Negative Charges, Not Necessary Phosphorylation, are Required for Ligand Recogni-

tion by 14-3-3 Proteins

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Running Title: Charge Contribution for Recognition of 14-3-3 Proteins

Abstract: Protein-protein interactions involving 14-3-3 proteins regulate various cellular activities in normal and pathological conditions. These interactions have mostly been reported to be phosphorylation-dependent, but the 14-3-3 proteins also interact with unphosphorylated proteins. In this work, we investigated whether phosphorylation is required, or, alternatively, whether negative charges are sufficient for 14-3-3ε binding. We substituted the pThr residue of pT(502-510) peptide by residues with varying number of negative charges, and investigated binding of the peptides to 14-3-3ε using MD simulations and biophysical methods. We demonstrated that at least one negative charge is required for the peptides to bind 14-3-3ε while phosphorylation is not necessary, and that two negative charges are preferable for high affinity binding.

Keywords: phosphopeptides; 14-3-3 proteins; binding motif; phosphate mimics.

Abbreviations: ASK1, apoptosis signal-regulating kinase 1; CCAAT, cytosine-anenosineadenosine-thymidine; CDC25A, cell division cycle 25 A; cSCC, cutaneous squamous cell carcinoma; DSF, differential scanning fluorimetry; ISA, interfacial surface area; LSCC, larynx squamous cell carcinoma; MD, molecular dynamics; SASA, solvent accessible surface area; SOS1, son of sevenless-1; SPR, surface plasmon resonance; VDW, van der Waals.

Introduction

Protein phosphorylation, one of the most common post-translational modifications, plays an important role in the regulation of cellular functions. Protein kinases commonly phosphorylate seryl, threonyl and tyrosyl residues of proteins, and this modification not only affects structural properties of the proteins but also their functions [1–3]. Phosphorylation modulates protein-protein interactions, and some proteins recognize binding motifs containing phosphorylated amino acid residues [4–8].

14-3-3 proteins are a family of adapter proteins that are ubiquitously expressed in eukaryotic cells. The 14-3-3 proteins have many binding partners with a broad range of functions. Thus, these proteins are involved in various cellular activities in both physiological and pathological conditions [9–12]. The 14-3-3ε isoform is upregulated and linked to abnormal cell growth in renal cancer [13]. Reduced expression of 14-3-3ε has been reported in gastric cancer [14]. The 14-3-3 proteins have also been associated with progression of larynx squamous cell carcinoma (LSCC) [15], and small cell lung cancer [16]. The 14-3-3 isoforms, ε, ζ and γ are expressed and mislocalized to cytoplasm in cutaneous squamous cell carcinoma (cSCC), where a heterodimer of the ε with either ζ or γ interacts with the cell cycle regulator, cell division cycle 25 A (CDC25A) to suppress apoptosis [17–19].

It is widely reported that interactions between 14-3-3 proteins and their binding partners are through binding motifs comprising a phosphoseryl (pSer) or a phosphothreonyl (pThr) amino acid residues [20,21]. The 14-3-3 proteins recognize two binding motifs, RSXp-SXP and RXY/FXpSXP; pS is pSer, and X is any amino acid [20]. Apoptosis signal-regulating kinase 1 (ASK1) [22], Raf-1 [23,24], CDC25A [17], cytosine-anenosine-adenosine-thymidine (CCAAT) -enhancer binding protein [25], and cystic fibrosis transmembrane conductance regulator [8] are among the phosphorylated binding partners of 14-3-3 proteins. However, not all 14-3-3 interactions with their binding partners are phosphorylation-dependent, as they also bind proteins such as CDC25B [26] and exoenzyme S [27,28] in a phosphorylation-independent manner.

Over the years, the binding mechanism of various phosphopeptides to 14-3-3 proteins has been studied using experimental and computational methods [20,29–31]. Muslin and colleagues [24] showed the phosphorylation-dependent binding of the Raf-1 derived peptide (pS-Raf-259) to 14-3-3ζ. The peptide occupies the amphipathic binding groove of 14-3-3ζ, and residues of the protein interacting with the peptide were identified [32], similar interaction were identified between 14-3-3ζ and phosphorylated myeloid leukemia factor 1 peptide [33]. Furthermore, the human SOS1-derived peptide binds 14-3-3ζ in a similar manner as pS-Raf-259 does [34]. In our previous work [19,30,31], we also developed CDC25A derived phosphopeptides that bind 14-3-3ε, and we identified basic amino acid residues and aromatic residues of 14-3-3ε (Lys⁵⁰, Arg⁵⁷, Arg¹³⁰, Tyr¹³¹) that interact with phosphorylated amino acid residues.

Phosphorylation-independent interactions between 14-3-3 proteins and peptides have also been elucidated. R18, the unphosphorylated peptide that has two negatively charged amino acid residues, discovered by phage display designed for 14-3-3τ binding has been shown to interact with 14-3-3 proteins in a manner reminiscent of phosphorylated peptides. Although, R18 has additional hydrophobic interactions with the protein [23,32]. Ottmann and colleagues [27] demonstrated that the exoenzyme S derived peptide interacts with 14-3-3 protein through ionic and hydrophobic interactions.

We have developed phosphopeptides that inhibit interactions between CDC25A and 14-3-3ε, and induce apoptosis of cSCC in animals [19]. We further improved binding affinities of the peptides for 14-3-3ε by 6.5 fold [30]. Although these phosphopeptides have high affinities for 14-3-3ε, it is likely that they are susceptible to dephosphorylation as various phosphatases have been shown to dephospharylate phosphopeptides in solution [35–37]. Therefore, we proposed replacing the pThr residue of the parent peptide (pT(502-510)) with pThr mimics to potentially improve their resistance to phosphatases. It has long been known that 14-3-3 protein interactions with their binding partners are dependent on phosphorylation of the binding partner. Thus, we aimed to elucidated whether the binding of peptides are dependent on the presence of negatively charged amino acid residues, and not necessarily their phosphorylation.

Here, we studied binding between 14-3-3ε and peptide analogs in which the pThr residue in our previously studied peptide, pT(502-510) [30], was substituted with residues carrying varying number of negative charges. We used biophysical methods to show that at least one negative charge is required for peptide binding to 14-3-3ε.

Materials and Methods

Molecular Dynamics (MD) simulations

Peptide – 14-3-3ε complexes preparation

The starting structure was obtained from our previously studied $14-3-3\varepsilon - pT(502-510)$ complex $[30]$. To obtain the various peptide analogs, $pThr⁵⁰⁷$ amino acid residue in the pT(502-510) was substituted with Thr, Glu, Gla, sThr or Pmb amino acid residues (Figure 1).

Non-standard residues were built in the YASARA program [38]. Binding of the peptides to 14- 3-3ε was studied using MD simulations.

MD simulations of peptide – 14-3-3ε complexes

MD simulations of the peptide – protein complexes were performed using the AMBER-FB15 force field as implemented in YASARA [39]. Initially, the 14-3-3ε – peptide complex was solvated with water in a cubic simulation box with minimal distance of 1.2 nm between the edge of the box and the complex. Then the system was energy minimized using standard parameters in YASARA. Then the energy-minimized structures of 14-3-3ε - peptide complexes were solvated again with water molecules and 150 mM NaCl, and the structure of the system was simulated using *md_run* macro of YASARA. The 500 ns simulations were performed at 310 K and 1 atm pressure, and the trajectories were saved in xtc format that is compatible with GROMACS [40]. The simulations were performed in two steps, (1) a 1 ns simulation using YASARA macro in the YASARA graphical user interface, (2) then the simulation was continued in a computer background to generate a 499 ns trajectory using *md_run* macro of YASARA.

Trajectory analysis

Using YASARA, the first frames of the trajectories were converted to PDB format and used as topology files for analysis using analysis utilities of GROMACS 2022.5 [40]. The trajectories were processed using *trjconv* module of GROMACS. The *rms* module of GROMACS was used to calculate Cα atoms root-mean-square deviation (RMSD). VMD software [41] was used to calculate salt-bridges and H-bonds between peptides and 14-3-3ε; 0.8 nm was used as the oxygen–nitrogen cut-off distance for salt bridges, and the donor–acceptor distance was

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set to 0.4 nm for H-bonds. The *sasa* module of GROMACS was used to calculate the solvent accessible surface area (SASA). The interfacial surface area (ISA) [42] was calculated as follow:

 $ISA = (SASA_{14-3-3ε} + SASA_{peptide}) - SASA_{14-3-3ε-peptide complex}$. The van der Waals (VDW) surface area of residues were calculated using YASARA.

Peptides.

The [sThr⁵⁰⁷]pT(502-510) peptide was from Biosynth International Inc.(Louisville KY, USA), other peptide analogs were synthesized in house. All *N*-α-Fmoc-protected amino acids were from CEM Corporation (Matthews NC, USA), except, *N*-alpha-(9-Fluorenylmethyloxycarbonyl)-gamma-Carboxy-L-glutamic-acid-gamma-di-t-butyl ester, (Fmoc-L-Gla(OtBu)2-OH) and (2S,3R)-2-((9-fluorenylmethyloxycarbonyl)amino)-4-(di-t-butylphosphonomethyl)-3 methylbutyric acid, (Fmoc-L-Pmb(tBu)2-OH) that were from Iris Biotech (Adalbert-Zoellner-Str 1 D-95615 Marktredwitz, Germany). Dimethylformamide (DMF) and *N*,*N′*-Diisopropylcarbodiimide (DIC) were from Sigma-Aldrich, the Rink amide protide resin and Oxyma Pure were from CEM.

Peptide synthesis

The peptides were synthesized using CEM Liberty blue 2.0 microwave peptide synthesizer, in a 0.1 mmol scale on Rink amide protide resin using standard Fmoc chemistry with Boc, tBut, Bzl, and Pbf side chain protections. A 1 min Fmoc deprotection was carried out using 20% piperidine in DMF, and 2 min coupling was performed in DIC/Oxyma Pure at 90 °C, except for the (Fmoc-L-Pmb(tBu)2-OH) that was coupled for 4 min. Peptide cleavage from the resin and side chains deprotection were simultaneously performed by stirring the peptide-

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resin in a cocktail containing TFA/Thioanisol/phenol/TIS/H2O/DODT in 81.5:5:5:1:5:2.5 (v/v/ v/v/v/v) for 30 min on ice and then 3.5 hours at room temperature. The resin was then separated from peptide by filtration and the peptide was precipitated using ice-cold ether. Crude peptides were purified by reverse-phase HPLC using C8 column (Phenomenex, Aeris 100 Å, 5 μM , 250 x 10 mm). A 0.1% TFA aqueous solution (v/v solvent A) and a 0.09% TFA acetonitrile (solvent B) were used to elute the peptides with a linear gradient of solvent B of 3 to 60% over 60 min at a flow rate of 4 mL/min. Identities of the peptides were confirmed by Thermo Q-Exactive Orbitrap mass spectrometer.

Differential scanning fluorimetry (DSF)

The (His)6-14-3-3ε was from Novus Biologicals (Centennial, CO, USA). Thermal unfolding of 14-3-3 protein and 14-3-3 – peptide complexes were performed as previously described [30]. Briefly, thermal unfolding was monitored using SYPRO-orange dye (Invitrogen, Carlsbad, CA, USA) in BioRad CFX384 Touch real-time PCR instrument (Hercules CA, USA). The protein and protein-peptide complexes were dissolved in TRIS buffer (150 mM NaCl, 20 mM TRIS, 1 mM DTT, pH 7.4) to the final concentrations of 30 μM of peptide, 2 μM of 14-3- 3ε, and 1000x dilution from SYPRO-Orange dye stock from the manufacturer. Melting temperature (T_m) was determined from the first derivative of unfolding curve and change in melting temperature (ΔT_m) was calculated as the difference between T_m of protein-peptide complex and T_m of protein alone. At least 3 replicates were performed per peptide.

Surface plasmon resonance (SPR)

To obtain binding affinity (K_D) of peptides for 14-3-3ε, SPR was performed as before [30]. Briefly, recombinant His6-14-3-3ε was immobilized on activated Biacore NTA chip (Cy-

tiva, Marlborough MA, USA), at a flow rate of 5 μL / min for 10 min and then washed with the running buffer (20 mM Tris, 150 mM NaCl, 50 μM EDTA, 0.005% Tween 20, pH 7.4) for 20 min. Following protein immobilization, solutions of peptides were injected at increasing concentrations ranging from 4 to 500 uM (3.9,7.8, 15.6, 31.2, 62.5, 125, 250 and 500 μM) for all analogs, except [Pmb⁵⁰⁷]pT(502-510) where concentrations ranging between 0.002 μM and 0.25 μM (0.002, 0.004, 0.008, 0.016, 0.031, 0.063, 0.125 and 0.25 μM) were used. The injections were carried out at a flow rate of 30 μL/min for 1 min, and dissociation time was 3 min in the running buffer. Steady state affinities of the peptide were obtained by fitting the data using Biacore Insight Evaluation Software (version 5.0.18.22102). For each peptide, at least 5 replicates were done.

Results

 In the present work, we elucidated the binding of peptide analogs of our previous studied peptide pT(502-510) [30], in which the pThr amino acid residue in position 507 was substituted with amino acid residues with uncharged Thr, the singly charged sulfothreonine (sThr) and Glu, or with the doubly charged γ-carboxyl glutamic acid(Gla) and 2-Amino-3-methyl-4 phosphonobutanoic acid (Pmb) residues (Table 1). Their chemical structures are shown in Figure 1. The sThr is a phosphatase resistant pThr mimetic with one negative charge, and Pmb is a phosphatase resistant pThr mimetic with two negative charges. For comparison, pThr residue, in our published work, [30] is also shown. Binding of each peptide analog to 14- 3-3ε was computationally investigated in 500 ns MD simulations.

The Cα-atoms RMSD indicated that the peptides did not dissociate from 14-3-3ε – peptide complexes throughout simulations (Figure S1). All 14-3-3ε – peptide complexes underwent structural rearrangement with RMSD between 0.15 nm and 0.4 nm, except the $14-3-3\varepsilon$ – $[Gla⁵⁰⁷]pT(502-510)$ complex that had RMSD of 0.15–0.6 nm (Figure S1). In all complexes, peptides showed less fluctuation than $14-3-3\varepsilon$, except the $[Pmb⁵⁰⁷]pT(502-510)$ peptide that showed higher fluctuation between 250 ns and 450 ns. Analysis of the interfacial surface area (ISA) between 14-3-3ε and peptide showed that the 14-3-3ε – peptide complexes had ISA of 13.7–17.3 nm² (Table 1). The $[Pmb⁵⁰⁷]pT(502-510)$ containing complex showed the largest ISA, while the [sThr⁵⁰⁷](502-510) containing complex had the smallest ISA. This suggests that all the peptides had a degree of interaction with 14-3-3ε during simulations. We further determined VDW surface area of the residue in position 507 of each peptide in each 14-3-3ε – peptide complexes. The results indicated that VDW surface area of the residues was between 1.386 nm² and 1.954 nm² (Figure S2). The isosteric replacements, Pmb, Gla, and sThr residues had VDW surface area of 1.954 nm², 1.824 nm² and 1.816 nm², respectively, and the nonisosteric replacements, Glu and Thr had VDW surface area of 1.605 nm² and 1.386 nm², respectively.

We identified several residues of the 14-3-3ε - peptide complex that interact with residues introduced in position 507 (Figure 2, and Table S1). In the 14-3-3ε – [Thr⁵⁰⁷]pT(502-510) complex, no interactions involving $Thr₅₀₇$ residue were observed. However, residues Arg⁵⁰², Lys⁵⁰⁴, Ser⁵⁰⁵, Arg⁵⁰⁶ and Trp⁵⁰⁸ of the peptide formed H-bonds with residues Lys¹²³, Glu¹³⁴, Glu¹⁸³, Asn¹⁷⁶, Asp²²⁶ and Asn²²⁷ of 14-3-3ε (Table S1). In all other 14-3-3ε – peptide complexes, H-bonds were observed between the 507 residue of the peptides and Lys⁵⁰, Arg⁵⁷, Arg¹³⁰ and Try¹³¹ of 14-3-3 ε (Table S1). Pmb residue in the [Pmb⁵⁰⁷]pT(502-510) peptide

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formed the longest lasting H-bonds with these residues of 14-3-3ε, followed by Gla residue in $IGIa⁵⁰⁷IpT(502-510)$. Additionally, Pmb formed H-bonds with Asn¹⁷⁶, Leu¹⁷⁵, Val¹⁷⁹ and Leu²²³ (Figure 2, and Table S1). The singly charged sThr and Glu in $[sThr⁵⁰⁷]pT(502-510)$ and $[Glu⁵⁰⁷]pT(502-510)$, respectively, showed less lasting H-bonds with Lys⁵⁰, Arg⁵⁷, Arg¹³⁰ and Try131 residues of 14-3-3ε, than Gla and Pmb. Analysis of ionic interactions revealed no interactions between the 507 residues of the peptides with residues of the 14-3-3ε, except the Glu residue in [Glu⁵⁰⁷]pT(502-510) peptide that formed salt-bridges with Lys⁵⁰, Lys¹²³, Arg¹³⁰.

The binding of synthetic peptides to 14-3-3ε was initially assessed using DSF. The unfolding curves of 14-3-3 – peptide complexes and 14-3-3ε, and their corresponding first derivatives are shown in Figures 3 and S3. Analysis of DSF results indicated that [Pmb⁵⁰⁷]pT(502-510) peptide caused a ΔT_m of 14-3-3ε of 3.6°C, while all other peptides showed ΔT_m of 0.0°C. SPR was used to quantitatively determine K_D of the peptides to 14-3-3ε. Figures 4 and S4 show steady-state binding isotherm, and sensograms of the peptides binding to 14-3-3ε. The results showed that $[Thr^{507}](502-510)$ peptide did not bind to 14-3-3ε. $[Glu^{507}]pT(502-510)$, [sThr^{507}]pT(502-510), and [Gla^{507}]pT(502-510) had K_D of 141.75 ± 24.96 μM, 86.71 ± 8.07 μM, and 24.06 \pm 8.43 µM, respectively. [Pmb⁵⁰⁷]pT(502-510) peptide had the highest affinity of $0.009 \pm 0.001 \,\mu M$ (Table 1).

Discussion

Interactions of 14-3-3 proteins with their binding partners have been primarily attributed to phosphorylated residues in the partners. However, 14-3-3 proteins have also been reported to bind proteins in phosphorylation-independent manner [24,27]. Therefore, here we studied the role of negative charges in binding partners for interaction with 14-3-3ε. We showed that phosphorylation is not necessary for binding of 14-3-3ε, but single or double negative charges are required. We used MD simulations to determine the stability of $14-3-3\varepsilon$ – peptide complexes.

All the peptides formed stable complexes with 14-3-3ε; no peptide dissociation was observed (Figure S1). The structural rearrangement was comparable to that of the phosphorylated peptides [30]. ISA calculations can be used to reveal interactions between binding partners [43] Our ISA results qualitatively confirmed that all the peptide had interactions with 14- 3-3ε in the protein – peptide complexes. The Pmb residue of the highest affinity peptide, [Pmb⁵⁰⁷]pT(502-510), had the largest ISA, while the residues from non-binding and weakly binding peptides had smaller ISA.

 Since the residues of 14-3-3 proteins that interact with the phosphorylated amino acid residues of their binding partners are known [32], we sought to determine if these interactions were maintained after residue substitutions in the parent peptide. In agreement with our previous results [19, 30], analysis of H-bonds involving pThr⁵⁰⁷ residue in the 14-3-3ε – pT(502-510) complex confirmed that the residues interacted with Arg¹³⁰, Tyr¹³¹, Arg⁵⁷ and Lys⁵⁰ for the most part of the simulation time (Table S1). As expected, the uncharged residue Thr in $[Thr⁵⁰⁷]pT(502-510)$ did not interact with the reported residues of the 14-3-3ε (Table S1). Hbond analysis further showed that the singly charged residues $sThr⁵⁰⁷$ and Glu⁵⁰⁷ in [sThr⁵⁰⁷] (502-510) and [Glu⁵⁰⁷]pT(502-510) peptides, respectively, exhibited short-lasting H-bonds with the residues, except the Arg¹³⁰ residues that lasted for more than 85% of the simulation time. Gla⁵⁰⁷ of the $IGIa^{507}$]pT(502-510) peptide interacted with Arg¹³⁰ and Tyr¹³¹ for > 95% of the simulations time, but temporarily also interacted with Arg⁵⁷ and Lys⁵⁰. Like the pThr⁵⁰⁷, Pmb⁵⁰⁷ residue in $[Pmb⁵⁰⁷]pT(502-510)$ peptide formed long-lasting interactions with all the four

residues. The Pmb showed additional H-bonds with Asn¹⁷⁶, Val¹⁷⁹, Leu¹⁷⁵ and Leu²²³. The Hbonds by residues in position 507 were formed by negatively charged side chain oxygen(s), this explains why no H-bonds were detected with Thr 507 residue of $pT(502-510)$ peptide, as well as the more and longer lasting interaction by the doubly charged residues than the singly charged residue. Pmb residues and pThr have larger VDW surface area (Figure S2), and subsequently larges ISAs, suggesting that besides charge, the larger of the side chain of residues in position 507 of the peptides might be of importance for the peptide binding to 14- 3-3ε.

DSF studies showed that the $[Pmb⁵⁰⁷]pT(502-510)$ peptide caused a shift of 3.6°C, while other peptides did not cause any shift in T_m of 14-3-3ε (Figure S3) at 30 μM. The concentration at which the parent peptide and its other analogs caused change in T_m of 14-3-3ε. Similar to DSF, in SPR experiments, the [Thr⁵⁰⁷]pT(502-510) did not bind 14-3-3ε at the studied concentrations (0–500 μM). This is in agreement with the fact that no interactions were observed between Thr⁵⁰⁷ residue of $pT(502-510)$ peptide and residues of 14-3-3ε. This also indicates the crucial importance of negatively charged amino acid residues in peptide binders of 14-3-3ε. The [sThr⁵⁰⁷](502-510) and [Glu⁵⁰⁷]pT(502-510) had K_D of 86.71 ± 8.07 µM and 141.75 \pm 24.96 µM, respectively. The peptide containing the doubly charged Gla, [Gla⁵⁰⁷]pT(502-510) showed a higher affinity (K_D of 24.06 \pm 8.43 µM) (Table 1). Furthermore, the peptide with the doubly charged residue (Pmb) that has a phosphono group instead of phosphate group, $[Pmb^{507}]pT(502-510)$ showed the highest affinity (K_D of 0.009 \pm 0.001 µM) for 14-3-3ε than all other analogs. $[Pmb⁵⁰⁷]pT(502-510)$ also has higher affinity for 14-3-3ε than our previously reported best binder for 14-3-3ε [30]. The biophysical results are in agreement with our computational results showing stronger interactions between the

[Pmb⁵⁰⁷]pT(502-510) peptide and 14-3-3ε. Similar to H-bond analysis, the [Gla⁵⁰⁷]pT(502-510) peptide showed weaker affinity than $[Pmb⁵⁰⁷]pT(502-510)$ yet both Gla and Pmb are doubly charged, this could be due to different distributions of electrons on the side chain of these residues, resulting in different location of net charges, as well as the slight difference in their VDW surface areas.

Although binding of 14-3-3 with other proteins has been mostly reported with phosphorylated binding partners [20,21,44], it also interacts with unphosphorylated proteins like exoenzyme S [27,28], and unphosphorylated peptides [23,32]. In a study conducted by Petosa and colleagues [32] comparing binding of the unphosphorylated pan inhibitor of 14-3-3 proteins, R18 and the RAF-1 derived phosphopeptide pS-RAF-1-259, a crystal structure of 14-3- 3ζ with R18 showed that a pentapeptide fragment of R18 (Trp-Leu-Asp-Leu-Glu) occupies binding site that overlaps with that of the phosphorylated peptides. The pentapeptide fragment contains two acidic amino acid residues (Asp and Glu), in the crystal structure, these residues are next to Lys⁴⁹, Arg⁵⁶, Arg⁶⁰, and Arg¹²⁷ residues of 14-3-3 ζ , the phosphate group of pS-RAF-1-259 peptide is next to the same residues. In a different study on a peptide that contains two Asp residues [27], one Asp is located far (0.48 – 0.573 nm) from the basic residues of 14-3-3ζ to form significant interactions, the other Asp residue contacts Lys⁴⁹. These studies are in agreement with our previous studies in which we reported similar residues (Lys⁵⁰, Arg⁵⁷, Arg¹³⁰ and Tyr¹³¹) of 14-3-3ε to interact with the phosphorylated residue in our phosphopeptide analogs [19,31]. The same residues of 14-3-3ε interacted with the 507 residues in the peptides presented in this work.

In the current work, we demonstrated that negative charge(s), not necessarily phosphorylation, are required for binding to 14-3-3 proteins and that two negative charges are fa-

vored for high affinity binding. The isosteric replacement of pThr with sThr with one negative charge on the side chain resulted in weak binder peptide. We also designed a peptide, [Pmb⁵⁰⁷]pT(502-510), having two negatively charged side chain, that has the highest, nanomolar, affinity for 14-3-3ε. In our previous work we designed phosphopeptide analogs that inhibit squamous cell carcinoma cell growth [19]. Unlike these pervious analogs, the [Pmb⁵⁰⁷]pT(502-510) is likely more stable, since it is not susceptible to dephosphorylation, making it a better candidate for inhibition of CDC25A – 14-3-3ε interactions to promote apoptosis of cSCC cells.

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Figure 1. **Structure of amino acid residues**. Amino acid residues in position 507 of the peptide analogs have varying negative charge number from 0 to -2. pThr, Phosphothreonyl; Thr, threonyl; sThr, sulfothreonyl; Glu, glutamic acid residue; Gla, γ-carboxyglutamic acid residue; Pmb, 2-Amino-3-methyl-4-phosphonobutanoic acid residue.

Figure 2. Residues of 14-3-3ε interacting with Pmb⁵⁰⁷ . Left, structure of 14-3-3ε- [Pmb⁵⁰⁷]pT(502-510) complex. The secondary structure of protein and peptide is shown in gray ribbon and red tube, respectively. Interacting residues are in stick representation. Hydrogen, white; carbon, cyan; nitrogen, blue; oxygen, red and phosphor, yellow. Right, a close-up view showing residues of 14-3-3ε that form H-bonds with Pmb⁵⁰⁷ residue of [Pmb⁵⁰⁷]pT(502-510) peptide. Residues of 14-3-3ε are indicated with one letter code, and Pmb⁵⁰⁷ indicated by letter Z. H-bonds were identified within 0.4 nm distance between acceptor and donor atoms and indicated by green dashed line.

Figure 3. Thermal unfolding of 14-3-3ε protein, and its complexes with peptides. Thermal unfolding curve of 14-3-3ε without (black) and with peptide (red), and their respective first derivatives from differential scanning fluorimetry (DSF). Left panel, [Pmb⁵⁰⁷]pT(502-510); right panel, [sThr⁵⁰⁷]pT(502-510). The binding of the peptide is determined as the change in melting temperature (T_m) between 14-3-3ε – peptide complex and 14-3-3ε alone.

Figure 4. Surface plasmon resonance-based dose response binding of peptides to 14- 3-3ε. A and B, sensograms and their corresponding binding isotherm of [Pmb⁵⁰⁷]pT(502-510); C and D, [sThr⁵⁰⁷]pT(502-510). The arrows indicate binding response of increasing concentrations of the peptides, concentrations range between 0.002 μM and 0.25 μM for $[Pmb⁵⁰⁷]pT(502-510)$, and 4 μM and 500 μM for $[SThr⁵⁰⁷]pT(502-510)$. 100 nM of 14-3-3ε was used in all experiments.

Table 1. Binding affinities of the peptides for 14-3-3ε, and interacting surface area (ISA)

between 14-3-3ε and the peptides. Residues that substituted pThr of the parent peptide $(pT(502-510))$ are highlighted in red. Ac, and NH₂, acetyl and amide, respectively, protecting groups. K_D values are average \pm SD of n \geq 5.

