due to its well understood regulatory mechanism. The Src domain structure consists of two Src Homology domains, SH3 and SH2, a catalytic kinase domain and a C-terminal tail containing a regulatory tyrosine (Tyr527) (Figure 1a). In the autoinhibited conformation of Src, phosphorylated Tyr527 binds to the Src SH2 domain intramolecularily and the Src SH3 domain interacts with the linker spanning the SH2 and kinase domains (Figure 1b). Dephosphorylation of Tyr527, and/or exogenous ligand binding to SH3 and SH2, liberates the kinase domain from the conformational restraints imposed by the SH3/2 domains, and the kinase domain shifts to the active conformation (Figure 1b). It is also established that the isolated kinase domains of Src family members (lacking the SH3-SH2 region) are active.

Kinases share the β3 strand lysine in the N-lobe that protrudes toward the active site, coordinates ATP, and forms a salt bridge with the conserved glutamate on the C-helix. In crystal structures of the autoinhibited form of Src, the C-helix is out of the active site, and the distance between the Lys ε-amino group and the Glu carboxylate group (Lys295/Glu310) is 14 Å (Figure 1c). In structures of active Src, the C-helix abuts the active site of the kinase domain leading to a short, 3.7 Å, distance between the Lys295 and Glu310 side chains (Figure 1d). The chemical environment of Lys295 differs between

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active and inactive conformations, and NMR chemical shift can therefore serve as a reporter of kinase activation status.

The $\epsilon$-NH$_2$ protons of lysine are not good NMR probes due to rapid exchange with water and resulting unfavorable relaxation properties. However, if the lysine side chain is modified in a manner that introduces a nonexchangeable NMR probe and retains the electrostatic properties of the native side chain, the conformational transitions between active and inactive kinases could be monitored in solution. Protein reductive methylation results in two methyl groups covalently attached to the terminal $\epsilon$-NH$_2$ of lysine side chains as well as the amino terminal $\alpha$-NH$_2$ (Figure 2a, inset). The positive charge on the lysine side chain is maintained, and so ion pair interactions are largely maintained, and so ion pair interactions are largely degenerate and likely correspond to closely spaced methyl groups attached to solvent-exposed lysines that experience similar chemical environments (Figure 2b). In contrast, several methyl resonances are well-resolved and evident in the spectrum and correspond to the methyl groups of the dimethylated amino-terminus (Supplementary Figure S1e). Systematic mutagenesis revealed a single missing resonance, allowing unequivocal assignment of the dimethylated amino-terminus ($\alpha$-NH$_2$) (Figure 2b,e). The same mutagenesis approach was used in the activity assays (Supplementary Figure S2a). There is ample precedence, most notably in the pseudokinase literature, for catalytically incompetent kinases being able to populate both active and inactive conformations. Moreover, inactive kinases are often employed for biophysical experiments, and so, despite the loss of catalytic activity for the methylated kinase, we proceeded with analysis using NMR spectroscopy.

Comparison of this two-dimensional NMR data set with that of unmethylated, $^{15}$N-labeled Src kinase domain shows that almost all resonances overlap for the two proteins, and only a small subset of peaks resonate at different frequencies, consistent with the overall kinase domain fold remaining intact following lysine methylation (Supplementary Figure S1e).

Kinase activity of methylated Src was dramatically reduced compared to unmodified enzymes (Supplementary Figure S2a). The loss of activity is likely due to impaired ATP coordination in the active site, since in addition to its interaction with Glu310, the $\beta$ strand Lys295 coordinates the $\alpha$- and $\beta$-phosphates of ATP. Indeed, addition of the stable ATP analogue, AMP-PNP, to $^{15}$N-labeled samples of unmodified and methylated Src KD shows loss of binding to the methylated protein at concentrations that match the ATP concentration used in the activity assays (Supplementary Figure S2b,c). There is ample precedence, most notably in the pseudokinase literature, for catalytically incompetent kinases being able to populate both active and inactive conformations. Moreover, inactive kinases are often employed for biophysical experiments, and so, despite the loss of catalytic activity for the methylated kinase, we proceeded with analysis using NMR spectroscopy.

We acquired a $[^1H,^{13}C]$ HSQC spectrum for the $[^{13}C]$-methylated Src SH3SH2KD protein (Figure 2a). This experiment measures the $^1$H and $^{13}$C resonance frequencies of each $^{13}$C $\text{C}^\text{H}$ functional group. A number of $^1$H-$^{13}$C crosspeaks are evident in the spectrum and correspond to the methyl resonances of the dimethylated lysine side chains and to concentrated buffer components due to $^{13}$C at natural abundance. Based on previously published work, the dimethyl resonances are centered at a $^{13}$C chemical shift of ~45 ppm. The dimethyl peaks that resonate downfield in the $^1$H dimension are largely degenerate and likely correspond to methyl groups attached to solvent-exposed lysines that experience similar chemical environments (Figure 2b). In contrast, several methyl resonances are well-resolved and shifted upfield, suggesting the presence of unique lysine side chain environments within Src.

Assignment of the $\beta$ strand lysine (Lys295 in Src) was accomplished by mutation of Lys295 to methionine in the Src SH3SH2KD protein. Methylation of the mutant protein followed by acquisition of the $[^1H,^{13}C]$ HSQC spectrum revealed a single missing resonance that is assigned to Lys295 in the original spectrum (Figure 2b,c). Additional peaks are shifted upfield in the $^1$H dimension, suggesting that at least one other lysine residue exists in a unique environment. Systematic mutation of candidate lysines, followed by methylation and acquisition of $[^1H,^{13}C]$ HSQC data for each mutant, led to assignment of the $^1$H-$^{13}$C cross peak at 2.25 and 42.5 ppm as dimethylated Lys315 (Figure 2b,d). The dimethylated $\alpha$-NH$_2$ was assigned by introducing a thrombin cleavage site between the N-terminal hexaHis tag and the first residue of Src SH3SH2KD. The methylated protein was then treated with thrombin and further purified using a Ni resin to remove uncleaved protein and the resulting His tag peptide. The $[^1H,^{13}C]$ HSQC spectrum for Src SH3SH2KD lacking the N-terminus reveals a single missing resonance, allowing unequivocal assignment of the dimethylated amino-terminus ($\alpha$-NH$_2$) (Figure 2b,e). The same mutagenesis approach was used to assign the corresponding resonances in the $[^1H,^{13}C]$ HSQC spectrum of Src KD (Supplementary Figure S3).
The resolved methyl resonances of Lys295 and 315 provide two separate probes within the Src SH3SH2KD protein to monitor conformational changes during the course of Src activation. Lys315 is one and a half turns away from Glu310 on the C-helix and in the autoinhibited Src structure projects away from the N-lobe of the kinase domain toward the linker between SH2 and kinase domains, making extensive contacts with Trp260 (Figure 3a). Src activation leads to a large shift in the position of the SH3/SH2 domains and the SH2-kinase linker region (Figure 3b). Peptide ligands that target the Src SH3 and/or SH2 domains compete with the autoinhibited form and activate the Src kinase.28 We titrated two peptides, VSLARRPLPPLP and pYEEIE (ligands for the Src SH3 and SH2 domains, respectively), into the NMR sample containing [13C]-methylated Src SH3SH2KD (Figure 3c,d). Addition of increasing concentration of the SH3 ligand causes spectral changes; specifically, the peak corresponding to Lys295 disappears over the course of SH3 ligand titration, due to line broadening and/or chemical shift change that results in overlap with the neighboring peak (Figure 3c). The Lys315 methyl peak exhibits slow exchange behavior as SH3 ligand concentration increases (Figure 3c). Saturation with SH3 peptide ligand results in the emergence of a new peak at a 1H frequency of 2.28 ppm and complete loss of the original Lys315 signal present in the spectrum of free Src SH3SH2KD.

In a separate titration, the SH2 peptide ligand, pYEEIE, was added to the methylated Src SH3SH2KD sample. Stepwise addition of the pYEEIE peptide causes no spectral change over a range of ligand concentrations (Supplementary Figure S4). This result can be reconciled with the fact that we are using a modified Src construct that contains a high affinity tail sequence surrounding pTyr527; the isolated pYEEIE phosphopeptide does not compete with the intramolecular ligand for SH2 binding at the concentrations used in the NMR experiment. We therefore explored whether the pYEEIE ligand might compete with the pTyr527 for binding to the SH2 domain of SH3SH2KD in the context of SH3 peptide ligand. Using the SH3SH2KD sample that is already saturated with SH3 ligand (last panel in Figure 3c), we added increasing concentration of the pYEEIE peptide. In this titration, addition of SH2 ligand results in further chemical shift perturbation of the Lys315 methyl resonance (Figure 3d). The finding that ligand occupancy of the SH3 domain in SH3SH2KD affects accessibility of the SH2 domain in the intact Src protein is reminiscent of crosstalk between regulatory domains explored previously using other methods.29–32

Figure 2. Chemical modification and NMR data acquisition and assignment. (a) [1H,13C] HSQC spectrum of [13C]-methylated Src SH3SH2KD. Dimethylated lysine resonances are labeled “DM,” and NMR signals from buffer components are labeled (identified by acquiring the identical [1H,13C] HSQC spectrum on buffer alone). The inset shows the structure of the protonated, dimethyl lysine epsilon amino group. (b) Select region of the [1H,13C] HSQC spectrum of methylated Src SH3SH2KD. Dimethyl Lys295, Lys315, and the dimethylated amino terminus (αNH₃) are assigned on the basis of mutational data shown in panels c–e. (c–e) Superposition of [1H,13C] HSQC spectra of [13C]-methylated SH3SH2KD (black) and Lys295Met mutant (red) (c); Lys315Arg mutant (red) (d); and SH3SH2KD lacking the amino-terminus (e). In panels c–e the missing resonance in the mutant spectrum is indicated by the arrow.
Upon saturation of SH3SH2KD with both SH3 and SH2 ligands, the dimethyl lysine region differs dramatically from the [1H,13C] HSQC spectrum acquired for free SH3SH2KD (Figure 3c,d (bolded spectra)). Superposition of the SH3/SH2 ligand-saturated SH3SH2KD spectrum with that of the isolated Src kinase domain (KD) shows an exact correspondence in the resolved region (Figure 3e). This finding provides direct evidence that SH3 and SH2 ligand binding results in release of the Src kinase domain, which then adopts the active conformation present in the free kinase domain. Finally, we subjected the ligand-saturated SH3SH2KD sample to repeated dialysis to remove peptide ligands and determine the reversibility of the conformational shift. Acquisition of a [1H,13C] HSQC spectrum following dialysis shows that the resonance belonging to Lys315 reappears at its original chemical shift (Figure 3d), suggesting that reduced peptide ligand binding to the SH3 and SH2 domains leads to a conformational shift toward the inactive conformation.

The proof-of-principle experiment in Figure 3 demonstrates that NMR spectroscopy, using [13C]-methylated lysine side chains as probes, reports on the conformational ensemble of the Src kinase domain under a range of conditions. In light of the inconvenient exchange regime/chemical shift degeneracy of the Lys295 resonance, the well-resolved Src Lys315 peak provides a convenient, alternative signal that monitors the complete conformational transition between inactive and active
Src (Figure 3c,d). While Src Lys315 is not strictly conserved across kinase families, we wondered whether other kinases would similarly exhibit multiple resolved dimethyl lysine resonances that, in the event of unfavorable spectral properties for the conserved β3 lysine, could be used to monitor the complete conformational transition between inactive and active kinase. To this end, we expressed and purified two additional kinases, Btk and Csk, and subjected these proteins to reductive methylation. Acquisition of [1H,13C] HSQC spectra for methylated Btk and Csk (Figure 4a,b) shows that these kinases, like Src, exhibit multiple nondegenerate dimethyl resonances. Moreover, a comparison of the isolated Btk and Csk kinase domains to their corresponding multidomain proteins (SH3-SH2-kinase domain constructs) (b). Non-degenerate resonances are indicated with a dashed box. (c) Btk kinase domain drug titrations. A 100 μM sample of [13C]-methylated Btk kinase domain (Tyr551 is mutated to Glu to correspond to protein used in the reported crystal structure35) was titrated with either B43 (top row) or dasatinib (bottom row) at the indicated molar equivalents. [1H,13C] HSQC spectra shown are acquired before addition of drug or at each titration point. The Btk kinase domain also carries the Y617F mutation that facilitates bacterial expression.39

In conclusion, NMR data acquired using a lysine methylated sample provide a spectral signature of the Src kinase activation/inactivation trajectory. The data report on the conformational ensemble that is present in solution, enabling the identification of intermediates along the course of kinase activation. We suggest this method will allow investigators to gain insight into the conformational ensemble of any pharmaceutically important, full-length protein kinase in solution, many of which rely on eukaryotic expression systems for production, are difficult to isotopically label, and show limited yield. The approach can be used to assess how allosteric and direct interactions, between a
kinase and its binding partner, be it a substrate, a regulator, or an adaptor protein, drives the conformation of the kinase domain toward the active or inactive state. Similarly, the method provides a rapid analysis of conformational status of a kinase upon inhibition with a small molecule, complementing previous applications of 13C NMR spectroscopy to kinase drug discovery. Finally, insight into how specific disease-causing mutations within or outside of a kinase domain affect conformational preferences of the catalytic site can also be gained with this method. Overall, this method promises to provide deeper insight into the molecular basis of kinase regulatory mechanisms.

■ METHODS

**Protein Production.** Src proteins were coexpressed with YopH and purified as described.12 15-Labeled proteins were expressed in minimal media as described previously.34 Reductive methylation followed previous published protocols.34 All mutations were introduced using site-directed mutagenesis kit (Strategene) and verified by sequencing at the ISU DNA Synthesis and Sequencing Facility.

**Characterization of Methylated Kinases.** For trypsin digestion, 15 μL of 50 mM NH4HCO3 and 1.5 μL of 100 mM DTT were added to 10 μL of 0.8 mg/mL kinase sample, and the volume was adjusted to 27 μL with H2O. The mixture was incubated at 95 °C for 5 min, and 3 μL of 100 mM iodoacetamide was added and incubated in the dark at RT for 20 min. Next, 1 μL of sequencing grade modified trypsin (Promega) at 0.1 μg/μL was added and incubated for 3 h at 37 °C. Digested samples were analyzed by MALDI-TOF and MS/MS using Q-Star XL quadrupole-TOF tandem mass spectrometer (ABI) in the Protein Facility at Iowa State University. Initial velocities for Src KD and SH3SH2KD (methylated and unmethylated) were measured using a generic peptide substrate, poly(Glu,Tyr), as previously described.34

**NMR Spectroscopy.** NMR spectra were acquired at 25 °C on a Bruker AVII700 spectrometer with a 5 mm HCN z-gradient cryoprobe operating at 1H frequency of 700.13 MHz using standard protocols (Bruker pulse program hsqcetgpsp2 and troys3ghpshs19). Sample concentration was 135 μM ([13C]-methylated Src SH3SH2KD and KD) or 200 μM (methylated and unmodified [13C]-Src KD) in the following buffer: 50 mM bicine pH 8.0, 100 mM NaCl, 2 mM DTT, 5% glycerol, and 0.02% NaN3. SH3 ligand VSLARRPLPPLP was from Tocris Bioscience. The 13C reductively methylated SH3SH2KD titration sample was dialyzed against 500 mL of buffer (50 mM bicine pH 8.0, 100 mM NaCl, 2 mM DTT, 5% glycerol, 0.02% NaN3) changed each day for 7 days prior to acquisition of the [1H,13C] HSQC spectrum. For AMP-PNP binding experiments, MgCl2 was added to the [13C]-labeled Src KD samples (methylated and unmodified) to a final concentration of 10 mM. B43 was purchased from Calbiochem and dasatinib was purchased from Selleckchem.

■ ASSOCIATED CONTENT

**Supporting Information** Additional characterization of the reductively methylated proteins used in this study, as well as further information on resonance assignments, ATP binding, and ligand titration. This material is available free of charge via the Internet at http://pubs.acs.org.

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**Notes**

The authors declare no competing financial interest.

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■ REFERENCES