Article

Dissection of Binding between a Phosphorylated Tyrosine Hydroxylase Peptide and 14-3-3 ζ : A Complex Story Elucidated by NMR

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ABSTRACT Human tyrosine hydroxylase activity is regulated by phosphorylation of its N-terminus and by an interaction with the modulator 14-3-3 proteins. We investigated the binding of singly or doubly phosphorylated and thiophosphorylated peptides, comprising the first 50 amino acids of human tyrosine hydroxylase, isoform 1 (hTH1), that contain the critical interaction domain, to 14-3-3 ζ , by ³¹P NMR. Single phosphorylation at S19 generates a high affinity 14-3-3 ζ binding epitope, whereas singly S40phosphorylated peptide interacts with 14-3-3z one order-of-magnitude weaker than the S19-phosphorylated peptide. Analysis of the binding data revealed that the 14-3-3 ζ dimer and the S19- and S40-doubly phosphorylated peptide interact in multiple ways, with three major complexes formed: 1), a single peptide bound to a 14-3-3 ζ dimer via the S19 phosphate with the S40 phosphate occupying the other binding site; 2), a single peptide bound to a 14-3-3z dimer via the S19 phosphorous with the S40 free in solution; or 3), a 14-3-3 ζ dimer with two peptides bound via the S19 phosphorous to each binding site. Our system and data provide information as to the possible mechanisms by which 14-3-3 can engage binding partners that possess two phosphorylation sites on flexible tails. Whether these will be realized in any particular interacting pair will naturally depend on the details of each system.

INTRODUCTION

Members of the 14-3-3 protein family are important modulators of several key signaling pathways that regulate critical biological activities, including progression through the cell cycle, growth, proliferation, and apoptosis ([1–3\)](#page-8-0). Extensive proteomics studies have revealed that the 14-3-3 family targets >200 distinct partners, ~0.6% of the entire cellular proteome $(4,5)$.

In humans, seven different 14-3-3 isoforms $(\beta, \gamma, \varepsilon, \eta, \sigma, \gamma)$ τ , and ζ) are found, each encoded by a distinct gene, that function as homo- or heterodimers of ~30-kDa proteins ([6,7](#page-8-0)). Crystal structures of all seven homodimeric mammalian proteins are available. The fold of the monomeric unit contains nine α -helices, arranged in an antiparallel fashion. Within the dimeric structure, two amphipathic grooves, one per monomer, constitute the binding pockets in which the conserved KRRY residues-Lys⁴⁹, Arg⁵⁶, Arg¹²⁷, and Tyr 128 in 14-3-3 ζ -are positioned for the recognition of a phosphorylated serine or threonine target [\(8](#page-8-0)). Three different phosphoserine (pS)- or phosphothreonine-containing sequence motifs can be recognized by 14-3-3 proteins: RSXp(S/T)XP (motif I), RXXXp(S/T)XP (motif II), and a carboxyl-terminal $p(S/T)XCOOH$ motif (motif III), where X is a nonproline amino acid $(9-11)$. Some 14-3-3 family members have also been reported to bind nonphosphory-

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lated peptides as well as motifs that do not strictly display the above canonical sequences $(6,11)$ $(6,11)$ $(6,11)$.

Tyrosine hydroxylase (TH) and tryptophan hydroxylase were the first proteins that were identified as binding partners for 14-3-3 proteins [\(12\)](#page-8-0). TH catalyzes the tetrahydrobiopterin-dependent hydroxylation of L-tyrosine to L-3,4-dihydroxyphenylalanine, which is the rate-limiting step in the biosynthesis of dopamine and other catecholamines ([13\)](#page-9-0). In humans, several isoforms of TH are expressed, with isoform 1 (hTH1; 497 residues, NP-000351.2) being the best characterized ([14–16](#page-9-0)). hTH1 can be phosphorylated at T8 and/or three serine residues (S19, S31, and S40) by several protein kinases [\(17](#page-9-0)) on its highly dynamic 50-residue N-terminal region that precedes the regulatory ACT domain, characteristic of the aromatic amino-acid hydroxylase family. 14-3-3z and other isoforms have been reported to bind to S19 phosphorylated TH [\(18–20](#page-9-0)). Although not strictly exhibiting the canonical binding motifs, recent crystallographic analysis of the complex between hTH1 S19-phosphorylated N-terminal peptide (1–43 residues) and 14-3-3 γ revealed that the phosphate interacts with the KRRY motif in the 14-3-3 γ binding grove, and the region surrounding S19 engages in similar interactions as seen for peptides containing canonical motifs I and II [\(21](#page-9-0)). In addition, phosphorylation of hTH1 on S40 stimulates TH activity [\(17](#page-9-0)), and induces binding to the yeast 14-3-3 proteins BMH1 and BMH2 and to sheep brain 14-3-3 proteins, but not to 14-3- 3ζ [\(18,19\)](#page-9-0). On the other hand, 14-3-3 ζ has been reported

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to interact with doubly S19- and S40-phosphorylated hTH1 ([18\)](#page-9-0). In a recent study ([22\)](#page-9-0) we have shown that the doubly phosphorylated hTH1 binds $14-3-3\gamma$ with similar stoichiometry and affinity as the singly phosphorylated enzyme on Ser¹⁹. However, 14-3-3 γ interfered with the phosphorylation of TH1-pS19 on Ser⁴⁰, supporting that Ser⁴⁰ becomes inaccessible in the hTH1:14-3-3 γ complex, although Ser⁴⁰ does not significantly contribute to the binding of $14-3-3\gamma$. Therefore the question of whether S40 phosphorylated hTH1 can be bound by 14-3-3 proteins has not been settled unequivocally.

The 14-3-3 proteins are thought to function by inducing a conformational change in the target proteins, thereby modulating the target proteins' activities. In this way, 14-3-3 family members impart additional control onto key cellular enzymes, in addition to common enzymatic control mechanisms such as feedback loops or product inhibition. The contemporary prevalent model for the binding of doubly phosphorylated peptides by 14-3-3 suggests that one phosphorylated residue of the target protein engages 14-3-3 first, linking the two proteins together, followed by binding of the second phosphorylated residue, to effect the conformational change ([23\)](#page-9-0). The notion that binding of 14-3-3 to two phosphorylation sites is required to induce the activity altering conformational change, has led some authors to consider 14-3-3 as a binary computer $(4,24)$ $(4,24)$ $(4,24)$.

We explored this model of 14-3-3 modulation using $\mathrm{^{31}P}$ NMR in a simplified peptide system. We investigated the binding of hTH1 peptides (region 1–50) with a single phosphorylation at $S19$ (pS19) or S40 (pS40) to 14-3-3 ζ and determined individual binding affinities. We found that $14-3-3\zeta$ can interact with the hTH1 peptide through either phosphorylated-S19 or -S40, although the binding affinity of $14-3-3\zeta$ for the pS40 peptide is significantly lower than that for the $pS19$ peptide. Furthermore, binding of $14-3-3\zeta$ to the doubly phosphorylated hTH1 peptide exhibits a complex binding profile involving three major complexes. Our results suggest that when considering 14-3-3 interactions with multiply phosphorylated targets in general, and hTH1 in particular, avidity considerations should be taken into account.

METHODS

Cloning, expression, and purification of hTH1 peptides and 14-3-32

The coding region for residues 1–50 of human tyrosine hydroxylase $(hTH1–50)$ ([Fig. 1](#page-2-0) A) was inserted into a modified pET32a vector, in frame with N-terminal thioredoxin and $6\times$ His tags, followed by a tobacco etch virus (TEV) cleavage site. A codon-optimized gene for 14-3-3z (GenScript, Piscataway Township, NJ) was inserted into pET15b with Cys^{25} and Cys189 codons converted to Ala using the QuikChange Site-Directed Mutagenesis Kit (Stratagene, La Jolla, CA). The DNA sequence for all constructs was confirmed by sequencing (Genewiz, South Plainfield, NJ).

Proteins were produced in Escherichia coli BL21(DE3) cells that were grown at 37 $\rm{^{\circ}C}$ in LB media to an OD₆₀₀ of 0.8 and induced with 0.5 mM IPTG for 18 h at 18°C. Cells were harvested, resuspended in 50 mM TBS (50 mM Tris, 150 mM NaCl, pH 8.0) buffer, and lysed with a microfluidizer (Microfluidics, Westwood, MA). The cell lysate was clarified by centrifugation (21,000g for 60 min at 4° C), and the supernatant was loaded onto a HisTrap column (GE Healthcare, Piscataway, NJ). Proteins were eluted using a linear imidazole gradient (0.05–1 M in a total volume of 15 mL).

Fractions containing tagged hTH1 peptide were treated with TEV (1 mg/ 20 mg of peptide) at 4° C overnight, and applied to a Vydac reverse-phase C4 column (Grace Davison Discovery Sciences, Deerfield, IL). hTH1–50 was eluted from the reverse phase column using a linear gradient of 10– 40% acetonitrile in 50 mL. Peptide-containing samples were further purified by gel filtration through a Superdex75 26/60 column (GE Healthcare) equilibrated in 20 mM Tris buffer, pH 7.5. The Trp-containing three aminoacid cloning artifact at the N-terminus of the hTH1–50 peptide [\(Fig. 1](#page-2-0) A) allowed for detection during purification and concentration determination at 280 nm.

Fractions containing 14-3-3 ζ were dialyzed into TBS buffer and treated with TEV overnight. Undigested protein, $6 \times H$ is tag cleavage products, and TEV protease were removed by passing the sample over the HisTrap column for a second time and the flow-through containing untagged $14-3-3\zeta$ was collected. The sample was further purified over anion exchange (HiTrap Q HP; GE Healthcare) followed by gel filtration on a Superdex75 26/60 column (GE Healthcare), equilibrated in 20 mM sodium phosphate buffer, pH 7.0.

The Cys^{25} Ala/Cys¹⁸⁹Ala 14-3-3 ζ variant exhibited essentially the same stability as wild-type 14-3-3 ζ protein, as evidenced by identical differential scanning calorimetry unfolding profiles.

Phosphorylation of hTH1–50

The serine residue at position 40 of the hTH1–50 peptide was phosphorylated using the catalytic subunit of cAMP-dependent protein kinase A (PKA; New England BioLabs, Ipswich, MA), whereas Ser^{19} was phosphorylated using p38 regulated/activated protein kinase (PRAK; obtained from the protein phosphorylation unit, University of Dundee, Dundee, Scotland). For phosphorylation at S19, the reaction mixture, containing hTH1–50 S40A peptide in 50 mM Tris buffer, pH 7.5, 1 mM ATP, 10 mM $MgCl₂$, and PRAK kinase (50 μ g for 10 mg of hTH1–50 S40A), was incubated for three days at 30°C. For phosphorylation at S40, hTH1-50 was incubated with PKA (0.5 μ g for 15 mg hTH1–50) in 50 mM Tris buffer, pH 7.5,10 mM $MgCl₂$, 200 μ M ATP at 30°C for 2 h. Reaction products were purified by gel filtration using a Superdex30 16/60 column (GE Healthcare) followed by anion exchange (HiTrap Q HP; GE Healthcare).

Doubly phosphorylated peptide was prepared using S40 phosphorylated hTH1–50 peptide (pS40-hTH1–50) and PRAK kinase, as described above. Approximately 70% of pS40-hTH1–50 was converted to the doubly phosphorylated product (pS19pS40-hTH1–50), which was purified as described above. For thiophosphorylation, adenosine 5'-[y-thio]triphosphate (ATPyS) was added to the PKA reaction mixture, instead of ATP. In all cases, the correct masses and purity of the final products were verified by matrix-assisted laser desorption/ionization-time of flight mass spectrometry.

³¹P NMR titrations

All ^{31}P NMR spectra were recorded at $37^{\circ}C$ on an AVANCE 600 MHz NMR spectrometer equipped with a broadband frequency probe (Bruker, Billerica, MA). Samples containing 14-3-3 ζ and singly or doubly phosphorylated hTH1–50 peptides were prepared in 20 mM sodium phosphate, pH 7.0, at a constant peptide concentration (0.83 or 0.71 mM) with molar ratios of $14-3-3\zeta$ to peptide of $0:1-8:1$.

Modeling of binding scenarios

Binding schemes for the different complexes between $14-3-3\zeta$ and singly or doubly phosphorylated peptides are presented in [Fig. 1](#page-2-0) C and [Fig. 2](#page-3-0) B, A SEW MPTPDATTPQ¹⁰ AKGFRRAVSE²⁰ LDAKQAEAIM³⁰ SPRFIGRRQS⁴⁰ LIEDARKERE⁵⁰

FIGURE 1 Binding of singly phosphorylated hTH1–50 peptides to 14-3-3 ζ . (A) Amino-acid sequence of the hTH1–50 peptide with S19 and S40 highlighted (boldface). (B) ³¹P spectra of pS19-hTH1–50 at 0.83 mM peptide concentration and molar ratios of $14-3-3\zeta$ to peptide (r) 0.5:1 (black) and 1:1 ($gray$). (C) Schematic depiction of pS19-hTH1–50 (denoted by 19–40) binding to the 14-3-3 ζ homodimer (free protein dimer is depicted by $P_{\emptyset}^{\emptyset}$, where \emptyset indicates an empty phosphoserine binding site). (Circled) Bound pS19 residues. (D) ^{31}P spectra of pS40-hTH1–50 at 0.71 mM peptide concentration and molar ratios (r) of $14-3-3\zeta$ to peptide of 0:1 (blue), 0.5:1 (red), 1:1 (dark green), 2:1 (purple), 4:1 (orange), and 8:1 (brown). (Inset) Chemical shift changes upon 14-3-3 ζ binding ($\Delta \delta$) are plotted as a function of molar ratio (r). The curve obtained by fitting $\Delta \delta$ to the quadratic equation shown in Eq. 2 is shown. To see this figure in color, go online.

respectively. Equations for individual binding equilibria and equations describing mass conservation are provided in [Fig. S1](#page-8-0) and [Fig. S2](#page-8-0) in the [Sup](#page-8-0)[porting Material](#page-8-0). The equations were solved numerically using the software MATLAB (The MathWorks, Natick, MA) and results yield populations of the individual species as a function of the $14-3-3\zeta$ to peptide molar ratios, r. Three different sets of solutions were obtained, only one of which was physically meaningful given that populations cannot be negative for any r value ≥ 0 .

RESULTS

Binding of singly phosphorylated hTH1 peptides to 14-3-3z

Binding between $14-3-3\zeta$ and the phosphorylated N-terminus of hTH1 was probed by ${}^{31}P$ NMR. hTH1 peptides, containing the first 50 residues (hTH1–50), were prepared and each serine in the peptide was phosphorylated individually, generating pS19-hTH1–50 and pS40-hTH1–50. PKA selectively and specifically phosphorylated only Ser^{40} , but PRAK phosphorylated both Ser¹⁹ and Ser⁴⁰ (J. Hritz and A.M. Gronenborn, unpublished results). Therefore, we introduced the S40A change, permitting the preparation of singly phosphorylated pS19-hTH1–50 peptide. Titration of $14-3-3\zeta$ into a solution containing pS19-hTH1–50 revealed the system to be in slow exchange on the NMR chemical shift timescale. Thus, resonances for both free and the 14-3-3 ζ -bound peptide can be observed separately (Fig. 1 B), and the relative fractions of the free and bound peptide can be determined from resonance intensities $(25,26)$. A dissociation constant (K_{d1}) of 0.15 \pm 0.03 mM for the 14-3-3 ζ /pS19hTH1–50 complex was derived, using

$$
K_{d1} = \frac{\left[pS19^{\text{free}}\right] * \left[14 - 3 - 3\zeta^{\text{free}}\right]}{\left[pS19^{\text{bound}}\right]}
$$

$$
= \frac{c_{\text{pep}} * A(pS19^{\text{free}}) * \left\{r - A(pS19^{\text{bound}})\right\}}{A(pS19^{\text{bound}})}, \quad (1)
$$

FIGURE 2 Binding of doubly phosphorylated peptide to 14-3-3 ζ . (A) ³¹P spectra of pS19pS40hTH1–50 at 0.83 mM and the molar ratio of 14- 3-3 ζ to the peptide (r) was 0:1 (blue), 0.5:1 (red), 1:1 (dark green), 2:1 (purple), 3:1 (yellow), 4:1 (orange), or 5:1 (light green). (B) Binding scheme of pS19pS40-hTH1–50 (denoted by 19–40) with respect to the 14-3-3 ζ homodimer ($P^{\emptyset}_{\emptyset}$). Dissociation constants K_{d1} and K_{d2} correspond to the values determined for the singly phosphorylated peptides. (Blue arrows) Equilibria for which slow exchange on the $31P$ chemical shift scale is observed; (red arrows) equilibria for which fast exchange on the $31P$ chemical shift scale is observed. (*Circled*) Bound phosphorous groups. Phosphorous groups contributing intensity to the same peak in the NMR spectra are shown in the same color. (C) Equations governing the relative peak intensities (denoted as ''A''), based on the scheme depicted in panel B; each phosphopeptide contributes to one or more of the five peaks. The specific phosphoserine that contributes to intensity of a particular peak is colored in the same color as in panel B. (D) Populations of phosphoserine species that contribute to the experimentally observed peak intensities (x) for different 14-3-3 ζ to peptide molar ratios, r. (Continuous lines) For comparison, the theoretically predicted populations, based on the binding model in panel B using $K_{d1} = 0.15$ mM, $K_{d2} = 1.0$ mM, and $K_{eq} = 1.5$, are plotted. To see this figure in color, go online.

where $c_{\rm pep}$, $A(pS19^{\rm free})$, and $A(pS19^{\rm bound})$ comprise the total peptide concentration, and the free and bound fractions of the pS19-hTH1–50 peptide, respectively, with r the molar ratio of $14-3-3\zeta$ to peptide. For both singly phosphorylated peptides, dissociation constants are reported for the $14-3-3\zeta$ monomer, assuming that binding to a single site in the 14-3- 3ζ homodimer is not influenced by binding to the second site.

To analyze our binding data, we generated a scheme for the interaction between $pS19-hTH1–50$ and the 14-3-3 ζ dimer ([Fig. 1](#page-2-0) C). A pS19-hTH1–50 peptide (represented by 19–40 in [Fig. 1](#page-2-0) C) can bind to the free 14-3-3 ζ dimer (depicted by $P_{\emptyset}^{\emptyset}$ in [Fig. 1](#page-2-0) C) at two different sites, with the dissociation constant being one-half of the observed K_{d1} ; the binding of a second pS19-hTH1–50 peptide to the $14-3-3\zeta$ dimer in which one site is already occupied

(depicted by P_{19-40}^{\odot} in [Fig. 1](#page-2-0) C) can only occur to the un-
occurried site and dissociation from a doubly occurried occupied site, and dissociation from a doubly occupied 14-3-3 ζ dimer (depicted by P_{19-40}^{19-40} in [Fig. 1](#page-2-0) C) can
occur from either site resulting in $2*K_{\text{H}}$. The equaoccur from either site, resulting in $2*K_{d1}$. The equations describing these two equilibria, taking mass conservation of the peptide and $14-3-3\zeta$ into account, are presented in [Fig. S1](#page-8-0) in the [Supporting Material.](#page-8-0) Based on this model, in a 1:1 mixture of pS19-hTH1–50 and 14-3-3 ζ (monomer concentration) with a K_{d1} of 0.15 mM, 34.4% of the peptide would be in the free state, 22.6% of peptide in the $\overline{P}_{19-40}^{\odot}$ bound form, and 43% in the P_{19-40}^{\odot} bound form the P_{19-40}^{19-40} bound form.

In contrast to the result with the pS19 peptide, titration of 14-3-3 ζ into a solution containing the pS40-hTH1-50 peptide revealed fast exchange on the NMR chemical shift scale, resulting in a single resonance at the weighted averaged chemical shifts of both the free and bound resonances ([Fig. 1](#page-2-0) D). By fitting the chemical shift change $(\Delta \delta)$ to Eq. 2, a K_{d2} value of 1.0 \pm 0.1 mM and a maximum chemical shift difference between free and bound peptide ($\Delta \delta_{\text{max}}$) of 0.94 p.p.m. were determined for the binding of 14-3-3 ζ to pS40hTH1–50:

$$
\Delta \delta = \frac{\Delta \delta_{\text{max}}}{2c_{\text{pep}}} \left(K_d + c_{\text{pep}} + r c_{\text{pep}} \right) - \sqrt{\left(K_d + c_{\text{pep}} + r c_{\text{pep}} \right)^2 - 4c_{\text{pep}}^2 r} \right). \tag{2}
$$

These values allowed us to calculate the $31P$ bound chemical shift, $\delta(pS40^{bound})$, which was 3.26 p.p.m. ([Fig. 1](#page-2-0) D).

Interestingly, the bound $31P$ chemical shifts for both singly phosphorylated hTH1–50 peptides are essentially identical $(3.26 \text{ p.p.m.}, \text{Fig. 1}, B \text{ and } D)$ $(3.26 \text{ p.p.m.}, \text{Fig. 1}, B \text{ and } D)$ $(3.26 \text{ p.p.m.}, \text{Fig. 1}, B \text{ and } D)$, irrespective of the different neighboring amino acids around the phosphorylated Ser in the hTH1–50 peptide. This suggests that the binding of the phosphorylated residues to the identical binding pockets of $14-3-3\zeta$ that are lined by residues Lys⁴⁹, Arg⁵⁶, Arg¹²⁷, and Tyr¹²⁸ is the major determinant of the chemical shift, irrespective of the local peptide sequence. We speculate that the side chain of Tyr^{128} most likely causes the upfield shift of the pS19 and $pS40$ ³¹P resonances.

Binding of doubly phosphorylated hTH1 peptide to 14-3-3z

Binding between doubly phosphorylated proteins and 14-3- 3 has previously been characterized using various methods, including isothermal titration calorimetry and fluorescence polarization [\(19,27,28\)](#page-9-0). Although these methods provide overall binding properties, such as K_d or stoichiometry, important details for a system involving a homodimer and two binding motifs on the target are lacking. For example, answers to the question of how individual phosphorylated amino acids interact with 14-3-3, and what kind of complexes are possible, are still incomplete. $31P$ NMR is ideally suited to provide such information, since individual ^{31}P signals can be monitored throughout the course of titration experiments between various interacting partners. We therefore prepared doubly phosphorylated hTH1–50 peptide $(pS19pS40-hTH1–50)$ and monitored $14-3-3\zeta$ binding to this peptide by $31P$ NMR. Our data reveal that a number of different bound species are present during the course of the titration (Fig. 2 A). The spectrum of free pS19pS40-hTH1–50 (blue spectrum in [Fig. 2](#page-3-0) A; $r = 0$) contains two well-separated resonances, at 3.76 p.p.m. and at 4.18 p.p.m. for the phosphorous atoms on $pSer¹⁹ (pS19^{free})$ and pSer⁴⁰ (pS40^{free}), respectively. At the end of the titra-tion (light green trace in [Fig. 2](#page-3-0) A; $r = 5$), the pS19pS40hTH1–50 peptide is saturated with $14-3-3\zeta$, and resonances at 3.26 p.p.m. and 3.43 p.p.m. for the phosphorous atoms on $p\text{Ser}^{19}$ (pS19bound) and $p\text{Ser}^{40}$ (pS40°), respectively, are observed. The bound resonances are broader than those of the free peptide, reflecting the slower rotational correlation time of the peptide/14-3-3 ζ complex (molecular mass $~52-67$ kDa).

The chemical shift of 3.43 p.p.m. is different from the completely saturated position (3.26 p.p.m.) obtained with a peptide that contains only phosphorylated Ser⁴⁰ (pS40-hTH1–50, [Fig. 1](#page-2-0) D), indicating that this bound resonance position contains a contribution from the free pS40 resonance (due to fast exchange on the chemical shift scale). This signal, therefore, is designated $pS40^c$, instead of pS40^{bound}. During the titration, the intensities of both the pS19free and pS40free resonances decrease to the same degree with increasing $14-3-3\zeta$ concentration, but only the bound $pS19$ ³¹P resonance at 3.26 p.p.m. increases in intensity at the same rate (Fig. 2 A , e.g., *red spectrum*). The bound $pS40$ ³¹P resonance at 3.43 p.p.m. is only visible for excess $14-3-3\zeta$ over peptide (e.g., *purple*, *yellow*, *or*ange, and light-green spectra). These data suggest that for excess peptide and little 14-3-3 ζ ($r < 1$), the phosphorous group on Ser^{19} (strong binder) is the one predominantly bound by $14-3-3\zeta$, and two $pS19pS40-hTH1-50$ peptides are likely bound by one $14-3-3\zeta$ dimer. For conditions where 14-3-3 ζ is in excess ($r > 1$), both phosphorous groups on a single peptide can be bound by one $14-3-3\zeta$ dimer.

The complete binding scenario, considering six possible complexes for the interaction between doubly phosphorylated peptide (free peptide is depicted as 19–40) and 14- 3-3 ζ (free 14-3-3 ζ dimer is depicted as $P^{\varnothing}_{\varnothing}$), is presented in [Fig. 2](#page-3-0) B. Based on this interaction scenario and taking into account that resonances can be in slow or fast exchange on the chemical shift scale (marked by blue and red arrows, respectively), five different resonances are expected to be observable in the $3^{1}P$ NMR spectra (see below). Phosphorous groups that exhibit identical resonance frequencies, thus contributing intensity to the same peak, are labeled in the same colors in [Fig. 2,](#page-3-0) B and C. The derived binding scheme is based on two assumptions:

- 1. The bound $3^{1}P$ chemical shifts of pS19 and pS40 are identical when saturated with $14-3-3\zeta$, irrespective of whether the second binding site on the $14-3-3\zeta$ dimer is occupied or not; and
- 2. The individual phosphorous resonances in pS19pS40 hTH1–50 exhibit the same NMR exchange regime as those for the singly phosphorylated peptides ([Fig. 1](#page-2-0), B and D , i.e., the pS19 resonance is in slow exchange and that of pS40 is in fast exchange on the ^{31}P NMR shift scale).

Note that the slow or fast exchange regimes apply not only to the interaction between free peptide (19–40) and the one site occupied 14-3-3 ζ dimer complex $(P_{19-40}^{\emptyset} \text{ or } P_{40-19}^{\emptyset})$ but
also to various additional 14-3-3 ζ complexes (Fig. 2.8). For also to various additional 14-3-3 ζ complexes ([Fig. 2](#page-3-0) B). For example, the pS40 phosphorous resonance of P_{19}^{40} , the complex in which both sites on the protein dimer engage the two phosphate groups on 19‒40, is in fast exchange with that of $\overline{P_{0,-40}^{\odot}}$, the complex in which one site on the dimer engages
the pS10 phosphate group of the pentide since very weak the pS19 phosphate group of the peptide, since very weak binding of the phosphate group of pS40 is involved. On the other hand, \hat{P}_{40}^{40} and \hat{P}_{40-19}^{0} are in slow exchange, since the exchange involves tight binding of the phosphate group of pS19. In agreement with the above binding scheme, five distinct peaks are observed in the ^{31}P spectra during the titration ([Fig. 2](#page-3-0) A).

Multiple complexes contribute to the intensity of each phosphorous resonance. All peptides with the pS19 phosphate group free of 14-3-3 ζ protein (19–40, $P_{40-19}^{\mathcal{O}}, P_{19-40}^{40-19}, P_{40-19}^{40-19}$)
contribute to the peak labeled "pS19^{free}" (19 colored *green* in [Fig. 2](#page-3-0), B and C). Free peptide $(19-40)$ as well as any peptides bound to $14-3-3\zeta$ via the pS40 phosphate group $(P_{40-19}^{\emptyset}, P_{19-40}^{40-19}, P_{40-19}^{40-19})$ (40 colored *orange* in [Fig. 2](#page-3-0), B and
C) will contribute to the peak labeled "pS40^{free}" in Fig. 2.4 C) will contribute to the peak labeled " $pS40$ ^{free}" in [Fig. 2](#page-3-0) A. Given that the same peptide can contribute intensity to two different resonances ($pS19^{free}$ and $pS40^{free}$), the total intensity of the two resonances is expected to be similar, as is indeed observed in the ${}^{31}P$ spectra [\(Fig. 2](#page-3-0) A).

Any complexes of 14-3-3 ζ that contain peptide bound via the phosphate group on pS19 $(P_{0-40}^{\circ}, P_{19-40}^{19-40}, P_{19}^{40}]$, and P_{0-40}^{40-19} , and P_{19-40}^{40-19} , and P_{19-40} intensity P_{19-40}^{40-19} ; 19 colored *blue* in [Fig. 2](#page-3-0), *B* and *C*) add intensity to the peak labeled "pS19^{bound}." The pS40^c peak arises from the pS40 phosphate groups in the P_{19}^{40} and P_{19-40}^{9} com-
plexes (40 colored *purple* in Fig. 2, B and C), which are in plexes (40 colored purple in [Fig. 2,](#page-3-0) B and C), which are in fast exchange. The contribution from the free pS40 phosphate group in P_{19-40}^{\odot} causes the "pS40^c" resonance to
slightly shift toward the free position (4.18 n n m) and slightly shift toward the free position (4.18 p.p.m.) and, therefore, it is observed at 3.43 p.p.m., slightly downfield from the bound position (3.26 p.p.m.) of the singly phosphorylated pS40 peptide (Fig. $1 D$). In contrast, the position of the $pS19^{bound}$ peak is at 3.27 p.p.m., identical to the bound position of the singly phosphorylated pS19 peptide (Fig. $1 B$). The reason for this is that the phosphate groups that contribute to the $pS19^{bound}$ peak are in slow exchange, thus no weighted chemical shift averaging comes into play.

Finally, the free pS40 phosphate groups in the P_{19-40}^{19-40} and P_{19-40}^{19-40} complexes (40 colored brown in Fig. 2, B and C) de- P_{10-19}^{19-40} complexes (40 colored *brown* in [Fig. 2](#page-3-0), B and C), de-
noted pS40^{*} reside on the pentide that is bound to 14.3.3% noted pS40*, reside on the peptide that is bound to $14-3-3\zeta$ via pS19, but they are not buried in the $14-3-3\zeta$ binding site and are free in solution, surrounded by solvent. Note that pS40* is observed as a distinct peak at 3.93 p.p.m., in slow exchange with the pS40 phosphate resonance (4.18 p.p.m.) of the free peptide ([Fig. 2](#page-3-0) A), because the peptides in these complexes are bound to $14-3-3\zeta$ via the pS19 phosphate group.

The relationships listed in Fig. $2 C$ imply that the total normalized area (denoted as " A ") of all peaks in [Fig. 2](#page-3-0) A corresponds to 2, because the peptide contains two phosphate groups:

$$
(A(pS19free) + A(pS40free) + A(pS19bound) + A(pS40c) + A(pS40*) * cpep = 2([19 – 40] + [P40-190]+ [P19-400] + [P1940]+ 2[P40-1940]+ 2[P40-1940]= 2 * cpep.
$$
 (3)

Assignment of phosphorous resonances by selective thiophosphorylation

The ³¹P resonance assignment of the pS19 and pS40 in the $pS19pS40-hTH1–50$ peptide ([Fig. 2](#page-3-0) A) was initially obtained using the assignment of singly phosphorylated hTH1–50 peptides ([Fig. 1](#page-2-0), B and D). We confirmed the assignment using a hTH1–50 peptide that was phosphorylated on Ser^{19} and thiophosphorylated on Ser^{40} (pS19tpS40-hTH1–50). Thiophosphorylation causes a large downfield shift of the ^{31}P resonance from ~4.2 p.p.m. (blue spectra in [Fig. 1](#page-2-0) D and [Fig. 2](#page-3-0) A) to 43.95 p.p.m. (blue spectra in [Fig. 3,](#page-6-0) A and B). Thus, in the spectrum of the pS19tpS40-hTH1–50 peptide [\(Fig. 3](#page-6-0) B), both resonances are clearly separated.

A thiophosphate group is assumed to functionally mimic a phosphate group, given its close chemical structure. To confirm that no gross changes to $14-3-3\zeta$ binding were introduced into the system by thiophosphorylation, we carried out titration experiments with tpS40-hTH1–50 and $14-3-3\zeta$ under the same conditions that were used for pS40-hTH1– 50. We determined a K_d value of 3.8 mM for tpS40-hTH1– 50 ([Fig. 3](#page-6-0) A), compared to 1.0 mM for the pS40-hTH1–50 ([Fig. 1](#page-2-0) D). This demonstrates that binding is slightly reduced by the replacement of an oxygen atom with a sulfur atom. Interestingly, binding of 14-3-3z to tpS40-hTH1–50 causes a downfield $3^{1}P$ chemical shift ([Fig. 3](#page-6-0) A), whereas binding to pS40-hTH1–50 induces an upfield shift (see above, and Fig. $1 D$), illustrating that chemical shifts are extremely sensitive to any changes and hard to predict.

Whereas 14-3-3z binding to the pS19pS40-hTH1–50 peptide yields three distinct pS40 phosphorous signals that

FIGURE 3 Binding of thiophosphorylated hTH1-50 peptides to 14-3-3 ζ . (A) ^{31}P spectra of tpS40-hTH1–50 peptide (0.71 mM) for varying molar ratios of 14-3-3 ζ to the peptide (r); 0:1 (blue), 0.5:1 (red), 1:1 (dark green), 2:1 (purple), 4:1 (orange), and 8:1 (brown). (Inset) The chemical shift changes upon 14-3-3 ζ binding ($\Delta \delta$) are plotted as a function of molar ratio (r). (Continuous line) Predicted $\Delta \delta$ from fitting the experimental $\Delta \delta$ to the quadratic equation shown in Eq. 2. (B) ³¹P spectra of pS19tpS40-hTH1-50 (0.83 mM) for varying molar ratios of 14-3-3 ζ to the peptide (r): 0:1 (blue), 0.5:1 (red), 1:1 (dark green), 2:1 (purple), 3:1 (yellow), 4:1 (orange), and 5:1 (light green). To see this figure in color, go online.

exhibit slow exchange, $pS40^{free}$, $pS40^*$, and $pS40^c$ [\(Fig. 2](#page-3-0) A), binding to the pS19tpS40-hTH1–50 peptide results in only two signals that are in fast-intermediate exchange (Fig. 3 B), caused in part by the smaller chemical shift difference between free ($pS40$ ^{free}) and bound ($pS40^\circ$) resonances (0.60 p.p.m) compared to that for pS19pS40-hTH1–50 (0.94 p.p.m.). However, even with this difference, it seems reasonable to assume that the binding scheme in Fig. $2 B$ can be applied to pS19tpS40-hTH1–50, with $K_{d2} = 3.8 \text{ mM}$ instead of 1 mM and $K_{eq} = 0.26$ (see below) rather than 1.5. Using the well-separated $pS19^{bound}$ ³¹P resonance in the thiophosphorylated peptide (Fig. $3 B$) proved beneficial for deconvoluting the overlapping pS19 resonances in the 14-3-3z-bound pS19pS40-hTH1–50 spectra (see [Fig. S3](#page-8-0)).

Analysis of the binding equilibrium between doubly phosphorylated hTH1 peptide to the 14-3-3 ζ homodimer

[Fig. 2](#page-3-0) B illustrates a comprehensive equilibrium binding scenario for the interaction between the doubly phosphorylated hTH1 peptide and the 14-3-3 ζ homodimer. Six possible complexes are considered. Because insufficient experimental data for the accurate determination of all individual dissociation constants are available, it is assumed that each subunit monomer in the $14-3-3\zeta$ homodimer binds equivalent phosphate groups on pS19pS40-hTH1–50 with identical affinity, irrespective of the state of the other 14-3-3z monomer, i.e., no cooperativity is present. Therefore, three independent equilibrium constants, K_{d1} , K_{d2} , and K_{eq} [\(Fig. 2](#page-3-0) B), come into play. K_{d1} (0.15 \pm 0.03 mM) and K_{d2} (1.0 \pm 0.1 mM) are the experimentally determined dissociation constants for the interaction between singly phosphorylated hTH1–50 peptides, pS19-hTH1–50 and pS40-hTH1–50, respectively, and the $14-3-3\zeta$ dimer (using the monomer at the concentration unit); K_{eq} is the equilibrium constant defined by the concentration ratio of the (P_{19}^{40}) complex to the (P_{19}^{0})
complex This is treated as the only variable parameter complex. This is treated as the only variable parameter in this simplistic equilibrium binding model.

We used three approaches for determining K_{eq} , as follows: In the first approach, we derived a set of six equations for the individual, independent equilibria, along with two equations that are necessary to ensure mass conservation of the peptide and $14-3-3\zeta$ (see [Fig. S2](#page-8-0) A). These equations were solved numerically in the software MATLAB. The resulting solutions provide the molar concentration of peptide complexes as a function of the $14-3-3\zeta$ to doubly phosphorylated peptide ratio, r (see [Fig. S2](#page-8-0) B). The equations in [Fig. 2](#page-3-0) C link the theoretically predicted molar concentrations of individual complexes (see Fig. $S2 \, B$) with the intensities of five experimentally observable resonances. By fitting the theoretical profiles (continuous lines in [Fig. 2](#page-3-0) D) with the experimentally observed intensities (individual data points depicted by x in [Fig. 2](#page-3-0) D), for several 14-3-3 ζ to peptide molar ratios, r, a K_{eq} value of 1.5 was determined.

In the second approach, the value of K_{eq} could, in principle, also be determined from the observed difference in the " $pS40^c$ " resonance position from that of " $pS40^{bound}$ ", if the resonance position of pS40 in the P_{19-40}° complex is known.
Unfortunately this is not known experimentally but Unfortunately, this is not known experimentally, but, assuming that it is identical to that of (pS40*), then the following equation holds:

$$
\delta(pS40^c) = \frac{[P_{19-40}^{\circ}]\delta(pS40^*) + [P_{19}^{40}]\delta(pS40^{\text{bound}})}{[P_{19-40}^{\circ}]+[P_{19}^{40}]}.
$$
 (4)

Considering that

then

$$
K_{eq} = \frac{\left[P_{19}^{40}\right]}{\left[P_{19-40}^{\oslash}\right]}
$$

$$
K_{eq} = \frac{\delta(pS40^c) - \delta(pS40^*)}{\delta(pS40^{\text{bound}}) - \delta(pS40^c)}.
$$
 (5)

If the values of $\delta(pS40^*), 3.93 p.p.m.; \delta(pS40^c), 3.43 p.p.m.;$ and $\delta(pS40^{\text{bound}})$, 3.26 p.p.m. are applied to Eq. 5, a K_{eq} value of 3.0 is found.

In the third approach, the value of K_{eq} can be estimated, based on K_{d2} and the local concentration of the pS40 phosphate group in the P_{19-40}° complex. In an extended chain model of the doubly phosphorylated pentide, the distance model of the doubly phosphorylated peptide, the distance between the Ser 19 and Ser 40 phosphate groups is estimated to be $~6$ nm. The local concentration of the Ser⁴⁰ phosphate that resides within a sphere of 6 nm radius, if the Ser^{19} phosphate from the same peptide is fixed to the first binding site on the 14-3-3 ζ dimer, would become $c_{\text{loc}} \sim 1/$ $(6.022*10^{23}*4*10^{3}*(6.10^{-9})^{3})$ M, i.e., 1.9 mM. Thus, an estimated value of K_{eq} is

$$
[P_{19}^{40}] / [P_{19-40}^{0}] = c_{\text{loc}} / K_{d2} = 1.9.
$$

The latter two estimated K_{eq} values of 3.0 and 1.9 are very similar to the value determined from fitting $(K_{eq} = 1.5)$, considering that K_{eq} values of 3.0, 1.9, and 1.5 result in relative populations of 75, 66, and 60%, respectively, of the P_{19}^{40} complex.

Using the simplified binding model [\(Fig. 2](#page-3-0) B), with $K_{d1} =$ 0.15 mM, $K_{d2} = 1.0$ mM, and $K_{eq} = 1.5$, populations of $P_{\text{40--19}}^{\text{20}}$, $P_{\text{40--19}}^{19-40}$, and $P_{\text{40--19}}^{40-19}$ complexes were found to be equipmently lower than other complexes (\leq 5.5%; see significantly lower than other complexes \langle <5.5%; see [Fig. S2](#page-8-0) B). For example, at a 1:1 molar ratio of doubly phosphorylated peptide to 14-3-3 ζ ($r = 1$), 37.0% of the peptide is unbound; 17.7% of the peptide is in the P_{19}^{40} complex; 11.8% is in P_{19-40}^{\odot} complex; and 24.1% (2 × 12.05%) is
in the P_{19-40}^{\odot} complex. Together, all other species account in the P_{19-40}^{19-40} complex. Together, all other species account
for 9.4% of the total population. At $r = 5$, the theoretical for 9.4% of the total population. At $r = 5$, the theoretical binding model predicts 52.5% of the peptide is in the P_{19}^{40} complex and 35% is in the P_{19-40}° complex. Note that the $3^{1}P$ resonances of the pS40 phosphate groups in these two $31P$ resonances of the pS40 phosphate groups in these two complexes are in fast exchange (Fig. $2 B$), resulting in a single $pS40^c$ peak [\(Fig. 2](#page-3-0) A).

DISCUSSION

In the prevailing model of 14-3-3 binding to doubly phosphorylated partners, the partners contain a high affinity, sometimes called a "gatekeeper" residue, which initially interacts with 14-3-3, and a secondary, weaker epitope, which only binds when the gatekeeper site is already bound to 14- 3-3 [\(23](#page-9-0)). hTH1 that is singly phosphorylated on Ser^{40} is thought to be unable to bind $14-3-3\zeta$, whereas the pS19 phosphate is the high affinity epitope [\(18,19\)](#page-9-0). Our NMR results with the N-terminal 50 residue peptide, pS40-hTH1– 50, indicate, however, that binding of $pS40$ to $14-3-3\zeta$ is indeed possible, as evidenced by the change from the free peptide phosphorous resonance to the bound one. This interaction is clearly weaker $(K_d = 1.0 \pm 0.1 \text{ mM})$ than that between $14-3-3\zeta$ and $pS19-hTH1–50$, containing the purported gatekeeper phosphate, which binds a factorof-10 tighter ($K_d = 0.15 \pm 0.03$ mM). Therefore, even under conditions where $14-3-3\zeta$ is present in excess over the

 $pS19pS40-hTH1-50$ peptide, two complexes, P_{19}^{40} and P_{0}^{\odot} = 40, form in a 60:40% (K_{eq} = 1.5) ratio, irrespective of the 14.3.3% concentration. Our results show that it is the $14-3-3\zeta$ concentration. Our results show that it is possible that multiple complexes for doubly phosphorylated binding partners of $14-3-3\zeta$ may exist, even if in the phosphorylated full-length hTH1 additional mechanisms may come into play and alter the relative binding affinities ([18\)](#page-9-0).

Indeed, for the full-length pS40-hTH1, the binding to 14- $3-3\gamma$ was almost undetectable by native mass spectroscopy and surface plasmon resonance, while for the doubly phosphorylated enzyme Ser⁴⁰ is involved in binding to 14-3-3 γ ([22\)](#page-9-0). In addition, binding of one $14-3-3\gamma$ dimer to one of the monomers in tetrameric hTH1—which is organized as a dimer of dimers—appears to interfere with the binding of an additional 14-3-3 γ dimer to the adjacent monomer in hTH1, but not to the dimer at the opposite end, as seen in electron micrographs of the complex (22) (22) . Thus, in the complex involving the full-length enzyme, steric factors appear to be responsible for moving the equilibrium toward a scenario where the pS19 site binds to a single monomer in the 14-3-3 dimer. Whether this is the case for all 14-3-3 interactions with phosphorylated regulatory regions is unclear. In general, the presence of additional interactions and their contributions to the overall regulatory mechanisms have to be ascertained and confirmed for each individual system.

At this point, it may be of interest to analyze the underlying thermodynamics that govern the observed interaction between $14-3-3\zeta$ and $pS19pS40-hTH1-50$ (or other similar peptides, denoted as $1-2$, with two phosphorous sites), with the P_1^2 bound form as the dominant complex when 14-3-3 ζ is in excess and P_{1-2}^{1-2} dominant when peptide is in excess.
For simplicity we assume that the binding of site 1 (primary For simplicity, we assume that the binding of site 1 (primary site) is independent of that of site 2 (secondary site) and that peptide binding to one binding site on the $14-3-3\zeta$ dimer leads to a global entropy decrease ($\Delta S_G < 0$), whereas engagement of the second site on the same peptide only results in a local entropy decrease ($\Delta S_L < 0$). The enthalpy changes corresponding to the binding of the primary and secondary sites to 14-3-3 protein are denoted as ΔH_1 and ΔH_2 . Given these assumptions, the condition where the amount of P_{1-2}^{1-2} is larger than P_{1}^{2} (when peptide is in excess over $14-3-3\frac{z}{r}$ $r < 1$) i.e. over 14-3-3 ζ ; $r < 1$), i.e.,

$$
\Delta G(P_{1-2}^{1-2})-\Delta G(P_1^2)|<0,
$$

results in

$$
\Delta H_1 - \Delta H_2 < T(\Delta S_G - \Delta S_L). \tag{6}
$$

The opposite inequality

$$
(\Delta H_1 - \Delta H_2 > T(\Delta S_G - \Delta S_L))
$$

is expected for the case when the amount of P_1^2 is larger than P_{1-2}^{1-2} (when $r > 1$). In the case that the two phosphorous groups exhibit comparable binding affinities (e.g., such is the case for the PKC ε -V3 peptide that contains tandem repeat sequences ([27\)](#page-9-0)),

$$
\Delta G(P_1^2) - \Delta G(P_{1-2}^{1-2}) = (\Delta H_2 - \Delta H_1) + T(\Delta S_G - \Delta S_L).
$$

Because $\Delta H_1 - \Delta H_2 \sim 0$ and $T(\Delta S_G - \Delta S_L) < 0$, the free energy difference

$$
\left(\Delta G(P_1^2)\right)-\Delta G(P_{1-2}^{1-2})\right)
$$

becomes negative, making the amount of P_1^2 complex larger than the amount of P_{1-2}^{1-2} complex, irrespective of the *r* value.
Considering the large number of 14.3.3 binding partners

Considering the large number of 14-3-3 binding partners and the above-described different binding modes, both cases are thermodynamically possible and may contribute to the regulation imparted by 14-3-3 on its targets.

CONCLUSIONS

A detailed analysis of the interaction between singly or doubly phosphorylated hTH1–50 peptides and 14-3-3 ζ is presented. Using phosphorous NMR, dissociation constants for the binding of singly phosphorylated peptides, pS19 hTH1–50 and $pS40-hTH1-50$, to 14-3-3 ζ were determined. They differ by a factor of 10, with the Ser^{19} phosphorylated peptide exhibiting a higher affinity ($K_d = 0.15 \pm 0.03$ vs. 1.0 ± 0.1 mM). Furthermore, for the doubly phosphorylated peptide, pS19pS40-hTH1–50, a mixture of distinct complexes was observed, which permitted us to derive a comprehensive equilibrium binding scheme. All six possible different complexes are taken into account, and the system is described by three equilibrium constants: $K_{d1} = 0.15$ mM, $K_{d2} = 1.0$ mM, and $K_{eq} = 1.5$. For excess pS19pS40-hTH1–50 peptide, the major complex is the 14- 3-3z dimer with two peptides bound via the phosphate group on Ser¹⁹ (P_{19-40}^{19-40}), whereas, for excess 14-3-3 ζ , two major
complexes are seen each one with a doubly phosphorylated complexes are seen, each one with a doubly phosphorylated peptide bound to one 14-3-3 ζ dimer, with the Ser¹⁹ phosphate group bound to one $14-3-3\zeta$ subunit, and the Ser⁴⁰ phosphate group either bound to the second subunit (P_{19}^{40})
or free in solution (P°_{1}) in a 60:40 ratio or free in solution $(P_{19-40}^{\circ\circ})$ in a 60:40 ratio.
For other systems, the underlying therm

For other systems, the underlying thermodynamics will contribute to whether a doubly phosphorylated partner will bind with a 1:1 stoichiometry to the $14-3-3\zeta$ dimer or if a mixture of complexes is formed. It is likely that most 14-3-3z binding partners will possess phosphorylated residues that exhibit different affinities for the binding sites on 14-3-3z. The presented methodology derived by monitoring individual $3^{31}P$ resonances has general applicability for any complex system involving 14-3-3 and doubly phosphorylated ligands, and it will be interesting to extend this approach to full-length hTH1 or other target proteins.

SUPPORTING MATERIAL

Three figures are available at [http://www.biophysj.org/biophysj/supplemental/](http://www.biophysj.org/biophysj/supplemental/S0006-3495(14)01012-1) [S0006-3495\(14\)01012-1.](http://www.biophysj.org/biophysj/supplemental/S0006-3495(14)01012-1)

AUTHOR CONTRIBUTIONS

J.H., A.M., and A.M.G. conceived this study; J.H. and I.-J.L.B. acquired experimental data; J.H., T.K., I.-J.L.B., and A.M.G. performed data analysis and interpretation; J.H., I.-J.L.B., and A.M.G. drafted the article; all authors contributed to the final article.

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Figure 1

A SEW MPTPDATTPQ¹⁰ AKGFRRAVSE²⁰ LDAKQAEAIM³⁰ SPRFIGRRQS⁴⁰ LIEDARKERE⁵⁰