Kinetic Mechanism of Fully Activated S6K1 Protein Kinase*

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S6K1 is a member of the AGC subfamily of serine-threonine protein kinases, whereby catalytic activation requires dual phosphorylation of critical residues in the conserved T-loop (Thr-229) and hydrophobic motif (Thr-389). Previously, we described production of the fully activated catalytic kinase domain construct, His$_{6}$-S6K1(II)(ΔAID)-T389E. Now, we report its kinetic mechanism for catalyzing phosphorylation of a model peptide substrate (Tide, RRRLSSLRA). First, two-substrate steady-state kinetics and product inhibition patterns indicated a Steady-State Ordered Bi Bi mechanism, whereby initial high affinity binding of ATP ($K_	ext{M}^\text{ATP}$ = 5–6 μM) was followed by low affinity binding of Tide ($K_	ext{M}^\text{Tide}$ = 180 μM), and values of $K_	ext{M}^\text{ATP}$ = 5–6 μM and $K_	ext{M}^\text{Tide}$ = 4–5 μM were expressed in the active ternary complex. Global curve-fitting analysis of ATP, Tide, and ADP titrations of pre-steady-state burst kinetics yielded microscopic rate constants for substrate binding, rapid chemical phosphorylation, and rate-limiting product release. Catalytic trapping experiments confirmed rate-limiting steps involving release of ADP. Pre-steady-state kinetic and catalytic trapping experiments showed osmotic pressure to increase the rate of ADP release; and direct binding experiments showed osmotic pressure to correspondingly weaken the affinity of the enzyme for both ADP and ATP, indicating a less hydrated conformational form of the free enzyme.

A key requirement for higher eukaryotic cells in sustaining prolific capacity is growth regulation, whereby increasing cellular mass and size prerequisite to division derive from coordinate macromolecular biosynthesis. The 70-kDa 40 S ribosomal protein S6 kinase-1 (S6K1)$^2$ is a key enzyme in coordinating cell growth with proliferation, as mitogen, nutrient, and energy status signaling pathways converge to activate S6K1 and initiate protein translation (1–5). Two S6K1 isoforms (accession no. NM003161, α and αII isoforms) are produced from a single gene by alternative mRNA splicing and the use of an alternative translational start site (6). The 525-residue αII isoform contains an N-terminal 23 residue segment that encodes a polybasic nuclear localization motif, whereas the cytoplasmic αII isoform starts at a Met residue equivalent to Met-24 in the αI isoform, and the sequences of both isoforms are identical thereafter.

S6K1 is a member of the AGC subfamily of serine-threonine protein kinases in which amino acid sequences are conserved in a segment of the catalytic kinase domain known as the activation loop or T-loop, as well as in a segment near the C terminus of the kinase domain known as the hydrophobic motif (7). Similar to other AGC kinase family members, catalytic activation of S6K1 minimally requires dual phosphorylation of a critical residue in both the T-loop and hydrophobic motif. For the full-length S6K1αI isoform these residues correspond to Thr-252 and Thr-412, respectively (8), whereas in the S6K1αII isoform the identical residues correspond to Thr-229 and Thr-389 (9). With combined knowledge from available amino acid sequence alignments and x-ray structures, molecular modeling and biochemical testing now provide strong evidence for a common AGC kinase activation mechanism in which the C-terminal phosphorylated hydrophobic motif interacts with a phosphate binding pocket located in the small N-lobe of the kinase (10). This intramolecular interaction acts synergistically with T-loop phosphorylation to stabilize the active conformation, whereby a critical Glu residue in the αC-helix forms an ion pair with the catalytic Lys that functions to position the terminal phosphate of ATP for phosphotransfer in the kinase reaction.

In recognizing the synergistic role of AGC kinase dual site phosphorylation, neither an x-ray three-dimensional structure nor a kinetic mechanism has been reported for any S6K1 isoform or domain construct. This derives largely from the inability to generate fully Thr-229 phosphorylated and activated S6K1. In previous work, we demonstrated by (i) Western analysis, (ii) electrospray ionization-time-of-flight, (iii) Mono Q anion exchange chromatography, and (iv) kinetic assays that the N-terminal His$_{6}$ affinity-tagged catalytic kinase domain construct of the αII isoform of S6K1 (His$_{6}$-S6K1αII(ΔAID)-T389E; residues 1–398) could be generated in its fully Thr-229 phosphorylated most highly active form (250 nmol/min/mg) by baculovirus-mediated expression and purification from S9 insect cells that are co-infected with recombinant phosphoinositide-dependent protein kinase-1, the upstream Thr-229 kinase (11).

Herein, we report that fully activated His$_{6}$-S6K1αII(ΔAID)-T389E catalyzes peptide phosphorylation by a Steady-State
Ordered Bi Bi mechanism, whereby high affinity ATP binding precedes low affinity peptide binding. Rapid chemical phosphorylation and release of phosphopeptide is followed by steady-state rate-limiting steps involving release of ADP. Pre-steady-state kinetic and direct binding experiments showed osmotic pressure to correspondingly weaken nucleotide affinity and increase the dissociation rate, indicating a less hydrated conformational form of the free enzyme.

**EXPERIMENTAL PROCEDURES**

**Materials**—His$_a$-S6K1elII-T389E was generated in its fully Thr-229 phosphorylated and activated form by baculovirus-mediated expression and purification from Sf9 insect cells that were co-infected with recombinant baculovirus expressing the catalytic domain of phosphoinositide-dependent protein kinase-1 (11). Minor amounts of high molecular weight protein impurities observed with His$_a$-affinity-purified His$_a$-S6K1elII-T389E were removed by heparin-Sepharose chromatography. Protein concentration was determined using the Bio-Rad Protein Assay Kit with bovine serum albumin as a standard. Fmoc-$\text{L}$-arginine, Fmoc-$\text{L}$-leucine, Fmoc-$\text{L}$-alanyl, Fmoc-$\text{L}$-serine, and Wang resin were from Advanced Chemtech (Louisville, KY). $\text{N}$-$\text{a}$ Fmoc-$\text{O}$-benzyl-$\text{L}$-phosphoserine (Fmoc-Ser[PO(OBzl)-OH]-OH) was from AnaSpec, Inc. (San Jose, CA). $[\gamma^{32}\text{P}]$ATP, $[8\text{-}^{14}\text{C}]$ATP, and $[8\text{-}^{14}\text{C}]$ADP were from MP Biomedical (Irvine, CA). All other chemicals, salts, and buffers were from Sigma.

**Peptide Synthesis**—The S6K/RSK model peptide substrate (Tide, RRRLS$\text{L}$LA) and phosphorylated peptide product (pTide, RRRL$\text{S}$pSLRA) were prepared by the solid-phase methodology on Wang resin (1.3 mmol/g). Briefly, coupling reactions were carried out under a 3-fold excess of Fmoc amino acids with diisopropycarbodiimide/1-hydroxybenzotriazole as activating agent in dimethylformamide. Intermediate deprotection was achieved with 20% (v/v) piperidine in dimethylformamide for 30 min. After an average coupling period of 1 h, the extent of acylation was monitored by the standard ninhydrin test. In the event of incomplete coupling, the coupling procedure was repeated. Cleavage of the peptide from the resin was performed with trifluoroacetic acid in the presence of 8% anisole, 2% diethyl sulfoxide, 2% $p$-cresol, and 2% thioresol at 0°C for 2 h, which also effectively removed the monobenzyl protecting group of phosphoserine. After removal of trifluoroacetic acid under a stream of nitrogen followed by vacuum, crude peptide was (i) precipitated with diethyl ether, (ii) filter washed with water and diethyl ether, (iii) extracted with 50% (v/v) aqueous acetic acid, and (iv) lyophilized. Crude peptide was purified by semipreparative reversed-phase high-performance liquid chromatography using a Vydac 219TP1010 column (diphenyl, 10 $\mu$m, 10-mm inner diameter $\times$ 250 mm) with 0.1% trifluoroacetic acid in water/0.1% trifluoroacetic acid in acetonitrile as the eluant system. Purified Tide ($\geq$95%) and pTide ($\geq$95%) were confirmed by analytical reversed-phase high-performance liquid chromatography (Vydac 219TP54, diphenyl, 5 $\mu$m, 4.6-mm inner diameter $\times$ 150 mm) and matrix-assisted laser desorption ionization time-of-flight mass spectrometry.

**Two-substrate Steady-state Kinetic Assays**—Steady-state kinetic assays were carried out for His$_a$-S6K1elII$(\Delta\text{AID})$-T389E-catalyzed phosphorylation of the S6K/RSK model peptide substrate (Tide, RRRL$\text{S}$LRA). The residue of the Tide substrate that undergoes phosphorylation is underlined. The 100-$\mu$m Tide phosphorylation reactions were performed at 25°C in 40 mM MOPS buffer, pH 7, containing 0.1% 2-mercaptoethanol, 10 mM MgCl$_2$, and 0.2 mM sodium vanadate. Initial velocities were measured for varying Tide concentrations (1, 2, 3, 5, 10, 25, and 50 $\mu$m) at different fixed concentrations of $[\gamma^{32}\text{P}]$ATP ($\sim$500–1000 cpm/pmol, 1, 2, 3, 5, 10, 25, and 50 $\mu$m). Data collection in this manner provided all information necessary to construct companion reciprocal plots for the two varied substrates. The assays were initiated by addition of 10–30 nm kinase.

For all kinase assays, 20-$\mu$l aliquots were removed at three different times (ranging from 5 to 15 min) and mixed with 20 $\mu$l of 75 mM phosphoric acid and applied to P81 phosphocellulose paper (2 × 2 cm). After 30 s, the papers were washed (3 ×) in 1 liter of fresh 75 mM phosphoric acid for 10 min, then rinsed with 50 ml of acetone, and placed in the hood ($\geq$5 min) to dry. The specific radioactivity of $^{32}\text{P}$-radiolabeled Tide (SA$\text{Tide}$, cpm/pmol) was determined from radioactivity detected by scintillation counting of the known amount of total Tide that was applied to the P81 paper; the micromolar amount of phosphorylated Tide product formed at each time point was determined by reference to the specific radioactivity of $[\gamma^{32}\text{P}]$ATP (SA$\text{ATP}$, $\sim$500–1000 cpm/pmol) and the volume of the aliquot removed for quenching (20 $\mu$l). Initial velocities ($v$, $\mu$m s$^{-1}$) were measured under conditions where total product formation represented $\geq$10% of the initial concentration of the limiting substrate. To better facilitate kinetic comparisons between steady-state and pre-steady-state kinetic results, initial velocities were normalized to enzyme concentration to yield apparent first order rate constants, $k$ (s$^{-1}$) = $v$(E$_{0\text{app}}$).

Control assays were carried out in which either the enzyme or Tide were omitted; these values were always $\leq$5% of the activity measured in the presence of both the lower- and upper-bound concentrations of these reagents (0.5 and 50 $\mu$m $[\gamma^{32}\text{P}]$ATP; 10 and 30 nm enzyme). Control assays containing the enzyme and only the $[\gamma^{32}\text{P}]$ATP substrate were further analyzed to measure ATPase activity. The amount of $[^{32}\text{P}]$inorganic phosphate released from $[\gamma^{32}\text{P}]$ATP was determined by addition of 50 $\mu$l of the reaction mixture to 100 $\mu$l of a quench solution containing a 21% suspension of acid-washed (HCl) activated charcoal in 75 mM phosphoric acid. The quenched solution was mixed and placed on ice for 5 min, which provided for effective removal of the $[\gamma^{32}\text{P}]$ATP nucleotide from solution by the charcoal. The charcoal with bound $[\gamma^{32}\text{P}]$ATP was pelleted by centrifugation for 10 min, and a 60-$\mu$l aliquot of the supernatant was analyzed for $[^{32}\text{P}]$inorganic phosphate. The $^{32}\text{P}$ radioactivity in the supernatant was determined in the absence of enzyme, indicating that the ATPase activity of His$_a$-S6K1$(\Delta\text{AID})$-T389E is significantly lower than activity to the Tide substrate.

**Product Inhibition Steady-state Kinetic Assays**—Diagnostic enzyme inhibition studies were carried out for His$_a$-S6K1$(\Delta\text{AID})$-T389E-catalyzed phosphorylation of Tide exactly as described for two-substrate kinetics. Both ADP and phospho-Tide (pTide, RRRL$\text{S}$LRA) were tested as true product inhibitors. For ADP inhibition, initial velocities were measured
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for varying ATP substrate concentrations (5, 10, 15, 20, 30, 50, and 100 μM) at different fixed concentrations of ADP (0, 5, 10, 15, and 20 μM) at a saturating concentration of Tide (200 μM). For pTide inhibition, initial velocities were measured for varying ATP substrate concentrations (0.5, 1, 2, 3, 5, 10, 15, and 20 μM) at different fixed concentrations of pTide (0, 100, 200, 300, and 400 μM) at a saturating concentration of Tide (200 μM).

Pre-steady-state Kinetic Assays—Pre-steady-state kinetic assays were carried out for His6-S6K1(ΔAID)-T389E-catalyzed phosphorylation of Tide using a KinTek Corp. (Austin, TX) Model RQF-3 rapid-quench-flow apparatus thermostatted at 25 °C. Reaction buffer refers to 40 mM MOPS buffer, pH 7, 0.1% 2-mercaptoethanol, 10 mM MgCl2, and 0.2 mM sodium vanadate. The left and right drive syringes in the rapid quench flow apparatus were filled with reaction buffer, and the middle loop was loaded with 15 μl of reaction buffer containing desig- nated amounts of Tide, [γ-32P]ATP, and ADP; the other sample loop contained 15 μl of His6-S6K1(ΔAID)-T389E enzyme in reaction buffer. The phosphorylation reaction was initiated by mixing the contents of the sample loops and acid quenched after varying reaction times (0.2, 0.3, 0.5, 0.7, 1.0, 1.5, 2.0, and 3.0 s). For each time point, two 20-μl aliquots of each quenched reaction solution were individually applied to P81 phosphocellulose paper (2 × 2 cm), and the amount of Tide phosphorylation was quantified as described for the steady-state kinetic assays. The time courses were performed in triplicate, and the data points represent averaged values.

Three companion sets of pre-steady-state kinetic experiments were carried out in which 2 μM enzyme (concentration after mixing) was reacted with the following three concentrations of other reactants: (i) [Tide] = 200 μM and [ATP] = 5, 10, 20, 50, and 200 μM; (ii) [ATP] = 200 μM and [Tide] = 5, 10, 20, 50, and 200 μM; and (iii) [ATP] = [Tide] = 200 μM and [ADP] = 0, 100, 200, 500, and 1000 μM. Data collection in this manner permitted global kinetic analysis to obtain exact or limiting fit- tness of the microscopic rate constants in the Ordered Bi Bi mechanism (Scheme 1). Control assays were carried out in which the Tide substrate was omitted for enzyme in the presence of 200 μM [γ-32P]ATP. The amount of 32P radioactivity detected on the filter paper was always ≤5% of the measured radioactivity in the presence of Tide, indicating that purified active His6-S6K1(ΔAID)-T389E catalyzes little or no nonspec- ific autophosphorylation during reaction times ≤3 s. In addition, the amount of 32P radioactivity hydrolyzed from [γ-32P]ATP during this time period was shown to be negligible by the charcoal-filtration assay.

Catalytic Trapping Assays—Catalytic trapping assays were carried out at 25 °C using the rapid-quench-flow apparatus exactly as described for pre-steady-state assays except that enzyme was preincubated with ADP prior to mixing with ATP and Tide substrates. The final concentrations after mixing were 2 μM active His6-S6K1(ΔAID)-T389E and 200 μM each of ADP, ATP, and Tide in reaction buffer either in the absence or presence of 5% (w/v) polyethylene glycol with an average molecular weight of 8000 (PEG-8000).

Nucleotide Binding Assays—Nucleotide binding to His6-S6K1(ΔAID)-T389E was measured using a filter binding assay, whereby 0.5 μM enzyme was preincubated at 25 °C in 70 μl of phosphorylation reaction buffer, either in the absence or presence of 5% (w/v) PEG-8000, containing varying concentrations of either [8-14C]ATP (~500 cpm/pmol, and 1, 2, 3, 5, 10, 25, and 50 μM) or [8-14C]ADP (~500 cpm/pmol, and 1.5, 2.5, 4, 7.5, 17.5, 37.5, and 62.5 μM). After 15 min, three separate 20-μl aliquots were removed and applied to three separate wells of a Bio-Rad Bio-Dot microfiltration apparatus with a fitted P81 phosphocellulose paper under vacuum. Radiolabeled nucleo- tide bound to the enzyme was retained by the P81 paper, whereas unbound nucleotide was collected in a vacuum flask. The P81 paper was removed from the apparatus and cut into individual sections. The amount of 14C background radiation was also determined for each nucleotide concentration by control assays carried out in which the enzyme was omitted. The amount of 14C background radiation (~10%) was subtracted from the total radioactivity measured in the presence of the enzyme. The corrected radioactivity was used to calculate the micromolar amount of bound nucleotide, [A]bound, according to the specific radioactivity of the nucleotide and the volume of the aliquot applied to the paper. Because titrated nucleotide exceeded enzyme concentration, the total nucleotide concentration, [A]total, approximated its free concentration, [A]free, for each measurement. Data points represent averaged values from triple measurements.

Data Analysis—Steady-state kinetic data were plotted and fitted using the GraFit 4.0 software (Erithacus Software, UK). The kinetic formulation of an Ordered Bi Bi system, taking into account an irreversible chemical phosphorylation step in the ternary complex (Scheme 1), is fully described in the supplemen- tary material. Initial rates determined in the two-substrate steady-state kinetic studies were globally fitted to Equation S8 (supplemental material), which yielded values for $k_{cat}$, $K_m^{ATP}$, $K_m^{Tide}$, and $K_d^{ATP}$. Secondary plots of $k_{cat(app)}$ determined from direct Michaelis-Menten fits of (i) $k$ versus [ATP] and (ii) $k$ versus [Tide] at different fixed concentrations of the other sub- strate were further analyzed by Equations S11 and S15, respectively, which confirmed the fitted values for $k_{cat}$, $K_m^{ATP}$, and $K_m^{Tide}$.

Initial rates determined in steady-state kinetic ADP and pTide inhibition studies were globally fitted to Equations S23 and S31, respectively, to yield values for $k_{cat}$, $K_m^{ATP}$, and the inhibition constant for the designated product (either $K_m^{ADP}$ or $K_p^{Tide}$). The secondary plot of $K_m^{ATP}$ determined from direct Michaelis-Menten fits of $k$ versus [ATP] at different fixed [ADP] was further analyzed by Equation S25, which confirmed the fitted values for $K_m^{ATP}$ and $K_m^{ADP}$; secondary plots of $k_{cat(app)}$ and $K_m^{ATP}$ determined from direct fits of $k$ versus [ATP] at different fixed [pTide] were analyzed by Equations S32 and S33, respectively, which confirmed the fitted values for $k_{cat}$, $K_m$, and $K_p^{Tide}$. 

\[ \begin{align*}
& E + ATP \overset{k_1}{\rightarrow} E-ATP \overset{k_2}{\rightarrow} E-ATP - Tide \overset{k_3}{\rightarrow} E-ADP + pTide \overset{k_4}{\rightarrow} E-ADP + pTide \overset{k_5}{\rightarrow} E + ADP \\
\text{SCHEME 1}
\end{align*} \]
Pre-steady-state kinetic data were analyzed using the DynaFit 3.28 software (Bio Kin, Ltd., Pullman, WA) (12). The time dependence of phosphopeptide product formation as a function of [ATP], [Tide], and [ADP] was globally fitted to obtain the microscopic rate constants depicted in Scheme 1. As demonstrated in the supplemental material, Equations 1–6 relate values of the microscopic rate constants in Scheme 1 with the kinetic constants determined in steady-state kinetic studies.

\[
    k_{\text{cat}} = \frac{k_{-2}k_{-3}k_{-5}}{k_{-2}k_{-4} + k_{-3}k_{-5} + k_{-4}k_{-5}} \quad (\text{Eq. 1})
\]

\[
    K_{d_{\text{ATP}}} = \frac{k_{-2}k_{-3}k_{-5}}{k_{-2}(k_{-3}k_{-4} + k_{-3}k_{-5} + k_{-4}k_{-5})} \quad (\text{Eq. 2})
\]

\[
    k_{d_{\text{ATP}}} = \frac{k_{-2}}{k_{-1}} \quad (\text{Eq. 3})
\]

\[
    K_{d_{\text{Tide}}} = \frac{k_{-3}k_{-5}(k_{-2} + k_{-3})}{k_{-2}(k_{-3}k_{-4} + k_{-3}k_{-5} + k_{-4}k_{-5})} \quad (\text{Eq. 4})
\]

\[
    K_{d_{\text{ADP}}} = \frac{k_{-5}}{k_{-5}} \quad (\text{Eq. 5})
\]

\[
    K_{d_{\text{Tide}}} = \frac{k_{-3}k_{-4} + k_{-3}k_{-5} + k_{-4}k_{-5}}{k_{-1}k_{-4}} \quad (\text{Eq. 6})
\]

With regard to ADP inhibition, \( K_{d_{\text{ADP}}} \) represents the true dissociation constant, \( K_{d_{\text{ATP}}} \). For pTide inhibition, \( K_{d_{\text{Tide}}} \) closely approximates the true dissociation constant of phosphopeptide product, \( K_{d_{\text{Tide}}} \), under conditions where both \( k_{-3} \) and \( k_{-4} \) exceed \( k_{-5} \).

The nucleotide dissociation constants (\( K_{d_{\text{ATP}}} \) and \( K_{d_{\text{ADP}}} \)) and binding capacity (\( C \)) were obtained from direct fittings of plots of concentrations of bound nucleotide, \([A]_{\text{bound}}\) versus concentrations of free nucleotide, \([A]_{\text{free}}\), to Equation 7.

\[
[A]_{\text{bound}} = \frac{C[A]_{\text{free}}}{K_{d} + [A]_{\text{free}}} \quad (\text{Eq. 7})
\]

**RESULTS**

Two-substrate Steady-state Kinetics—Fig. 1 shows double reciprocal plots of all steady-state kinetic data for titration of active His6-S6K1II(DAID)-T389E with varying concentrations of one substrate at different fixed concentrations of the other substrate. Due to the results obtained in product inhibition studies (see below), analysis of these data was best approximated and described for a Steady-State Ordered Bi Bi system, whereby binding of ATP precedes binding of peptide substrate (Scheme 1). Global fitting of the data to Equation S8 yielded values of \( k_{\text{cat}} = 0.192 \pm 0.002 \text{ s}^{-1}, K_{d_{\text{ATP}}} = 5.1 \pm 0.2 \text{ mM}, K_{d_{\text{Tide}}} = 4.1 \pm 0.1 \text{ microM}, \) and \( K_{d_{\text{ADP}}} = 5.0 \pm 0.4 \text{ microM} \).

To better illustrate this analysis, these data are further given in direct and secondary plots, where both the global fit to Equation S8 and individual direct fits to the Michaelis-Menten equation are displayed and tabulated (supplemental Fig. S1 and Tables S1 and S2).

Product Inhibition Steady-state Kinetics—The pattern of intersecting lines shown in two-substrate steady-state kinetics (Fig. 1) provided little evidence toward the kinetic mechanism of peptide phosphorylation other than ruling out both (i) Ping-Pong and (ii) Rapid Equilibrium Ordered Bi Bi systems. For these mechanisms, respectively, either parallel lines are observed or lines intersect on the y-axis for the second binding substrate. Discrimination between other possible mechanisms (e.g. Steady-State Ordered, Steady-State Random, or (Partial) Rapid Equilibrium Random Bi Bi systems) may be achieved by comparing the effects of product inhibitors in double reciprocal plots constructed for varying one substrate concentration at a fixed concentration of the other substrate.

Fig. 2 shows double reciprocal plots of both ADP and pTide product inhibition steady-state kinetic data for titration of active His6-S6K1II(DAID)-T389E, whereby [ATP] was varied using a fixed [Tide] (200 mM) far exceeding \( K_{d_{\text{ATP}}} \) (4.1 mM). Fig. 2A shows the ADP product to be competitive with respect to the varied ATP substrate, and global fitting of the data to Equation S23 yielded very close approximations of the true values of \( k_{\text{cat}} = 0.190 \pm 0.003 \text{ s}^{-1}, K_{d_{\text{ATP}}} = 5.2 \pm 0.4 \text{ microM}, \) and the inhibition constant \( K_{d_{\text{ADP}}} = 5.5 \pm 0.5 \text{ microM} \). In this case, the measured \( K_{i_{\text{ADP}}} \) equates with the true dissociation constant for ADP product release, \( K_{d_{\text{ADP}}} \) (Scheme 1 and Equation S5).

Fig. 2B shows the pTide product to be uncompetitive with respect to the varied ATP substrate, and global fitting of the
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Figure 2. **Product inhibition steady-state kinetics.** Double reciprocal plots of \(1/k\) versus \(1/[ATP]\) performed at 25 °C. A, ADP product inhibition was measured using (i) varying [ATP] = 5, 10, 15, 20, 30, 50, and 100 μM, (ii) saturating fixed [Tide] = 200 μM, and (iii) [ADP] = 0 μM (○), 5 μM (□), 10 μM (▲), 15 μM (△), and 20 μM (▽). Solid lines were generated using the kinetic constants determined from the global fit of the data to Equation S23 and its reciprocal Equation S23. B, pTide product inhibition was measured using (i) varying [ATP] = 0.5, 1, 2, 3, 5, 10, 15, and 20 μM, (ii) saturating fixed [Tide] = 200 μM, and (iii) [pTide] = 0 μM (○), 100 μM (□), 200 μM (▲), 300 μM (△), and 400 μM (▽). Solid lines were generated using the kinetic constants determined from the global fit of the data to Equation S31 and its reciprocal Equation S34.

Table 1. **Comparison of kinetic constants determined for phosphorylation of Tide by His<sub>6</sub>S6K1(ΔAID)-T389E**

| Constant | Steady state<sup>a</sup> | Pre-steady state<sup>b</sup> | Equation |
|----------|----------------|----------------|----------|
| \(k_{\text{cat}}\) (s<sup>-1</sup>) | 0.18–0.19 | 0.22 | 1 |
| \(K_{\text{m}}^{\text{ATP}}\) (μM) | 4.9–5.5 | 5.9 | 2 |
| \(K_{\text{m}}^{\text{pTide}}\) (μM) | 5.0–5.5 | 5.9 | 3 |
| \(K_{\text{d}}^{\text{pTide}}\) (μM) | 4.1 | 5.1 | 4 |
| \(K_{\text{d}}^{\text{Tide}}\) (μM) | 5.5–6.0 | 5.8 | 5 |
| \(K_{\text{d}}^{\text{Tide}}\) (μM) | 170–180 | 185 | 6 |

<sup>a</sup> These numbers represent the range of values obtained from global and secondary fits of data obtained in both two-substrate and product inhibition steady-state kinetics. A detailed listing of all determined values ± S.E. is given in supplemental Table S1.

<sup>b</sup> These numbers were calculated using the designated equation and microscopic rate constant values in Table 2.

To better illustrate this analysis, the ADP and pTide inhibition data are further given in direct and secondary plots, where both the global fits to Equations S23 and S31 and individual direct fits to the Michaelis-Menten equation are displayed and tabulated (supplemental Fig. S2 and Tables S1 and S3). Table 1 summarizes the range of values obtained for the steady-state kinetic constants obtained from all two-substrate and product inhibition analyses.

Pre-steady-state Kinetics—Fig. 3 shows three companion plots of pre-steady-state time progress curves measured for pTide product formation, which were simultaneously fitted to yield microscopic rate constants of the Ordered Bi Bi mechanism (Scheme 1 and Table 2). The rapid-quench-flow apparatus was used to measure three companion sets of pre-steady-state kinetic data, whereby free enzyme from one sample loop was reacted at 25 °C with the following final concentrations of reactants from the other sample loop: A, [Tide] = 200 μM and [ATP] = 5 μM (○), 10 μM (□), 20 μM (▲), 50 μM (△), and 200 μM (▽); B, [ATP] = 200 μM and [Tide] = 5 μM (○), 10 μM (□), 20 μM (▲), 50 μM (△), and 200 μM (▽); and C, [ATP] = [Tide] = 200 μM and [ADP] = 0 μM (○), 100 μM (□), 200 μM (▲), 500 μM (△), and 1000 μM (▽). The curves are from global fitting of all time progress data to the Ordered Bi Bi mechanism in Scheme 1 with DynaFit (12), which yielded the rate constants given in Table 2.
TABLE 2
Microscopic rate constants for phosphorylation of Tide by His$_6$-S6K1(ΔAID)-T389E

| Reaction step | Rate constant | Value |
|---------------|---------------|-------|
| $E + ATP \rightarrow EATP$ | $k_1$ (m$^{-1}$s$^{-1}$) | $3.8 \pm 0.2 \times 10^4$ |
| $EATP \rightarrow E + ATP$ | $k_2$ (s$^{-1}$) | $0.22$ |
| $EATP + Tide \rightarrow EATP-Tide$ | $k_{12}$ (m$^{-1}$s$^{-1}$) | $\leq 10^4$ |
| $EATP-Tide \rightarrow EATP + Tide$ | $k_{23}$ (s$^{-1}$) | $\geq 190$ |
| $EATP-Tide \rightarrow EATP$ | $k_2$ (s$^{-1}$) | $8.6 \pm 1.0$ |
| $EATP + pTide \rightarrow EATP + pTide$ | $k_3$ (s$^{-1}$) | $\leq 10^5$ |
| $EATP + ADP \rightarrow EATP + ADP$ | $k_4$ (s$^{-1}$) | $0.23 \pm 0.01$ |
| $E + ADP \rightarrow EADP$ | $k_5$ (m$^{-1}$s$^{-1}$) | $4.0 \pm 0.2 \times 10^4$ |

This was used to mix free enzyme from one sample loop with varying concentrations of ATP, Tide, and ADP from the other sample loop. In all cases the final active His$_6$-S6K1(ΔAID)-T389E concentration after mixing was 2 μM.

Figs. 3A (varying [ATP]) and 3B (varying [Tide]) show time progress curves, whereby the final concentration of one substrate was fixed at 200 μM and the other substrate was varied from 5 to 200 μM. Fig. 3 (A and B) shows significant burst kinetics appearing at very high substrate concentrations, indicating that the rate-limiting step occurs after chemical phosphorylation. In addition, the burst amplitude approaches the concentration of the enzyme (2 μM), indicating fully active enzyme.

Further inspection of the data in Fig. 3 (A and B) revealed that burst kinetics required very high concentrations of both [ATP] and [Tide], indicating unusually slow association of the first binding substrate, ATP. This is consistent with the measured values of $k_{cat}/K_m^{ATP} = 3.7 \times 10^4$ M$^{-1}$s$^{-1}$ and $k_{cat} = 0.19$ s$^{-1}$ (Table 1), which approximate the microscopic rate constants for association ($k_{+1}$) and dissociation ($k_{-1}$) of ATP (Scheme 1) under conditions where $K_m^{ATP} \approx K_m^{ADP}$ (Equations 1–3).

Fig. 3C shows time progress curves, whereby the final concentrations of ATP and Tide were fixed at 200 μM and ADP was varied from 0 to 1 mM. In this case, decreasing linear steady-state velocities were observed with increasing [ADP] in accordance with increasing values of $K_m^{ADP}$ (in the absence of [ATP]). In addition, decreasing burst amplitudes were observed with increasing [ADP], reflecting the competition between ATP and ADP for binding to the free enzyme upon mixing. Most significantly, it can be seen that the burst amplitude was reduced ~50% when [ADP] = [ATP] = 200 μM. Making note of the approximately equal values measured for $K_m^{ADP}$ and $K_m^{Tide}$, the ADP association (k$_{+3}$) and dissociation rate constants (k$_{-3}$) must approximate those of ATP (k$_{-3} = k_{-1} \sim k_{cat}/K_m^{ATP}$; k$_{+3} \sim k_{+1} \sim k_{cat}$), strongly suggesting that release of ADP is the rate-limiting step.

Preliminary analysis of the pre-steady state kinetic data to the mechanism in Scheme 1 revealed that good fitted values could not be obtained for $k_{-1}, k_{+2}, k_{-2}, k_{-4}$, and $k_{-4}$. The insensitivity of the rate constants for association and dissociation of Tide (k$_{+2}$ and k$_{-2}$) and pTide (k$_{+4}$ and k$_{-4}$), together with the measured weak inhibition constant of pTide (Table 1, $K_{pTide}^{Tide} = 180$ μM), are consistent with rapid equilibrium peptide binding. Using the measured value of $K_{pTide}^{Tide} = 180$ μM, ratio fixed values of $k_{+4} = 1.8$, 18, 180, and 1800 s$^{-1}$ with $k_{-4} = 10^4$, 10$^5$, and 10$^7$ M$^{-1}$s$^{-1}$, respectively, were assigned and held constant during the fitting analysis. Assuming that pTide and Tide association rates are approximately equal, fixed values of $k_{+2} = k_{-4} = 10^4$, 10$^5$, and 10$^7$ M$^{-1}$s$^{-1}$ were also assigned and held constant during the fitting analysis. Finally, the calculated value of $k_{-1} = k_{cat} = 0.19$ s$^{-1}$ was assigned and held constant.

In all cases where $k_{+2} = k_{-4} \geq 10^6$ (and $k_{+4}/k_{-4} = 180$ μM), global fitting of the pre-steady state kinetic data (Fig. 3) yielded identical values of $k_{+4} = (3.8 \pm 0.2) \times 10^4$ M$^{-1}$s$^{-1}$, $k_{-4} = 8.6 \pm 1.0$ s$^{-1}$, $k_{+5} = 0.23 \pm 0.01$ s$^{-1}$, and $k_{-5} = (4.0 \pm 0.2) \times 10^4$ M$^{-1}$s$^{-1}$ (Table 2). In addition, the fitted values of $k_{-2} = 190 \pm 27$ s$^{-1}$ and 1900 ± 270 s$^{-1}$ when $k_{+2} = k_{-2}$ were correspondingly fixed at 10$^6$ and 10$^7$ M$^{-1}$s$^{-1}$, respectively) yielded $k_{d}^{Tide} = k_{-2}/k_{+2} = 190$ μM, which approximates the measured value of $K_{d}^{Tide}$. In cases where $k_{+2} = k_{-2} \leq 10^5$ such that $k_{+5} \leq 18$ s$^{-1}$, global fitting procedures were increasingly unable to converge, yielding systematic deviation of the fitted lines from the data and surely indicating that release of ADP ($k_{-5} = 0.23 \pm 0.01$ s$^{-1}$) is the rate-limiting step. Thus, limiting lower values were assigned to rate constants for association and dissociation of Tide ($k_{+2}$ and $k_{-2}$) and pTide ($k_{+4}$ and $k_{-4}$) (Table 2). Using the microscopic rate constants in Table 2, Equations 1–6 were used to calculate values of the steady-state kinetic constants; and these values agree well with the values measured in steady-state kinetics (Table 1). According to Equations 1–3, the calculated value of $k_{cat} = 0.22$ s$^{-1}$ was assigned to be the value of $k_{-1}$, because $K_m^{Tide} \approx K_m^{ATP}$ (Table 2). In addition, this kinetic mechanism is consistent with conditions where both $k_{+3}$ and $k_{+4}$ exceed $k_{-1}$ so that the $K_{j}^{Tide}$ (Equation 6) approximates $K_{d}^{Tide} = k_{-1}/k_{-4}$.

Effects of Cosolutes on ADP Release and Nucleotide Binding—"Stressing" cosolutes are known to exert effects on enzymatic equilibrium and kinetic constants by any number of a variety of mechanisms, including (i) specific interactions between the cosolute and enzyme and (ii) changes in solution properties such as water activity, dielectric constant, and viscosity (13). Most typical to enzymatic kinetic studies, a microviscosogen cosolute (e.g. sucrose and glycerol) is used to detect rate-limiting diffusional release of products, whereby $k_{cat}$ is decreased proportionately with increasing solution viscosity (14). To rule out effects from specific interactions, the observed rate decreases should be determined to be the same when using amounts of different microviscogens that produce the same relative viscosity. One further important control involves testing the reaction in the presence of a high molecular weight macroviscosogen (e.g. PEG-8000), which should not affect the rate of diffusion of a small molecule product.

During preliminary steady-state kinetic studies, we made the unusual observation that reaction mixture solutions utilizing increasing amounts of either sucrose or glycerol as microviscosogens increased $k_{cat}$; the small amounts of rate enhancements were nonlinear with measured buffer viscosities. Moreover, this same effect was observed in the presence of much lower amounts of the macroviscosogen PEG-8000. Although the rate of diffusion of ADP away from enzyme was surely decreased in the presence of the microviscogens, it certainly remained faster than a rate-limiting step preceding ADP release; and this rate-limiting step was activated by cosolutes of differing chemical nature (i.e. sucrose, glycerol, and PEG). In addition, we early noted that preparations of His$_6$-S6K1αII(ΔAID)-T389E were
To account for observations (i) that the rates of nucleotide association and dissociation were unusually slow and (ii) that nucleotide binding kinetics and equilibria were sensitive to water activity, the deduced kinetic mechanism of Scheme 1 is sensitive to enzyme inactivation and that stability could be conferred by storage in buffers containing a known protein “stabilizing” agent such as sucrose (12). Hence, the activating effect of cosolute in pre-steady state kinetic and equilibrium binding experiments (Fig. 4) will be discussed in terms of it acting as “protecting” osmolyte.

In cases where the free and ligand-bound forms of a protein are differentially solvated, osmotic pressure will shift the binding equilibrium toward the protein conformation that is in the less hydrated form (15–18). For osmotic pressure studies, PEG-8000 is an optimal “nonbinding” high molecular weight cosolute compared with either sucrose or glycerol on several accounts (15–18). First, it tends to be excluded from the layer of water molecules near the protein surface. As an osmolyte, PEG-8000 reduces the activity of protein surface water by forcing water molecules to migrate from solvent-exposed surfaces, interfacial surfaces, and internal clefts into the bulk solution. In contrast, numerous other small molecule osmolytes are known to exert conformational effects more proximate to the protein surface. Second, PEG-8000 is a very strong osmolyte so that minimal amounts can be used to obtain significant osmotic pressure levels. Finally, small molecule diffusional processes will not be complicated since PEG-8000 behaves as a macroviscogen.

To better probe the nature of the rate-limiting step, the rapid-quench-flow apparatus was used to determine and compare pre-steady-state time progress curves obtained in the absence and in the presence of PEG-8000. As might be expected, the presence of 5% (w/v) PEG-8000 in the reaction buffer exerted no effect on the initial chemical burst for pTide formation. In accordance with steady-state kinetic observations, the slope of the subsequent rate-limiting linear phase increased ~2-fold (Fig. 4A), suggesting that release of ADP generates an enzyme conformation that is less hydrated.

To more directly observe and confirm osmolyte-induced activation of ADP product release, the rapid-quench-flow apparatus was used to determine and compare time progress curves in “catalytic trapping” experiments. In this case, enzyme preincubated with a saturating amount of ADP from one sample loop was reacted with saturating amounts of ATP and Tide from the other sample loop. That the overall process associated with release of ADP is rate-limiting is clearly indicated by complete loss of the burst phase, which became a significant lag phase. Again, the presence of 5% (w/v) PEG-8000 in the reaction buffer increased the subsequent linear steady-state accumulation of product by 2-fold (Fig. 4A).

Finally, a radiometric filter-binding assay was used to measure the effect of PEG-8000 in the reaction buffer on equilibrium binding of both [8–14C]ATP and [8–14C]ADP (Fig. 4B). In the absence of PEG, the enzyme showed similar values of $K_{d}^{ATP} = 5.6 \pm 0.6 \mu M$ and $K_{d}^{ADP} = 5.7 \pm 0.7 \mu M$, which agreed with the values determined in steady-state and pre-steady state kinetics (Table 1). But in the presence of 5% (w/v) PEG-8000, the enzyme exhibited ~2-fold weaker affinities ($K_{d}^{ATP} = 13 \pm 1 \mu M$ and $K_{d}^{ADP} = 15 \pm 2 \mu M$), consistent with the 2-fold increased rate of ADP dissociation measured in pre-steady state kinetics (Fig. 4A). Thus, the weakened nucleotide affinity and correspondingly increased dissociation rate most likely result from osmolyte-induced stabilization of a less hydrated nucleotide-free form of the enzyme. That ADP release is coupled to a rate-limiting conformational transition is further supported by the unusually slow rate constants measured for association and dissociation of both ATP substrate ($k_{-1} = 3.8 \times 10^{4} M^{-1} s^{-1}$ and $k_{1} = 0.22 s^{-1}$) and ADP product ($k_{-5} = 4.0 \times 10^{4} M^{-1} s^{-1}$ and $k_{5} = 0.23 s^{-1}$) (Table 2).
elaborated to that shown in Fig. 5. In specific, the single-step ATP and ADP forward binding reactions (Scheme 1; $K^\text{ATP}_{d}(\text{app}) = k_{+1}/k_{-1}$ and $K^\text{ADP}_{d}(\text{app}) = k_{-5}/k_{+5}$) are now modified to behave as sequential two-step binding reactions, whereby initial rapid equilibrium weak association of enzyme with either ATP ($K^\text{ATP}_{d} = k_{+1a}/k_{-1a}$) or ADP ($K^\text{ADP}_{d} = k_{5a}/k_{-5a}$) is converted to an overall high affinity complex by a fast and favorable conformational change ($K^\text{ATP}_{d} = k_{+1b}/k_{-1b}$ and $K^\text{ADP}_{d} = k_{-5b}/k_{+5b}$). Thus, the degrees to which such nucleotide “clamping” yields apparently higher affinity complexes ($K^\text{ATP}_{d}(\text{app})$ or $K^\text{ADP}_{d}(\text{app})$) is represented by the amount that either $k_{-1b}$ exceeds $k_{-1a}$ (ATP) or $k_{-5b}$ exceeds $k_{-5a}$ (ADP), according to Equations 8 and 9, respectively.

$$K^\text{ATP}_{d}(\text{app}) = \frac{k_{-1b}k_{-1a}}{k_{+1b}k_{+1a}}$$

(Eq. 8)

$$K^\text{ADP}_{d}(\text{app}) = \frac{k_{+5b}k_{+5a}}{k_{-5b}k_{-5a}}$$

(Eq. 9)

The two-step nucleotide binding reactions in Fig. 5 can account for the apparently slow bimolecular association rates (Table 2, $k_{+1(\text{app})}$ and $k_{+5(\text{app})}$) in that fast first order rates of clamping ($k_{+1b}$ and $k_{-5b}$) are augmented by initial rapid equilibrium weak binding ($K^\text{ATP}_{d}$ and $K^\text{ADP}_{d}$), according to Equations 10 and 11,

$$k^\text{ATP}_{+1(\text{app})} = \frac{k_{+1}[\text{ATP}]}{K^\text{ATP}_{d} + [\text{ATP}]}$$

(Eq. 10)

$$k^\text{ADP}_{+5(\text{app})} = \frac{k_{-5}[\text{ADP}]}{K^\text{ADP}_{d} + [\text{ADP}]}$$

(Eq. 11)

whereas the apparently slow first order nucleotide dissociation rates (Table 2, $k_{-1(\text{app})}$ and $k_{-5(\text{app})}$) represent the overall rate-limiting conformational change that precedes a more rapid diffusion separation ($k_{\text{cat}} = k_{-1b} = k_{-5b}$). In addition, the two-step nucleotide binding reactions most readily explain how rate-limiting ADP release can be increased in the presence of cosolutes, which are typically used as viscosogens to decrease diffusion rates of product release. In this case, rate-limiting conformational “unclamping” (Fig. 5, $k_{\text{cat}} = k_{-1b} = k_{-5b}$) is enhanced by osmotic pressure induced by added cosolute, which favors a less hydrated nucleotide-free form of the enzyme.

Protecting osmolytes are routinely added to enzyme preparations to increase enzyme stability during freezing and storage. The emerging view is that protecting osmolytes raise the free energy of a more hydrated partially “unfolded” state, which favors a less hydrated and more “folded” population (15–18). Accordingly, we may postulate that nucleotide clamping results in some peptide region(s) becoming more unfolded and solvent exposed.

After utilizing a conformational “clamp” for sequential nucleotide and peptide binding, fully activated S6K1 then utilizes fast and favorable phosphoryl transfer as a chemical clamp ($K^\text{3}$) (Fig. 5) in the transient ternary complex, which converts rapid equilibrium weak binding of the peptide substrate ($K^\text{m(tide)} = k_{-3}/k_{+3} \approx 180 \mu M$) to apparently high affinity (Equation 4, $K^\text{m(tide)} \approx 4–5 \mu M$). Adams and coworkers first noted and coined “catalytic clamping” in the reactions of (i) yeast Sk1p protein kinase with the RNA carrier protein Np13 (19) and (ii) the human C-terminal Src kinase with Src (20); we agree that natural selection of the fast chemistry coupled to weak peptide binding mechanism has been advantageous, because it facilitates high peptide specificity but with quick production and release of the phosphorylated downstream protein target (21).
Kinetic Mechanism of S6K1 Kinase

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