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Chemical Genetic Approach for Kinase-Substrate Mapping by Covalent Capture of Thiophosphopeptides and Analysis by Mass Spectrometry

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Abstract

Mapping kinase-substrate interactions demands robust methods to rapidly and unequivocally identify substrates from complex protein mixtures. Towards this goal we present a method in which a kinase, engineered to utilize synthetic ATP γ S analogs, specifically thiophosphorylates its substrates in a complex lysate. The thiophosphate label provides a bio-orthogonal tag that can be used to affinity purify and identify labeled proteins. Following the labeling reaction proteins are digested with trypsin, thiol containing peptides are then covalently captured and non-thiol containing peptides are washed from the resin. Oxidation promoted hydrolysis, at sites of thiophosphorylation, releases phosphopeptides for analysis by tandem mass spectrometry. By incorporating two specificity gates: kinase engineering and peptide affinity purification, this method yields high confidence substrate identifications. This method gives both the identity of the substrates and phosphorylation site localization. With this information investigators can analyze the biological significance of the phosphorylation mark immediately following confirmation of the kinase-substrate relationship. Here we provide an optimized version of this technique to further enable widespread utilization of this technology.

Keywords

phosphorylation; chemical genetics; analog specific kinase; kinase substrate identification; thiophosphate labeling

Introduction

Members of the kinase superfamily are integral components of many signaling pathways. Kinases play pivotal roles in regulating growth and development and are misregulated in many diseases including cancer. Kinases transduce extracellular signals by altering the functions of substrate proteins through phosphorylating specific sites on these proteins. It is

estimated that a third of all proteins are phosphorylated, and therefore regulated by kinases, however there are only 518 known human kinases (Cohen, 2001; Manning et al., 2002). Therefore many kinases must phosphorylate multiple substrates. The mapping of these relationships is complicated by the shared enzymology of kinases and therefore it is extremely difficult to assign a phospho-site to a specific kinase.

We have developed bio-orthogonal chemical reactions that allow the assignment of a particular phospho-site to a specific kinase (Bishop et al., 2000; Shah et al., 1997). In this approach the active site of the kinase of interest (KOI) is engineered by creating a new active site pocket that allows the kinase to accept a bulky ATP analog (Figure 1A). This engineered “lock and key” provides selectivity for the KOI, as the engineered kinase will accept the N⁶ substituted ATP analog whereas the vast majority of wild-type kinases cannot use these analogs.

In order to identify the substrates of a particular enzyme, many of the most widespread proteomic methods rely on affinity based approaches to enrich for proteins of interest. To apply this approach to kinase-substrate mapping, we developed a modified gamma-phosphate analog (thiophosphate) capture and release strategy for thiophosphate purification. The ATP analog used to identify direct kinase substrates is modified in two ways: a bulky group is added in the N⁶ position providing exclusive recognition by the engineered kinase of interest, and the gamma phosphate is replaced with a thiophosphate moiety (Figure 1). The modified analog-specific (AS) kinase will use these analogs most efficiently, therefore the substrates of the AS kinase will be uniquely labeled with the thiophosphate affinity tag. This tag can then be used to purify and identify the tagged proteins (Figure 1B).

We have recently published two different approaches for the affinity purification of thiophosphorylated substrates (Allen et al., 2007; Blethrow et al., 2008). In this report we provide an updated protocol for the application of these techniques, and demonstrate how they can be used in a complementary manner to rapidly identify the substrates of one kinase from a complex protein mixture.

In the first technique we react the thiophosphorylated proteins with a thiol specific alkylating agent that generates a bio-orthogonal thiophosphate ester. We affinity purify the tagged proteins by using a thiophosphate ester specific antibody 51-8 (Allen et al., 2005; Allen et al., 2007) also see Figure 2B. The antibody is able to distinguish between a thiophosphate ester and a cysteine thioether, and in this way we are able to specifically visualize and identify tagged substrates by immunoblotting (IB) and immunoprecipitation (IP). This approach cannot reliably allow for the identification of the site of phosphorylation because the alkylating agent can prevent normal collision induced dissociation of the modified peptide. Given that it is currently difficult to produce the monoclonal antibody 51-8 in sufficient amounts for IP, we now routinely use this technique for two optimization steps. 1. To determine the optimal N⁶ substituted ATP γ S analog for a KOI and 2. To optimize N⁶ substituted ATP γ S, ATP, and GTP concentrations to achieve the lowest background labeling in cell lysates. Once labeling conditions are optimized, a candidate protein may also be verified as a KOI substrate by immunoprecipitating the candidate substrate and immunoblotting with monoclonal antibody 51-8.

After optimization of labeling conditions using the thiophosphate specific antibody, our second technique provides unambiguously assigned sites of modification on substrates of the engineered kinase. In this technique, the modified peptides are directly affinity purified, and then subjected to tandem mass spectrometry to identify the parent protein and to assign phosphorylation sites (Blethrow et al., 2008) also see Figure 2B. To accomplish the affinity

purification, we first digest the labeled protein lysate and then covalently capture the peptides by reacting the digested peptides with iodoacetyl agarose beads, thus capturing all thiol containing peptides. Unbound peptides are washed away, and the beads are then treated with Oxone to oxidize the sulfur in the thiophosphate ester to a sulfoxide. The peptides linked by a thiophosphate ester bond to the agarose beads are then released by spontaneous hydrolysis, while those linked by a cysteine thioether are retained on the resin. In this protocol we provide the most updated and optimized form of these techniques and provide detailed positive and negative controls to ensure their successful implementation by any lab.

Strategic Planning

Engineering the kinase of interest

To apply this method the KOI must be able to catalytically transfer thiophosphate to substrate proteins, and tolerate the mutation of the normally bulky gatekeeper to a smaller residue. We believe most kinases will be able to use ATP γ S as a phosphorylation cofactor; in preliminary experiments 13 out of 15 kinases were able to use ATP γ S to thiophosphorylate substrate proteins (Allen et al., 2007). To determine whether a KOI can transfer thiophosphate, the thiophosphate ester specific antibody is used to rapidly determine the thiophosphorylation state of a substrate protein after incubation with ATP γ S and the KOI. The first step is therefore to set up an *in vitro* kinase reaction with the KOI, substrate protein and ATP γ S followed by IB with the thiophosphate ester specific antibody (Support Protocol 1). The bioinformatic technique for identifying the gatekeeper as well as the generation of an AS kinase have been described previously (Blethrow et al., 2004; Buzko and Shokat, 2002; Gregan et al., 2007) As described in (Buzko and Shokat, 2002), the kinase database is an online tool that can be used to easily find the gatekeeper residue for their kinase (<http://sequoia.ucsf.edu/ksd/>). After identifying the gatekeeper residue (equivalent to I338 in v-Src) conventional site directed mutagenesis techniques can be used to mutate the gatekeeper to either a glycine (analog sensitive-1, as1) or alanine (analog-sensitive-2, as2). These space creating mutations will allow the kinase to utilize bulky ATP analogs (Buzko and Shokat, 2002; Gregan et al., 2007).

Certain protein kinases lose catalytic activity upon mutation of their gatekeeper residue to Gly or Ala. In such cases, second-site mutations can be used to rescue activity of the kinase [Zhang, Nature Methods 2005]. Identification of second-site suppressor mutations have been successfully executed through bioinformatic analysis or genetic selection. Mutations at a few positions were found to rescue activity in multiple kinases and thus can be considered as top candidate sites for rescue mutations [CZ, KS unpublished results]. For example, one position is immediately N-terminal from the conserved DFG motif (DFG-1) and Ala was found beneficial for kinases activity this position. If the natural residue in a kinase is different from Ala at DFG-1, one can change it to Ala attempting to rescue activity of the kinase. Another position is typically 11 residues N-terminal from the DFG motif (DFG-11) and Leu was found beneficial for kinase activity in general here. If different from Leu, the residue at DFG-11 could be mutated to Leu for rescue of activity. It should be noted that successful rescue mutations varies from kinase to kinase and that there seems to be no universal rescue mutation as of now. Once the AS kinase protein has been expressed and purified, the kinase assay is repeated to identify the N⁶ substituted ATP analog that is preferentially utilized by the AS-KOI (Support Protocol 1a and Figure 1B).

Lysate Optimization

Optimization of N⁶ substituted ATP γ S analog concentration in each specific lysate is also required to ensure the lowest levels of background and therefore the highest confidence in the identified substrates. There are several different parameters required to optimize the

labeling conditions. Although the N⁶ substituted ATP γ S analogs are preferentially used by the engineered AS kinase, at high enough concentrations other kinases can also utilize these analogs. Providing sufficient unmodified ATP and GTP ensure that these kinases are occupied and will therefore decrease the non-specific use of the N⁶ substituted ATP γ S analogs. In addition, each lysate will have a different N⁶ substituted ATP γ S analog that will produce the lowest level of background. The conditions that give the highest signal-to-noise should be determined empirically by varying the levels of AS kinase, N⁶ substituted ATP γ S analog, ATP, and GTP in the presence of AS kinase. Utilizing the thiophosphate ester specific antibody to detect background thiophosphorylated proteins is the best method for lysate optimization. The amount of kinase is critical to the efficiency of the labeling. When using a recombinant form of the AS kinase, up to 1% (by total weight) of the AS kinase can be added. For example 10 μ g of AS kinase in 1 mg of total lysate (100 μ l of 10 mg/mL lysate). If using a transfected cell than the level of kinase can be varied by using different expression vectors, but usually the highest level of expression is necessary to achieving the best signal to noise. Varying the levels of ATP (50–200 μ M), GTP (1–3 mM), and N⁶ substituted ATP γ S analog (100–500 μ M) should give an optimal set of specific conditions in which the lowest background and the highest levels of AS specific labeling is observed (Support Protocol 2).

Generating positive control peptides and protein

We recommend generating a hyper-thiophosphorylated control protein (myelin basic protein (MBP)) and control peptide (GSK3 substrate peptide CREB) which can be accomplished using commercially available materials (Support Protocol 3). The labeled protein will serve as a control for the retention of the thiophosphate modification during the digestion steps, and both controls will serve to ensure that the covalent capture, release, and mass spectrometric analysis of thiophosphorylated peptides are working properly.

During the labeling procedure two negative controls are essential; minus N⁶ substituted ATP γ S analog and minus AS kinase. These controls will provide information about which background substrates are detected and therefore excluded from the list of identified substrates. Using these controls to troubleshoot and optimize each step of the procedure is critical to the success of the protocol.

Basic Protocol: Digestion and Covalent Capture of Thiophosphorylated Peptides

Materials

Solutions and Reagents:

2X Denaturation Buffer: 200 mM NH₄HCO₃, 4 mM EDTA store indefinitely at 22°C

99% pure Urea store indefinitely at 22°C

Siliconized microcentrifuge tubes

1 M *tris* 2-carboxyethyl phosphine (TCEP) in H₂O store for 6 month at –80°C

1 M Dithiothreitol (DTT) store for 6 months at –80°C

Iodoacetyl Agarose Beads (Pierce) store for 6 months at 4°C

200 mM HEPES pH 7.0 store for 4 months at 22°C

50% Acetonitrile 50% 20 mM HEPES pH 7.0 store for 4 months at 22°C

Ziptips (10 μ L and 100 μ L) store for indefinitely at 22°C

Small Disposable Columns (Isolute SPE Accessories Double Fritted Column 120–1021-A and

Single Fritted Res 120-1111-A) Biotage store for indefinitely at 22°C

5 M NaCl store for indefinitely at 22°C

5% Formic acid store for 3 months at 22°C

50% Acetonitrile: 50% H₂O store for indefinitely at 22°C

C-18 Sep Pak (Waters) store for indefinitely at 22°C in dessicator

Oasis SPE (Waters)

0.1 % TFA 50 % Acetonitrile H₂O store for indefinitely at 22°C

0.1 % TFA H₂O store for indefinitely at 22°C

Special Equipment:

QSTAR Elite Mass spectrometer (Applied Biosystems Foster City, Ca.) or another Tandem LC

MS/MS capable mass spectrometer

Step 1. Digestion of labeled protein lysate—In addition to the experimental samples, we recommend the following five control samples:

1. 100 pmol of hyper-thiophosphorylated MBP (see Support Protocol 3)
2. Unlabeled lysate.
3. Unlabeled lysate plus 100 pmol hyper thiophosphorylated MBP.
4. Lysate plus ATP γ S analog minus kinase.
5. An additional covalent capture reaction control (see step 2-1) and the experimental sample: Lysate plus ATP γ S analog plus kinase.

These controls will ensure that all the steps of this protocol are working correctly. The MBP alone and MBP plus lysate will lead to recovery of only one peptide (see Expected Results Section for MW). The lysate controls will provide a list of non-specific substrates.

- 1 Prepare 500 μ L complete (1X) denaturation buffer (8M Urea, 10 mM TCEP, 100 mM

NH₄HCO₃, 2 mM EDTA) fresh by mixing the following:

250 μ L 2X Denaturation Buffer

5 μ L 1 M TCEP

68 μ L H₂O

240 mg Urea

Making the denaturation buffer fresh will limit carbamate formation and will ensure the TCEP is fully reduced. Using a non-thiol reducing agent will reduce non-peptide side reactions with the iodoacetyl beads. Performing the digestion in siliconized tubes will ensure that minimal sample is lost to non-specific adsorption to the tube walls.

- 2 Add 1X denaturation buffer to the labeled lysate or to the sample of labeled MBP such that the final concentration is 6 M Urea (20 μ L of lysate should be dissolved in 60 μ L of 1X buffer) : Incubate the sample at 55°C for 1 hour then cool the sample at room temperature for 10 minutes.

Do not incubate the sample with an alkylating agent such as iodoacetamide. This will react with the thiophosphate group thereby destroying the reactivity of the tag.
- 3 Dilute the sample by adding 50 mM NH_4HCO_3 such that the final concentration of Urea is 2 M and add 1 M TCEP to a final concentration of 10 mM. Then add trypsin in a 1:50-1:10 ratio by weight (1:10 would be 10 μ g trypsin for 100 μ g of lysate protein).

As no iodoacetamide is used in this protocol, high concentrations of urea (2 M) and reducing agent (10 mM TCEP) are used to ensure that proteins in the lysate remain unfolded during the digestion. One easy way to optimize the amount of trypsin and the incubation time is to add BSA to the lysate and then to analyze the digested peptides to determine the sequence coverage. Coverage of at least 60–80% of BSA or around 30–35 peptides should be accomplished.
- 4 After incubation at 37°C for 6 hrs to overnight, acidify the sample by adding 2.5% TFA to attain 0.1% TFA.

The thiophosphate modification is very labile to acid-promoted hydrolysis so do not acidify to 2% TFA as is common.
- 5 Wash a C-18 Sep Pak (or a Oasis HLB) with 10 mL of 0.1% TFA 50% Acetonitrile:H₂O, followed by 10 mL of 0.1% TFA:H₂O. Load the acidified sample to the column and pass it through the column 5 times. Wash the column with 10 mL 0.1% TFA:H₂O. Elute the peptides with 1 mL 0.1% TFA in 50% Acetonitrile:H₂O. Concentrate to near dryness using a speed vacuum.

Different Solid Phase Extraction columns can lead to differential retention of peptides. We have seen success with both types of column (the more hydrophobic C-18 as well as the more hydrophilic Oasis HLB type column) however each lysate will require optimization. Do not take the sample to dryness, as the peptides may be difficult to resuspend into solution. The peptides should be in around 50 μ L after evaporation so that the correct reaction volume can be attained.

Step 2. Covalent Capture of Thiophosphorylated Peptides

- 6 Prepare the Iodoacetyl agarose beads for each sample by pipeting 100 μ L of the 50% resin slurry into a siliconized 0.5 mL tube. Spin on a tabletop centrifuge for 30 seconds at 10,000 x g and remove the supernatant. Wash with 200 μ L 200 mM HEPES pH 7.0, spin and remove supernatant. Add 150 μ L 50% Acetonitrile 50% 20 mM HEPES pH 7.0 and add 5 μ L 5 mg/mL BSA, mix by vortexing briefly and block the beads for ten minutes in the dark.

The beads are supplied in an acidic buffer; washing with 200 mM HEPES pH 7.0 is critical to achieving the correct final pH. The addition of BSA helps by blocking the agarose beads and prevents sample loss by non-specific adsorption. Mixing the slurry for 5 minutes with gentle rocking before pipeting should mix the slurry to homogeneity.

2-1. In this step set up an additional covalent capture reaction by directly adding several pmols of the thiophosphorylated CREB peptide to the beads in 50% Acetonitrile 50% 20 mM HEPES pH 7.0 with BSA. (10–50 pmol should give a very robust signal)

- 7 Adjust the pH of the digested labeled peptide mixture by adding 200 mM HEPES pH 7.0 to a final concentration of 20 mM (for example add 15 μ L 200 mM HEPES pH 7.0, 10 μ L H₂O and 75 μ L acetonitrile to 50 μ L digested peptides for a final volume of 150 μ L). Spin the beads down again, remove the supernatant and add each sample to one tube of bead mixture followed by room temperature incubation in the dark with gentle rocking for 12–16 hours.

Adding the additional labeled peptide control at this point will simplify troubleshooting later. To prevent light from generating radicals within the sample we recommend wrapping the tubes in aluminum foil.

- 8 Prepare one disposable column for washing and elution of each reaction by washing with 1 mL 50% Acetonitrile: 50% H₂O and 1 mL H₂O.

At this point two methods can be used, a column method or a spin method. The protocol is the same in each case however the column method is detailed below. For the spin method all the washes should be performed by addition of the same volume followed by a short 30 second spin at 10,000 x g.

- 9 Add the entire reaction to the top of the column and let it drain into a 1.5 mL microcentrifuge tube. Wash the reaction tube with 100 μ L 50% Acetonitrile 50% 20 mM HEPES pH 7.0 being sure to resuspend all the beads, and add the residual beads to the column.

- 10 Wash the beads by adding 1 mL of each wash liquid to the top of each column. Then push through the wash liquid using a P1000 pipet.

Wash the beads with 1 mL each in the following order:

1. H₂O
2. 5 M NaCl
3. 50% Acetonitrile
4. 5% Formic Acid

If the level of wash liquid drops below the top of the beads the recovery of phosphopeptides is severely reduced. In addition, the order of washing is critical to the success of the washing steps.

- 11 Prepare a solution of 10 mM DTT and add 1 mL to the column, allowing it to drain half way, and then incubating for 10 minutes with the solution in the column.

The addition of DTT to the beads will react with any available iodoacetyl groups left on the beads and will ensure that no I₂ forms during the oxidation step which can iodinate tyrosine residues (unpublished data).

- 12 Prepare a fresh solution of 1 mg/mL Oxone pH ~3.5 in water and add 100 μ L of this solution to the column being sure to resuspend the beads. After draining the column, add an additional 100 μ L and incubate for 10 minutes.

Be sure to resuspend the beads at this point to ensure the complete oxidation of the thiophosphate ester.

- 13 Immediately desalt and concentrate the phosphopeptides with a 10 μ L C-18 ziptip. Wash the ziptip 3X with 0.1 % TFA 50 % acetonitrile H₂O, then 3X with 0.1 % TFA H₂O. Attach the ziptip to the end of a P200 tip and draw the sample through the tip using the P200 on a setting of 100 μ L passing the sample through the ziptip a total of 4 times. Wash 2X with 20 μ L 0.1 % TFA H₂O and elute by washing 3X with 20 μ L of 0.1 % TFA 50 % Acetonitrile H₂O.

Passing the entire sample through the ziptip multiple times ensures the retention of low abundance peptides. The 10 μ L ziptip should fit on the end of a P200 or P1000 tip to simplify the passage of the entire sample across the C-18 column.

- 14 Concentrate the sample to 10 μ L on a speed vacuum and analyze 5 μ L by tandem mass spectrometry. Typically we use a QSTAR ELITE that includes coupled liquid chromatography on a reverse phase C18 column utilizing a 3–32% acetonitrile gradient in 0.1% formic acid. Analyze the phosphopeptides in positive ion mode. MS spectra are acquired for 1 s. For each MS spectrum the two most intense multiply charged peaks are selected for generation of subsequent collision-induced dissociation MS.

This analysis will provide the identity of both the substrate protein and the phosphorylation site.

- 15 Analyze the data by centroiding using Analyst QS software or another appropriate software. Search the MS/MS spectra against the entire UniProt database utilizing Protein Prospector or another appropriate software. Typically we select no constant modifications and allow the following variable modifications: Phospho: Serine, Threonine, Oxidized: Methionine.

Protein Prospector is available free of charge online: <http://prospector.ucsf.edu>

Support protocols

Materials

N-6 substituted Adenosine 5'-[γ -thio]triphosphate (exclusively from Biolog: www.biolog.de), store 10 mM stock in aliquots at -80°C for one year and avoid freeze thaw cycles p-nitrobenzyl mesylate (PNBM) (exclusively from Epitomics: www.epitomics.com) store solid for 1 year at 4°C . A 50 mM stock in DMSO may be stored frozen for one year in one time use aliquots.

Thiophosphate Ester Rabbit Monoclonal Antibody clone 51-8 (1:5,000 in 5% Skim Milk) (exclusively from Epitomics: www.epitomics.com) store for 1 month at 4°C or indefinitely at -20°C

Secondary anti-Rabbit HRP conjugated antibody (Epitomics: www.epitomics.com) store for 1 month at 4°C or indefinitely at -20°C

Materials for Support Protocol 2 (May also require materials from Support Protocol 1)

Plasmid containing KOI suitable for expression of kinase in mammalian cells

Transfection reagents (e.g., Lipofectamine, FuGene)

Phosphate buffered saline (PBS) store indefinitely at 4°C

Cell scrapers

Protease inhibitor cocktail – EDTA (Roche) store for 1 year at 4°C

Phosphatase inhibitor cocktail (Roche) store for 1 year at 4°C

ATP (Sigma) store solid indefinitely at –20°C

GTP (Sigma) store solid indefinitely at –20°C

2X RIPA Buffer (100 mM Tris-HCl pH 8.0, 150 mM NaCl, 1% NP-40, 0.1% SDS) store indefinitely at 4°C

EDTA (Sigma) store solid indefinitely at 22°C

Protein G Magnetic beads (Invitrogen Dynabeads) store for 1 year at 4°C

Magnetic stand which holds 1.5 mL microcentrifuge tubes

Materials for Support Protocol 3 (May also require materials from Support Protocol 1)

Purified active KOI or Plasmid containing KOI suitable for expression of kinase in *E.coli*

Recombinant GSK3β

CREB peptide (KRREILSRRPS (p) YR) store for 1 year at –80°C

Dephosphorylated Myelin Basic Protein (2.5 mg/mL) (Millipore)

10 mM stock in water of Adenosine 5'-[γ-thio]triphosphate tetralithium salt (Sigma) store for 1 year at –80°C

Myelin Basic Protein, Histone H1, or other general kinase substrate (Millipore)

10X HEPES buffered saline (HBS) (200 mM HEPES pH 7.4, 1.5 M NaCl) store for 6 months at 22°C

100X MgCl₂ (1 M MgCl₂) store for 6 months at 22°C

Optional: 33X MnCl₂ (100 mM MnCl₂) store for 6 months at 22°C

5X Laemmli sample buffer store for 1 month at 4°C

OMIX 100 μL Ziptip (Varian)

0.1 % TFA 50 % Acetonitrile store for indefinitely at 22°C

0.1 % TFA H₂O store for indefinitely at 22°C

Tris buffered saline 0.5% Triton TX100 (TBST) store for 1 month at 22°C

Speed Vacuum

Equipment necessary for SDS-PAGE and western blotting (gel running apparatus, transfer apparatus, Nitrocellulose or PVDF)

Support Protocol 1: Kinase Reaction with ATPγS followed by Western Blotting Utilizing Thiophosphate Specific Antibody

Detection of thiophosphorylated substrate proteins is a critical analysis tool to determine whether this technique will work for a KOI in a particular lysate. An in-vitro kinase reaction

with a KOI, substrate and ATP γ S is used here to determine if the kinase will use ATP γ S to thiophosphorylate substrate proteins. If the kinase is able to use ATP γ S to thiophosphorylate substrate proteins the band corresponding to the substrate should be immunoreactive with the thiophosphate ester specific antibody (51-8).

1. On ice prepare a master mix of 1X HBS (or other kinase reaction buffer suitable for the KOI) 1M MgCl₂ (to 10 mM) substrate (or general kinase substrate at 1 mg/mL) and KOI between 50–200 ng per kinase reaction.

Order of addition: H₂O, HBS, MgCl₂ Substrate, KOI. These general conditions will work for most kinases, but check the literature for kinase specific reaction conditions.

2. Label 4 tubes A-D and aliquot 27 μ L of master mix into each tube.

Tubes A-C are controls for non-specific alkylation of cysteine residues by the thiol specific alkylating agent PNBM.
3. Add 3 μ L 10 mM ATP γ S (final concentration of 1 mM) in two of the samples (samples C and D)

Flick the tubes to mix, and then briefly centrifuge to collect the kinase reaction in the bottom of the tube.
4. Let the reaction proceed for 30 minutes at room temperature (or suitable temp for KOI)
5. Add 1.5 μ L of 50 mM PNBM to samples B and D to afford a final concentration of 2.5 mM. The alkylating reaction should proceed for one hour at room temperature. If desired the kinase reaction may be quenched by adding EDTA to 20 mM.

The 50 mM PNBM should be prepared immediately before adding to the samples by dissolving PNBM in DMSO (50 mM is equal to 12 mg/mL).
6. Add 5X Laemmli sample buffer to the samples. Run the samples on a SDS-PAGE gel. Transfer to nitrocellulose or PDVF membrane. Block the membrane for 1 hr with 5% Skim milk (in TBST).
7. Incubate the blot overnight at 4°C with 1:5,000 Thiophosphate Ester Rabbit Monoclonal Antibody, clone 51-8 (in 5% skim milk TBST).

Incubating the blot with the antibody for 1–3 hours at room temperature can also lead to detection of modified proteins with lower sensitivity.
8. After washing 4 times with TBST, incubate blot for 1 hr with anti Rabbit HRP secondary, wash 4 times and then add ECL reagent to visualize PNBM alkylated/thiophosphorylated proteins.

Support Protocol 1a. Identifying optimal N⁶ substituted ATP γ S analog

Each analog specific kinase will have a slightly different substrate specificity. To determine the preferred N⁶ substituted ATP γ S each kinase must be tested with several ATP analogs. Following this protocol to perform a kinase assay with several N⁶ substituted ATP γ S analogs will provide the optimal ATP analog for further studies to identify substrates.

1. Follow support protocol 1, utilizing several N⁶ substituted ATP γ S analogs in place of ATP γ S to identify which analog is used optimally by the AS KOI.

The optimal analog is the one that results in the most substrate labeling as detected by Thiophosphate Ester Rabbit Monoclonal Antibody, clone

51-8. Wild-type KOI should not be able to utilize this N⁶ substituted ATP γ S analog.

Support Protocol 2: Thiophosphorylation of a candidate kinase substrate in cell lysate

The following method is used to determine if a candidate protein is a substrate of a KOI in a complex protein mixture. First the AS KOI is transiently expressed in cells and activated, then cell lysate is prepared and N⁶ substituted ATP γ S is added to initiate the substrate labeling reaction. Thiophosphorylated proteins are alkylated and the candidate substrate is immunoprecipitated using an antibody against that protein. If a band is detected by IB with Thiophosphate Ester Rabbit Monoclonal Antibody then the candidate is a substrate of the KOI.

1. Grow appropriate mammalian cells on 10-cm dishes.
2. Transfect with plasmid encoding WT or AS KOI using transfection reagent of choice, following manufacturer's protocol.
3. Allow cells to grow for 24–48 hours.
4. If appropriate, stimulate cells to activate KOI.
For example, mitogen activated protein kinase ERK may be activated by epidermal growth factor.
5. Remove media and rinse once with 5 mL cold PBS.
6. Using a cell scraper, lyse and scrape cells on ice in 500 μ L 1X RIPA buffer + 1X protease inhibitor + 1X phosphatase inhibitor.
7. Centrifuge at 10,000 \times g for 10 min at 4°C to remove cell debris. Keep supernatant.
8. Add 100 μ M N⁶ substituted ATP γ S, 100 μ M ATP, and 3 mM GTP to each sample (order does not matter).

Recommended concentration ranges for optimizing labeling conditions:

- N⁶ substituted ATP γ S: 50–500 μ M
 - ATP: 50–200 μ M
 - GTP: 1–3 mM
9. Allow for thiophosphorylation of kinase substrates to occur for 20 minutes at room temperature.
 10. Quench reaction with 500 μ L 1X RIPA buffer + 40 mM EDTA (*final [EDTA] = 20 mM*) + 5 mM PNBM (*final [PNBM] = 2.5 mM*), then alkylate with PNBM for 1 hour at room temperature on a rotator.
Follow Support Protocol 1 for preparation of PNBM stock solution.
 11. Immunoprecipitate candidate substrate using antibody according to manufacturer's recommendation. Incubate overnight at 4°C.
Save 20 μ L input for western blotting to check immunoprecipitation efficiency.
 12. Add 40 μ L of 50% slurry of appropriate (protein A or G) magnetic beads to each sample. Incubate 3–4 hours at 4°C on a rotator.

Before use, wash beads once with 1 mL 1X RIPA buffer. Then resuspend in original slurry volume and add to each sample.

13. Using a magnet to collect the beads, wash 5 times w/1 mL 1X RIPA + protease inhibitor + phosphatase inhibitor.

Note: Before first wash, save 20 μ L supernatant for western blotting to check immunoprecipitation efficiency.

14. Resuspend beads in 20 μ L 1X RIPA + protease inhibitor + phosphatase inhibitor +1X Laemmli sample buffer.
15. Heat samples at 95°C for 2.5 minutes.
16. Load the samples onto an SDS-PAGE gel and proceed with IB as described in steps 6–8 of Support Protocol 1, blotting with Thiophosphate Ester Rabbit Monoclonal Antibody.

Support Protocol 3

Making a thiophosphorylated positive control peptide and protein The two control samples, a thiophosphorylated peptide and protein, can be prepared from readily purchasable materials. A kinase reaction is used to make the thiophosphorylated substrates. Mass spectrometric analysis using a MALDI can be used to confirm whether the CREB peptide has been thiophosphorylated. After labeling, the thiophosphorylated myelin basic protein (MBP) can be run on a denaturing polyacrylimide gel followed by an immunoblot using 51-8 (support protocol 1). After this experiment the peptide can be used to evaluate whether the covalent capture reaction is working properly. Labeled MBP can be added to a lysate as a control for the digestion and covalent capture reactions.

- 1 On ice prepare two kinase reactions that each contain:

6 μ L 10X HBS

0.6 μ L 1M MgCl₂ (to 10 mM)

6 mL 10 mM ATP γ S (to 1 mM final) and either 25 μ L 2.5 mg/mL MBP (5 nmol total) or 18 μ L 0.5 mg/ml CREB peptide (5 nmol) and water to a final volume of 60 μ L.

Order of addition: H₂O, HBS, MgCl₂ Substrate, KOI.

- 2 Add recombinant GSK3 β 500 ng and incubate the reaction at room temperature for 4 hours

Incubate for longer reaction times to ensure stoichiometric thiophosphorylation of MBP

- 3a Aliquot the reaction containing MBP into 0.5 mL Eppendorf tubes (5 μ L/tube), snap freeze them in liquid N₂ and store them at –80°C until needed

- 3b Use a large capacity ziptip (OMIX 100 μ L) to desalt the CREB thiophosphorylation reaction, eluting into 200 μ L 0.1% TFA 50% Acetonitrile. Concentrate the sample using a speed vacuum to 60 μ L. Aliquot into 0.5 mL Eppendorf tubes (5 μ L/tube) and snap freeze them in liquid N₂. Store them at –80°C for 3 months or until needed

Utilize the ziptip according to product literature: wash 2X with 0.1% TFA 50% Acetonitrile, 2X with 0.1% TFA H₂O. Pass the sample through the tip 5X being sure not to introduce any air into the tip. Wash

2X with 0.1% TFA H₂O and then elute 2X with 0.1% TFA 50% Acetonitrile.

Support Protocol 4: Chemical synthesis of N⁶ substituted ATP_γS

The synthesis of N⁶ substituted ATP_γS is presented to those interested in making additional analogs. The synthesis will require good knowledge of organic chemistry synthesis procedures and a well-equipped laboratory. This protocol will allow the synthesis of larger quantities of N⁶ substituted ATP_γS analogs, and the synthesis of those that are not available from Biolog. This synthesis can be found in the supplementary information in (Allen et al., 2007)

Materials

Ethanol (Sigma)
Alkyl amine containing desired N⁶ modification
Triethylphosphate (TEP) (Sigma)
POCl₃ (Sigma)
H₃PO₄ (Sigma)
1,8-diazabicyclo[5.4.0]undec-7-en (DBU) (Sigma)
Triethylamine (Sigma)
DOWEX® 501-X8 ion exchange resin
trisodium thiophosphate
3-chloropropionamide
tri-*n*-octylamine
dimethylformamide DMF
diphenyl phosphorochloridate
tri-*n*-butylamine
dioxane
petroleum ether
β-mercaptoethanol
NaOH
CO₂ Tank
PTFE syringe filter (0.45 μm, Acrodisk)
small plastic column
Buchi Rotovapor Model R-200 or equivalent rotary evaporator
Oil bath
Filter paper
Buchner funnel
Acta FPLC (Amersham Biosciences)

Small molecule capable LCMS instrument (Possibly from Waters)

- 1 Dissolve 1 g (3.5 mmol) of 6-chloropurine ribonucleoside in 21 mL of ethanol while stirring.
- 2 Heat to at 95°C then add seven equivalents of the alkyl-amine desired to be the bulky substituent in the N⁶ position (in this case phenethylamine, 2.5 g (21 mmol)) and stir under a reflux condenser for 12 hrs.
- 3 Remove solvent under reduced pressure in Buchi Rotovapor.
- 4 Add 150 mL of Ethanol and place at 4°C for 5 hours.
- 5 Filter the crystals and wash with cold Ethanol to obtain N⁶ Phenethyl (or additional alkyl) Adenosine

Synthesize N⁶ Phenethyl ADP

- 6 Dissolve 2 mmol of the N⁶ Phenethyl (or additional alkyl) Adenosine (0.76 g) in 5 mL of triethylphosphate (TEP)
- 7 Cool to 0°C in an ice bath while stirring.
- 8 Add 0.6 g (4 mmol) of POCl₃ dropwise and continue to stir at 0°C for two hours.
- 9 In a separate container dissolve first dissolve 8 mmol of 1,8-diazabicyclo[5.4.0]undec-7-en (DBU) in 5 mL TEP then add 8 mmol of H₃PO₄.
The order of addition is critical here, first dissolve the DBU then add the H₃PO₄
- 10 Add the entire H₃PO₄ DBU mixture to the mixture of the N⁶ substituted Adenosine and POCl₃.
A yellow solid should form immediately
- 11 Continue to stir for two minutes and then quench the reaction by adding 30 mL of 0.1M triethylammonium bicarbonate (TEAB) and stir for 30 minutes.
Make 2M TEAB by mixing 557.5 mL triethylamine (TEA) with 1,442.5 mL H₂O. Bubble CO₂ through this mixture until the TEA is dissolved.
- 12 Filter the mixture through a PTFE syringe filter (0.45 μm, Acrodisk)
- 13 Load the mixture onto two HiPrep 16/10 QFF anion exchange columns (Amersham Biosciences) in series using a peristaltic pump.
- 14 Run a 80 minute gradient from 100% 0.1M TEAB to 50% 2M TEAB:50% 0.1MTEAB
- 15 Pool the fractions containing the diphosphate
Analyze the fractions by mass spec (LCMS) on a small molecule capable LCMS instrument and pool those fractions containing the diphosphate
- 16 Lyophilize fractions containing the diphosphate
N⁶ phenylethyl ADP calculated mass 530.10 m/z

Synthesize Disodium S-2-Carbamoylethyl Phosphorothioate

- 17 Dissolve 18.6 mmol of trisodium thiophosphate in 28 mL of water
- 18 Dissolve 28 mmol of 3-chloropropionamide in 5.6 mL of DMF and add mixture to trisodium thiophosphate mixture followed by stirring for 48 hours at room temperature.
- 19 Filter the reaction and add 100 mL of ethanol to the filtrate and then cool to 4°C overnight (16 hours)
- 20 Filter the precipitate, wash with ethanol and dry to afford disodium S-2-carbamoylethyl phosphorothioate

Convert disodium S-2-carbamoylethyl phosphorothioate and N⁶ phenylethyl ADP (TEA salt) to pyridinium salt

- 21 Swell 5 mL of DOWEX® 501-X8 ion exchange resin in 50 mL 5% Pyridine (95% water)
- 22 Add the DOWEX® 501-X8 ion exchange resin to a small plastic column and let the 5% pyridine run through until the pH is greater than 10.
When the pH is above 10 all of the sites on the resin are occupied with pyridine.
- 23 Dissolve 2.0 mmol of disodium S-2-carbamoylethyl phosphorothioate or 0.3 mmol N⁶ phenylethyl ADP (TEA salt) in 10 (or 5) mL 1:1 MeOH: H₂O
- 24 Add the mixture to the top of the DOWEX® 501-X8 ion exchange resin column (pyridinium form)
- 25 Elute with another 10 (or 5) mL 1:1 MeOH: H₂O
- 26 Dry in vacuo on a Buchi rotary evaporator to afford the pyridinium salt of S-2-carbamoylethyl phosphorothioate and N⁶ phenylethyl ADP

Synthesize O-diphenyl phosphoro S-2-carbamoylethyl phosphorothioate

- 27 Add 20 mL methanol to 2.0 mmol of the pyridinium salt of S-2-carbamoylethyl phosphorothioate
- 28 Add 2 mmol of tri-*n*-octylamine to convert to the mono (tri-*n*-octylammonium) salt.
- 29 Remove solvent in vacuo on a Buchi rotary evaporator. Repeat evaporation (two times) after addition of 10 mL aliquots of dry DMF.
- 30 Dissolve the residue in 14 mL of dry dioxane, then add 2.9 mmol diphenyl phosphorochloridate followed by 3.8 mmol of tri-*n*-butylamine and stir for two hours at room temperature
- 31 Remove solvent in vacuo on a Buchi rotary evaporator. Add 20 mL ether and after dissolution add 40 mL warm petroleum ether.
- 32 Place mixture at 4°C for 30 minutes until a higher density oily mixture separates and can be decanted into an additional roundbottom flask
- 33 Dry the oily residue on a high vacuum for 1 hour to afford *O*-diphenyl phosphoro S-2-carbamoylethyl phosphorothioate

Synthesis of N⁶ phenylethyl ATP_γS

- 34 Add 12 mL methanol to the pyridinium salt of N⁶ phenylethyl ADP (from step 26)
- 35 Add 0.32 mmol tri-*n*-octylamine and 0.32 mmol tri-*n*-butylamine until complete dissolution occurs.
- 36 Remove solvent in vacuo on a Buchi rotary evaporator. Dry the residue by repeated (three times) evaporation of 5 mL aliquots of dry pyridine.
- 37 Dissolve residue in 3.3 mL dry pyridine and add this to the dry residue of O-diphenyl phosphoro S-2-carbamoylethyl phosphorothioate (from step 33).
- 38 Stir at room temperature for two hours,
A precipitate will form during this time
- 39 Remove solvent in vacuo on a Buchi rotary evaporator. Add 20 mL 0.2 M NaOH then heat the reaction to 100°C for 10 minutes.
- 40 Quench the reaction by adding 10 ml DOWEX® 501-X8 ion exchange resin (pyridinium form) and 0.4 ml β-mercaptoethanol.
- 41 Filter the reaction and then purify on the two HiPrep 16/10 QFF anion exchange columns (Amersham Biosciences) exactly as in step 13.
- 42 Fractions should be analyzed by LCMS and fractions containing the N⁶ phenylethyl ATP_γS should be pooled and lyophilized.
Redissolving the dried fractions in 0.1M TEAB and another round of lyophilization can be done to concentrate and to produce a white solid.
- 43 After lyophilization the compounds should be assumed to be in the TEA form. To obtain accurate concentrations the absorbance at 280 nm should be compared to several standard solutions made up of the N⁶ modified adenosine.

Commentary

Background Information

Identifying the substrates of a kinase is a critical step towards understanding its biological role. Traditional approaches for kinase substrate mapping depend on knockout experiments followed by phosphopeptide enrichment and mass spectrometry. While useful, this method fails to identify all kinase substrates as there exists redundancy in which a single substrate may be phosphorylated by multiple kinases. Methods for phosphopeptide enrichment include metal affinity chromatography using titanium oxide columns (Trinidad et al., 2008) and identifying new phospho proteins by enriching thiophospho peptides (Kwon et al., 2003). A quantitative mass spectrometry approach can more precisely define the effects of stimulation or knockout of a particular kinase. In an application of this technique, a SILAC labeling approach followed by TiO₂ enrichment was used to determine the effects of growth factors on global phosphorylation status (Olsen et al., 2006). All of these approaches fail to specifically assign a phosphorylation site to a particular kinase but are useful for defining the global changes in phosphorylation and for mapping novel phosphorylation sites.

Our approach allows for the labeling of the substrates of a single kinase in a chemical genetic approach, and can therefore unambiguously assign kinase substrate relationships. Originally the N⁶ modified ATP analogs were radioactively labeled in the gamma phosphate position. Putative substrates were then identified by coupling this approach with affinity tagged libraries. This approach was successfully used to identify Cdk1 and Pho 85 substrates in *S. cerevisiae* (Dephoure et al., 2005; Ubersax et al., 2003). Although this engineered

“lock and key” provides selectivity for a single kinase, the transfer of a radioactively tagged phosphate by the N⁶ modified ATP substrate only assists in visualization and traditional biochemical purification of direct kinase substrates from tagged libraries that are not available in more complex eukaryotes.

Introduction of a bio-orthogonal tag onto the substrate proteins of a particular kinase simplifies the affinity purification and identification of the substrates of any protein kinase in a complex lysate. This approach relies on two separate specificity gates: the kinase is engineered to accept bulky ATP analogs, and the substrates are tagged with a bio-orthogonal phosphate analog. The affinity purification of the tagged substrates also depends on two specificity gates: only thiol containing peptides are pulled down, and of those peptides that are covalently bound to the affinity matrix, only those linked by thiophosphate ester bonds are cleaved off the resin. These specificity elements allow for the purification of very low abundance substrates, while also increasing the confidence in the identification of the identified substrates.

Critical Parameters

Utilization of the N⁶ modified ATP analogs by other kinases can lead to false identification of kinase substrate relationships. It is therefore crucial to fully optimize the labeling reaction. Using a wide range of ATP, GTP, and N⁶ modified ATP analogs during the lysate optimization steps will provide the best conditions for these reactions. Nonspecific binding of peptides to the agarose beads can also lead to the false identification of substrate proteins. Generating several positive and negative control samples will help control for the non-specific purification of peptides.

Several controls are essential for distinguishing background from AS kinase specific labeling. First, it is essential to demonstrate the AS kinase dependence of purification of any phosphopeptides. When labeling is observed, or phosphopeptides are affinity purified from samples that do not contain AS kinase but do contain the N⁶ modified ATP analog, these proteins must be subtracted from any list of putative substrates. When non-phosphorylated peptides are identified in the absence of AS kinase it may indicate that the washing steps are not effective. Increasing the stringency of the wash steps by increasing the volume or incubating the washes for 10 minutes should lower background. Secondly, the positive control thiophosphorylated peptide and protein are very useful for diagnosing the digestion and affinity purification steps. In typical tryptic digests, complex lysates are reduced and alkylated before digestion to ensure the complete digestion of all proteins. The alkylation step with iodoacetamide is incompatible with our technique, thereby complicating the digestion of thiophosphorylated protein lysates. Also the thiophosphate mark is susceptible to acid promoted hydrolysis. It is therefore important to include the hyper-thiophosphorylated MBP as a digestion control to ensure the mark is retained on substrate proteins. A rapid way to determine whether the covalent capture and release reaction is working is by including a separate covalent capture reaction with the CREB peptide followed by MALDI TOF analysis.

Troubleshooting

A trouble-shooting guide is presented in Table 1.

Anticipated results

The application of this technique to appropriately labeled lysates should result in the identification of phosphopeptides. After subtraction of any phosphopeptides found in the control lysates, these peptides should predominantly be the substrates of the AS kinase that was used in the labeling reaction. The location of the phosphorylation mark on the substrate

proteins will also allow the investigator to produce non-phosphorylatable substrates and thus to analyze the biological significance of the phosphorylation mark immediately following confirmation of the kinase-substrate relationship.

Thiophosphorylation of CREB (KRREILSRRPS (p) YR) (1795.84 MH+) by GSK3B should lead to a predominant peak at 1891.986 (MH+). After covalent capture with iodoacetyl agarose beads followed by specific hydrolysis the only product ion present should be a double phosphorylated CREB peak at 1875.986 (MH+).

The hyper-thiophosphorylation of Myelin Basic Protein followed by digestion with trypsin and analysis by LC MS/MS will lead to the detection of a thiophosphorylated peptide 1800.8939 (MH+) NIVT (Thiophospho) PRTPPPSQGKGR or 1587.7951 NIVT (Thiophospho) PRTPPPSQGK or 796.4044 (MH+) NIVT (Thiophospho) PR. After covalent capture with iodoacetyl agarose beads followed by specific hydrolysis the only multiply charged ions present will be the same peptide a phospho group instead of a thiophospho modification, a loss of 16 Daltons. 1784.8939 (MH+) NIVT (phospho) PRTPPPSQGKGR, 1587.8001 (MH+), NIVT (phospho) PRTPPPSQGK, or 780.403 (MH+) NIVT (phospho) PR. In addition, incubating the labeled MBP with the lysate to be used for substrate ID and then performing the covalent capture reaction should yield these same peptides, but no background should be seen as there should be no source of thiophosphate in the lysate for background labeling to occur.

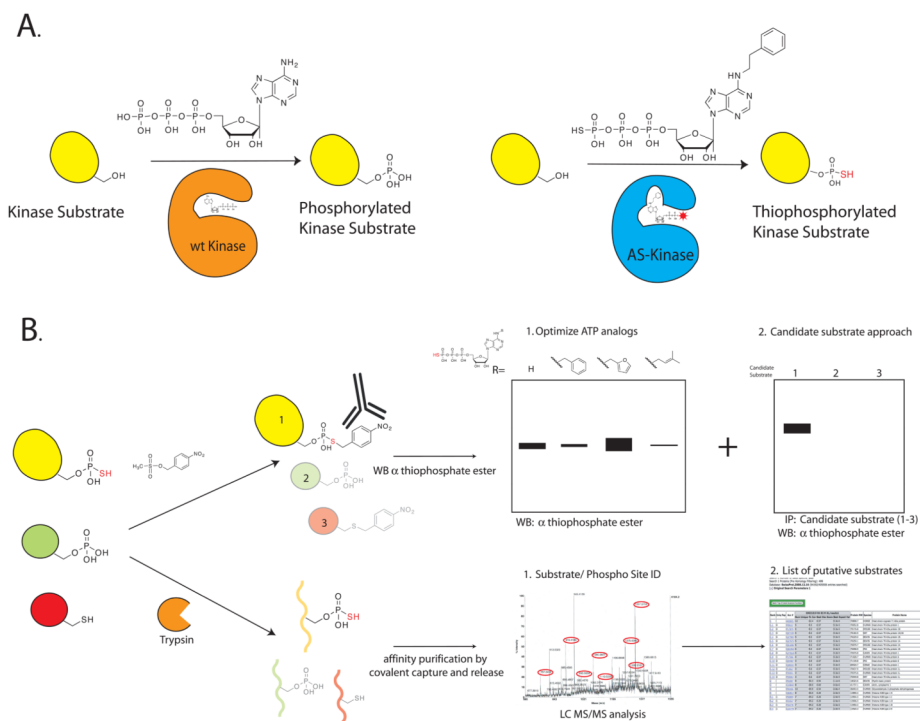
Time considerations

A significant investment of time and resources is required for the design and production of an AS kinase. Once a kinase has been engineered, optimization of labeling conditions and identification of putative substrate proteins should be rapidly achieved in 1–2 weeks. Production of the labeled positive controls should take 1–2 days depending on how long the digestion of labeled MBP is permitted. Once comfortable with labeling, digestion and preparation of covalent capture reactions an experienced investigator should be able to prepare and process several samples in 2–3 days.

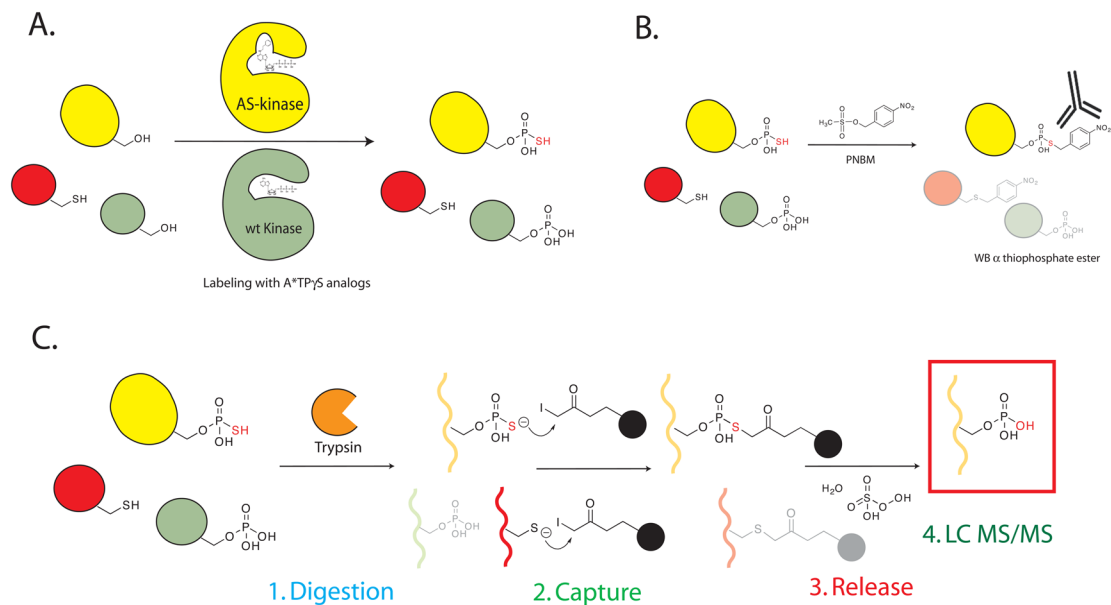
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**Figure 1.**

Thiophosphopeptide purification scheme. A. wt kinase utilizes ATP to phosphorylate its substrates. In contrast, an engineered AS kinase has a new active site pocket that allows it to accept an unnatural bulky ATP analog. This “lock and key” allows the AS kinase to transfer the thiophospho group to its substrates. B. Detail of the two different affinity purification methods: The thiophosphorylated substrates are found in a background of phosphorylated and unlabeled proteins. In the first technique we react the thiophosphorylated proteins with a thiol specific alkylating agent that generates a bio-orthogonal thiophosphate ester. Labeled proteins are detected by using a thiophosphate ester specific antibody. In the second approach the lysate is first digested to generate tryptic peptides. Thiol containing peptides are captured by reaction with iodoacetyl agarose beads. All non-thiol containing peptides are then washed away, and the remaining peptides are treated with Oxone, which releases thiophosphate ester linked peptides by spontaneous hydrolysis.

**Figure 2.**

A. Labeling of a cell lysate by incubating AS kinase and N⁶ substituted ATP γ S analogs generates thiophosphorylated and phosphorylated proteins. B. Alkylating thiol containing proteins generates the bio-orthogonal thiophosphate ester. Thiophosphate ester labeled proteins are detected by utilizing the thiophosphate ester specific antibody (clone 51–8). C. Thiol containing peptides are captured by reaction with iodoacetyl beads. During washing all other non-thiol containing peptides are washed away. The bound peptides are treated with Oxone to convert the thiol group to a sulfoxide. The presence of an electrophilic phosphoester in the thiophosphate ester linked peptides catalyzes the spontaneous hydrolysis of these peptides, and after concentration these peptides are analyzed by tandem LC MS/MS.

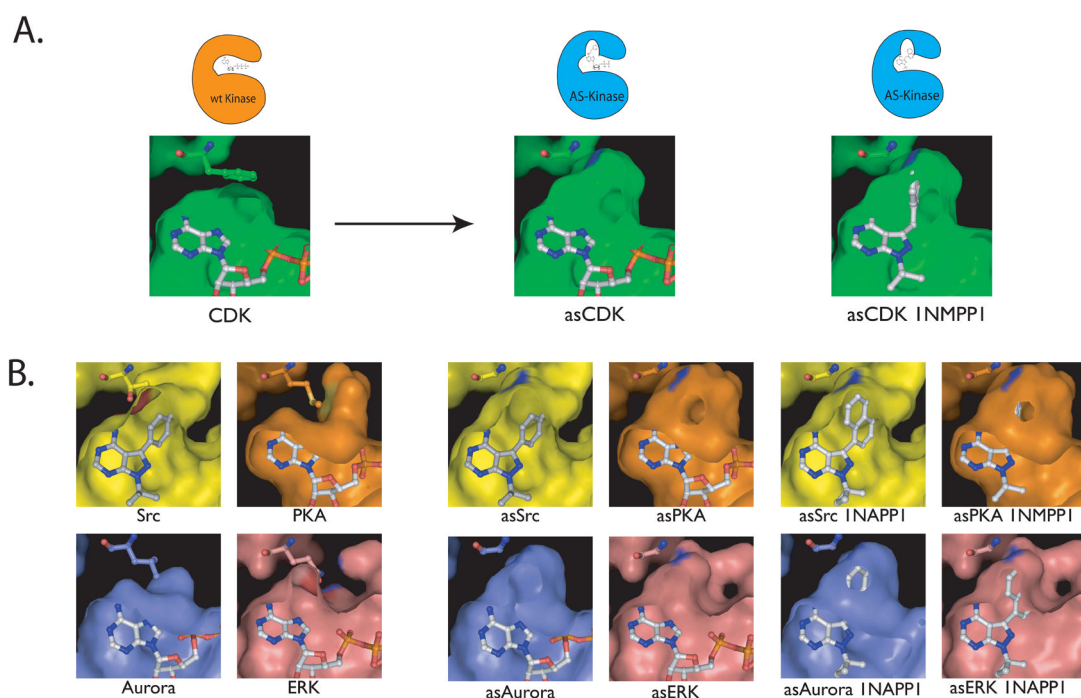
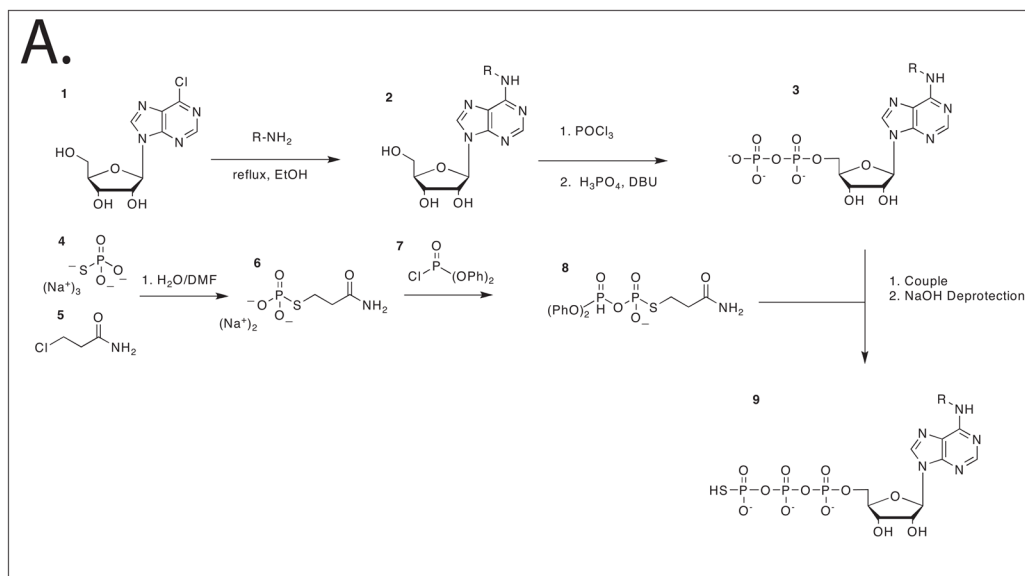


Figure 3.

A. The gatekeeper residue is shown in CDK1 as a phenylalanine above the bound ATP in the CDK1 active site. After mutation to the much smaller glycine, a pocket is opened that allows for a bulky substituent to bind (pictured as asCDK1). A “bumped” (N^6 substituted) ATP analog or inhibitor (in this case INMPP1) can now bind to the engineered kinase by utilizing the extra space above the N^7 position. B. The active sites of SRC, PKA, aurora and ERK are shown along with their gatekeeper residues and the successful mutation of these bulky residues to smaller ones. The mutation of the gatekeeper to a smaller residue creates a similar pocket in each of these four kinases. These kinases are shown with ATP bound or with bumped inhibitors that target these engineered kinases, but not the wild type. The similarity in the engineered pocket among disparate kinases demonstrates the wide applicability of this method.

**Figure 4.**

A. Raw data showing an expected MALDi analysis of thiophosphorylated CREB peptide. Both the singly phosphorylated (KRREILSRRPS(p)YR) (1795.84 MH+) and the thiophosphorylated (KRREILS(thiophosphate)RRPS(p)YR) 1891.986 (MH+) peptide can be seen in the first pane. Analysis of the flow-through after covalent capture with iodoacetyl agarose beads in the second pane shows the complete reaction of the thiophosphorylated (1891.986) peptide, but no reaction with the parent peptide (1795.84 MH+). Analysis of the flow-through with additional HeLa lysate after covalent capture with iodoacetyl agarose beads in the third pane shows many peptides, but the CREB peptide is not seen. B. Following hydrolysis the only peptide present is the double phosphorylated CREB peak at 1875.986 (MH+) demonstrating the specificity of the technique. C. After the covalent capture reaction with digested thiophosphorylated myelin basic protein only one peptide (NIVT(phospho)PRTPPPSQGK) is observed. MSMS analysis of the 1587.8001 (MH+), NIVT(phospho)PRTPPPSQGK (seen as the triply phosphorylated peptide at 524.6 m/z) shows good sequence coverage of this peptide with loss of phosphate from the y10 ion.

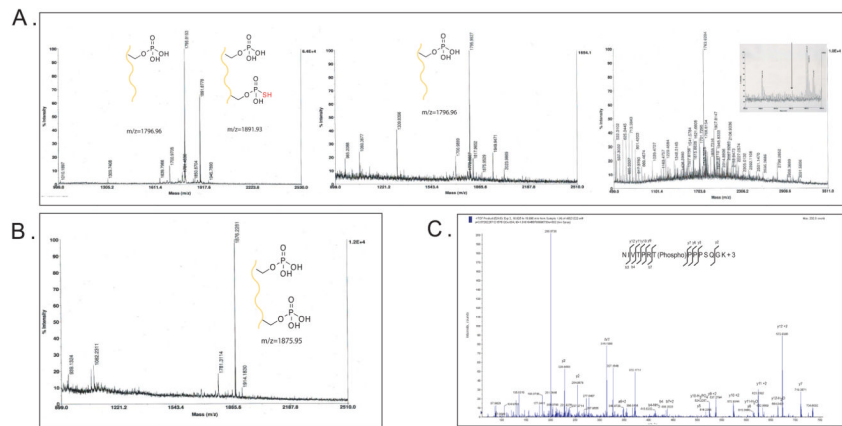


Figure 5.
A. Synthetic scheme for the production of N⁶ substituted ATPγS analogs

Table 1

Problem	Cause	Solution
No thiophosphorylation of substrate seen during in-vitro ATP γ S kinase reaction	Inactive kinase or kinase doesn't utilize ATP γ S	Generate sufficient quantities of pure kinase—check activity with ATP γ S35 kinase assay
	Inactive kinase	Generate a constitutively active kinase
Singly charged polymer or detergent ions seen during mass spectrometric analysis	Concentration of detergents is too high in the lysis buffer	Typically the lysis buffer should only contain low amounts of a non-ionic non denaturing detergent like 0.1% NP-40
No phospho-CREB detected after test covalent capture reaction	pH is too low or high in covalent capture buffer	Check the pH of the flow through after the covalent capture reaction. Ensure the pH of the agarose beads is correctly adjusted by washing the agarose beads carefully with 200 mM HEPES pH 7.0
	Insufficient reaction time for the covalent capture reaction	Incubate the thiophosphopeptides overnight (11–16 hrs, especially if using a complex lysate).
	Insufficient incubation with 10 mM DTT	Incubate with 10 mM DTT for at least 120 minutes following the 5% Formic acid wash step
	Non-specific adsorption of peptides to Eppendorf tubes	Using siliconized Eppendorf tubes and BSA will reduce non-specific adsorption
	Thiophospho CREB levels are too low	Increase thiophospho CREB to see a signal, and then adjust other parameters to diagnose where the sample loss is occurring
No thiophospho peptide is detected after digesting the labeled MBP	Levels of TCEP are too low during digestion	Make a fresh solution of 1 M TCEP before the digestion and add to 10 mM concentration in both the reduction and the digestion steps.
	Levels of urea are too low during digestion	Do not dilute to 1M urea, only dilute to 2M as the additional denaturant will assist in unfolding proteins during the digestion
	Digestion not allowed to proceed long enough	Incubate digestion for a longer time (11–16 hrs)
	Insufficient trypsin added	Add trypsin to 1:10 w/w
Phosphopeptides detected in no AS kinase control	Background kinases are utilizing the ATP analogs	Follow all optimization steps to determine the correct amount of ATP, GTP and N6 modified ATP to use during the labeling
	Background kinases are utilizing the ATP analogs	Identify which background kinases are utilizing the ATP analogs and include inhibitors for these kinases during the labeling reaction
No phosphopeptides detected in the plus AS kinase reaction	Kinase activity is low	Increase the amount of AS kinase in the labeling reaction
	Stoichiometry of labeling is low	Increase the effective concentration of substrates by fractionating the lysate before the labeling reaction
	Stoichiometry of labeling is low	Increase the effective concentration of substrates by isolating specific sub cellular compartments prior to labeling
	N ⁶ ATP analog concentration is too low	Increase the amount of ATP analog in both the negative control and the sample with AS kinase