Recognition of high-risk HPV E6 oncoproteins by 14-3-3 proteins studied by interactomics and crystallography

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Abstract

In tumors induced by high-risk mucosal human papillomaviruses (hrm-HPVs), HPV E6 oncoproteins inhibit apoptotic processes and sustain cell proliferation. E6 from all hrm-HPVs harbor a C-terminal short PDZ domain-binding motif (PBM), whose phosphorylation down-regulates PDZ binding but triggers E6 binding to 14-3-3 proteins. Here we classify PBMs of E6 proteins depending on their principle ability to be phosphorylated and subsequently acquire a 14-3-3-binding motif III consensus, (pS/pT)XX-COOH. Systematic competitive fluorescence polarization measurements show that the PBMs from four selected E6 oncoproteins bind all seven human 14-3-3 isoforms with distinct, wide-ranging affinities, obeying remarkable trends assigned to 14-3-3 isoform specificity and small E6 sequence variations. We crystallized the hrm-HPV18 E6 PBM bound to 14-3-3ζ, revealing a 14-3-3-motif III complex at 1.9 Å resolution. Using fluorescence polarization and crystallography, we also demonstrate that fusicoccin, a molecule that reinforces many known 14-3-3 complexes, destabilizes the 14-3-3-E6 interaction, indicating the druggability of that complex.

Keywords:
Protein chimera; protein-protein interactions; phosphopeptides; phosphorylation; crystal structure; binding affinity

INTRODUCTION

Papillomaviruses (PVs) are a large group of small oncogenic DNA viruses infecting various vertebrates including humans (human papillomaviruses, HPVs). PVs target mucosal or cutaneous epithelium cells and induce their proliferation, to amplify their circular double-stranded DNA genome and package it into virion particles ready for a new infectious cycle. 225 HPV types currently annotated in the Papillomavirus Episteme (PaVE) database (https://pave.niaid.nih.gov/) are classified into five phylogenetically distinct genera (alpha, beta, gamma, mu and nu), of which alpha HPVs display almost exclusively mucosal tropism 1. While most HPV only generate benign proliferative events such as warts, lesions or condylomas, a subset of about 20 HPV types belonging to two alpha species have been associated to cancers and are therefore epidemiologically categorized as “high-risk” 2, 3, 4. HPV16 and 18 are the most prevalent, causing up to 80% of squamous cervical carcinomas 5, whereas HPV16 is also highly prevalent in HPV-positive tumors of the oropharynx and the anus 6.

The E6 protein is one of the two main HPV oncoproteins expressed at the early stage of infection. In HPV-transformed cells, E6 participates in counteracting apoptosis, altering differentiation pathways, polarity and adhesion properties, and thereby sustains cell proliferation 1, 7. Inhibition of E6 in HPV-positive cell lines results in the cell growth arrest and induces apoptosis or rapid senescence 8, 9, 10, 11.

E6 has been found to interact with numerous distinct cellular target proteins involved in a variety of cellular functions 12. Importantly, E6 proteins from distinct HPV species recognize distinct subsets of the full panel of potential E6 targets 13. This likely contributes to the particular biological traits of each HPV type in terms of tropism, viral cycle, or pathogenicity.
E6 proteins of most mammalian PVs comprise two zinc-binding domains (E6N and E6C) separated by a linker helix \(^1\) (Fig. 1A), which altogether form a charged-hydrophobic pocket that recognizes LxxLL motifs interspersed with acidic residues, found in a variety of host cellular proteins \(^2\). Hrm-HPV E6 recruits simultaneously the LxxLL motif of ubiquitin ligase E6-Associated Protein (E6AP) and, via a distinct interface, the central “core” domain of tumor suppressor p53 \(^14\). The resulting E6-E6AP-p53 trimer drives the ubiquitinylation and subsequent proteasomal degradation of p53 \(^15\). In HPV-transformed cells, this leads to low levels of p53 protein that probably contribute to the tumorigenic phenotype. It is supposed that E6 similarly stimulates the E6AP-mediated degradation of other host proteins \(^15,16\).

In addition to their LxxLL-binding pocket and p53-binding surface, hrm-HPV E6 proteins harbor a C-terminal PDZ domain-binding motif (PBM) \(^17\) (Fig. 1A), which remarkably is absent in most low-risk non-cancer-causing HPV E6 proteins \(^18\). The intrinsically disordered PBM enables the E6 interaction with a variety of host PDZ domain-containing proteins involved in the regulation of cell polarity, adhesion, differentiation or survival \(^19\). These interactions of E6 with PDZ proteins are potentially modulated by PBM phosphorylation. Several E6 PBMs are phosphorylatable \textit{in vitro} at variable levels of specificity and efficiency by basophilic kinases, such as PKA \(^18,20,21\), PKB/Akt \(^18,21\), CamKII \(^21\), or Chk1, and by the glutamine-directed ATM/ATR kinases \(^22\). E6 PBMs from high-risk mucosal HPV types 16, 18, 33, 35 (Fig. 1B) are good substrates for PKA \textit{in vitro} \(^18,21,23\). In addition, E6 of HPV16 and HPV18 (hereinafter called 16E6 and 18E6) are metabolically labeled by \(^32\)P upon transient transfection \textit{in vivo}, and forskolin promotes such labeling \(^23\). Phosphorylation of several HPV-E6 proteins at the PBM was confirmed by using phospho-specific antibodies raised against the C-terminal PBM of 18E6 \(^18,21\). It was also shown that the level of 18E6 phosphorylation remains low in the normal cell cycle, until the activation of stress-response kinases and those involved in the DNA damage response (see in particular \(^22\)).

While phosphorylation of HPV-E6 PBMs generally reduces their binding to PDZ domains \(^24\), it turns several of them into candidate binding sites for 14-3-3 proteins \(^18,21\). Despite being encoded by seven distinct genes, human 14-3-3 proteins share highly similar sequences and biochemical properties. Hence they are commonly referred to as "isoforms", individually named \(\beta, \gamma, \varepsilon, \zeta, \eta, \sigma, \text{and } \tau\) (beta, gamma, epsilon, zeta, eta, sigma and tau) \(^25\), not to be confused with their splice variants. 14-3-3 proteins form homo- and heterodimers \(^26\) characterized by their ability to bind phosphopeptides \(^27,28\). 14-3-3 are involved in the regulation of multiple cellular processes including apoptosis, cell division and signal transduction \(^29\). Phosphorylated 14-3-3-binding sequences usually correspond to internal motifs \(\text{I RSX(pS/pT)XP(G)}\) and \(\text{II RXY/ FX(pS/pT)XP(G)}\) \(^28\) and to the C-terminal motif \(\text{III (pS/pT)}X_{0-2} \text{COOH} \) \(^30,31\), where pS/pT denotes phosphorylated serine or threonine and X denotes any amino acid. The regulation by 14-3-3 binding typically protects 14-3-3 targets from dephosphorylation and degradation, affects their activity, intracellular localization, and interactions with other proteins by occlusion of the phosphorylated segments \(^32\). Given the involvement of 14-3-3 in the development of various pathological conditions associated with the deregulation of the corresponding protein-protein interactions, 14-3-3/phosphotarget complexes are considered with increasing attention \(^33\).

Phosphorylated 16E6, 18E6, and 31E6 have been previously shown to bind to 14-3-3 proteins, albeit with different efficiencies \(^18,21,24\). In contrast, the binding of unphosphorylated or phosphomimetic E6 PBMs to 14-3-3 proteins was not detectable \(^21,24\).
This suggests that the phosphorylation-dependent interaction with 14-3-3 proteins is a general feature of various E6 PBMs.

Here we classified PBMs of hrm-HPV E6 proteins depending on their principle ability to be phosphorylated and subsequently acquire a 14-3-3-binding motif III consensus, (pS/pT)XX-COOH. We performed systematic quantitative binding assays of four phosphorylatable E6 PBMs with all seven human 14-3-3 isoform homodimers and solved a high-resolution crystal structure of the HPV18 E6 PBM complex with 14-3-3ζ. This combination of approaches has allowed us to rationalize differences in the affinities of tested PBMs and fully confirm this by mutated peptides. Finally, we used binding assays and crystallography to analyze the impact of natural toxin fuscoccin on the 14-3-3-E6 PBM interaction, which proved that the 14-3-3-E6 complexes are nominally druggable by small molecules.

**RESULTS**

PDZ-binding motifs of E6 proteins can be classified according to their phosphorylation and 14-3-3-binding propensity

Among all 225 HPV type E6 proteins curated in the PaVE database (https://pave.niaid.nih.gov/), 31 E6 proteins from the α-genera HPV, having predominantly mucosal tropism, possess a C-terminal PDZ-binding motif (PBM) that corresponds to the class 1 PBM [X(S/T)X(L/V/I/C)-COOH, where X is any amino acid residue]. It has previously been shown that the E6 PBMs are susceptible to phosphorylation by protein kinases on the conserved Ser/Thr residue at the antepenultimate C-terminal position, which contributes to the class 1 PBM consensus. This phosphosite is preceded by arginine residues in most of the HPV-E6 PBM sequences with recognizable patterns of the two common basophilic kinase substrate consensus motifs of R(X/R)X(S/T) and RXRXX(S/T). Additional Arg residues, e.g. at position -6 are most probably less involved in kinase recognition and more important for binding to other partners. For instance, Arg -6 of 18E6 is important for PDZ recognition but its mutation does not affect its phosphorylation efficiency by PKA, while mutation of Arg -5 and -4 blocks 18E6 PBM phosphorylation. Considering the basic consensus motifs, these 31 E6 PBMs can be readily divided into three subgroups: Subgroups 1 and 2 having sequences prone to phosphorylation by the basophilic kinases and the third, orphan subgroup with a less predictable phosphorylation propensity (Fig. 1C). Accordingly, the PBM of HPV66 E6, which in our classification falls into the orphan group, is normally calcitrant to phosphorylation by PKA in cellulo but becomes prone to it upon introduction of Arg residues at its positions -5 and -4. In addition, more than half of the identified HPV-E6 PBM sequences harbor candidate phosphorylation sites of DNA damage response ATM/ATR kinases with a consensus of (S/T)Q.

Therefore, the regulation of subgroup 1 and 2 E6 PBMs by phosphorylation is likely a rather general phenomenon. In line with earlier observations, the phospho-PBM sequences from subgroups 1 and 2 can ideally match the C-terminal 14-3-3-binding motif III (Fig. 1B and C). It is therefore likely that phosphorylation of the C-terminal PBMs from HPV-E6 types belonging to subgroups 1 and 2 by different kinases triggers their recognition by 14-3-3 proteins.
Fig. 1. Features of the structure and the C-terminal PDZ-binding motif of E6 proteins. A. The domain organization (top) and three dimensional structure (bottom) of HPV16 E6 (16E6) showing the location of the phosphorylatable PDZ-binding motif (PBM). N-terminal (E6N) and C-terminal (E6C) domains are color coded, structural Zn atoms are in black, PBM is in cyan, phosphorylatable Thr -2 is in red. B. E6 PBMs from high-risk mucosal HPV types overlap with the 14-3-3-binding motif III. The positions are numbered above, according to conventional PBM numbering. C. Classification of 31 PBM-containing HPV-E6 proteins based on the correspondence of their C-terminal PBMs to the consensus motifs phosphorylatable by basophilic kinases (subgroups 1 and 2). The third, orphan subgroup comprises PBMs of HPV-E6 proteins whose phosphorylation is less certain. Bold font indicates the E6 PBMs used in this work. Note that many PBMs overlap with recognition motifs (S/TQ) for phosphorylation by DNA damage response kinases ATM/ATR (TQ and SQ sites highlighted). Below are shown Weblogo diagrams for the PBMs within each of the three subgroups (positions are numbered according to the PBM convention).
E6 PBMs from four distinct hrm-HPV types display wide-ranging affinities to the seven 14-3-3 proteins with parallel binding profiles

Four phosopho-PBMs from E6 proteins of HPV types 16, 18, 33 and 35 belonging to subgroups 1 and 2 (as defined in Fig. 1C) were analyzed for their interaction with the all seven human 14-3-3 homodimers. For comparison, we also measured two non-viral phospho-PBMs originating from protein kinase RSK1. We used a competitive fluorescence polarization assay that measures the displacement of a fluorescent tracer 14-3-3-binding peptide in the presence of an increasing amount of a competitor molecule (all binding curves are shown in Supplementary Fig. 1A).

All phospho-PBMs (hereinafter called p16E6, p18E6, p33E6, p35E6, RSK1_-1P, and RSK1_-2P) detectably bound to 14-3-3 proteins, in sharp contrast to their unphosphorylated counterparts. The interactions between E6 phospho-PBMs and 14-3-3 proteins spanned very wide affinity ranges, from just below 1 μM (p33E6-14-3-3y) to above 300 μM (Fig. 2A and Supplementary Fig. 1). Such large binding affinity differences are noteworthy since the four E6 PBM sequences are very similar (Fig. 1B), and all 14-3-3 isoforms share highly conserved phosphopeptide-binding grooves.

Remarkably, the six phospho-PBMs showed consistent hierarchy in their relative binding preferences towards any of the seven 14-3-3 isoforms, albeit with a systematic overall shift in affinity from one peptide to another. The seven 14-3-3 isoforms clustered as four groups of decreasing affinity, in a conserved order from the strongest to the weakest phospho-PBM binder: gamma, eta, zeta/tau/beta and epsilon/sigma (γ, η, ζ/τ/β, and ε/σ) (Fig. 2B). These conserved relative affinity shifts can be quantified by calculating, for two distinct 14-3-3 isoforms, their differences of free energy of binding (ΔΔG) towards each individual phosphopeptide, then calculating the average difference (ΔΔG_{av}) with its standard deviation (Fig. 2C). Between the strongest and the weakest binders (isoforms γ and σ, respectively) the average phosphopeptide-binding energy difference is ΔΔG_{av} = -5.1 ± 1.3 kJ/mol, roughly corresponding to a 11-fold K_{D} ratio.

Noteworthy enough, another study also measured the binding of seven human 14-3-3 isoforms to the completely unrelated cystic fibrosis transmembrane conductance regulator (CFTR) phosphopeptide CFTR-R7, representing an internal 14-3-3-binding motif. A roughly comparable hierarchy of affinities was observed, also showing a 11-fold K_{D} ratio between the strongest binder (gamma isoform, K_{D} = 1 μM) and the weakest binder (epsilon isoform, K_{D} = 11 μM); both values being remarkably close to those obtained herein for the strongest-binding E6 phospho-PBM, p33E6. Similar trends were also detected upon interaction of the Leucine-Rich Repeat Kinase 2 (LRRK2) internal motif phosphopeptides with 14-3-3 isoforms, when gamma and eta were the strongest and epsilon and sigma were the weakest binders. The K_{D} ratio for the gamma and sigma isoforms averaged for the three of LRRK2 phosphopeptides (~20) is remarkably similar to the 11-fold difference described here for the E6 PBM interaction with the strongest and weakest human 14-3-3 isoforms. Therefore, our data support the idea that specificity of 14-3-3 isoforms for protein partners follows a general trend.

The seven 14-3-3 isoforms also showed consistent profiles in their relative binding preferences towards any of the four E6 phospho-PBMs. For each 14-3-3 isoform, the four phospho-PBMs systematically rank the same way from the strongest to the weakest binder: p33E6, p18E6, p16E6 and p35E6 (Fig. 2C). The average 14-3-3 binding free energy difference for two distinct phospho-PBMs systematically rank the same way from the strongest to the weakest isoform, the four E6 PBM sequences are very similar (Fig. 1B), and all 14-3-3 isoforms share highly conserved phosphopeptide-binding grooves.

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Energy difference between p33E6 and p35E6 was $\Delta \Delta G_{\text{av}} = -10.9 \pm 0.7$ kJ/mol, roughly corresponding to a 100-fold $K_D$ ratio.

| K_D (μM) ± std | 14-3-3γ | 14-3-3η | 14-3-3ζ | 14-3-3τ | 14-3-3β | 14-3-3ε | 14-3-3σ |
|----------------|--------|--------|--------|--------|--------|--------|--------|
| pHV33E6        | 0.94 ± 0.25 | 1.98 ± 0.15 | 2.72 ± 0.39 | 3.52 ± 0.28 | 4.05 ± 0.65 | 10.3 ± 2.39 | 6.77 ± 1.13 |
| pHV18E6        | 11.1 ± 0.6 | 23.6 ± 6.2 | 22.4 ± 1.3 | 37.8 ± 6.0 | 42.2 ± 6.3 | 101 ± 20 | 139 ± 24 |
| pHV16E6        | 37.2 ± 3.0 | 80.5 ± 16.9 | 73.6 ± 19.1 | 144 ± 28 | 159 ± 45 | >300 | >300 |
| pHV35E6        | 125 ± 6 | 163 ± 19 | 191 ± 18 | 233 ± 14 | >300 | >300 |
| pHV35E6_{1,4R} | 71.2 ± 6.7 | 85.0 ± 20.6 | 139 ± 21 | 194 ± 37 | 179 ± 29 | 270 ± 20 | >300 |
| pHV35E6_{1,4R, E-1A} | 9.31 ± 0.72 | 12.4 ± 1.5 | 17.6 ± 3.6 | 21.5 ± 1.2 | 20.2 ± 1.4 | 31.9 ± 4.2 | 44.1 ± 13.3 |
| RSK1_{-1P}    | 0.31 ± 0.03 | 0.46 ± 0.06 | 0.96 ± 0.10 | 0.78 ± 0.13 | 0.74 ± 0.04 | 1.58 ± 0.21 | 1.71 ± 0.23 |
| RSK1_{-2P}    | 0.20 ± 0.01 | 0.66 ± 0.05 | 1.54 ± 0.09 | 0.45 ± 0.08 | 0.96 ± 0.03 | 1.22 ± 0.16 | 3.57 ± 0.32 |

Fig. 2. Summary of affinity measurements carried out in this study. A. Affinities of four selected HPV-E6 phospho-PBMs, p35E6 mutants and RSK1 phosphopeptides towards the seven human 14-3-3 isoforms as determined by fluorescence polarization using FITC-labeled HSBP6 phosphopeptide as a tracer. Apparent $K_D$ values determined from competitive FP experiments are presented. B. The heatmap representation of the data on panel A showing the affinity trends in the interaction profiles between 14-3-3 isoforms and four HPV-E6 phospho-PBMs from weakest (white) to strongest (red). C. Averaged $\Delta \Delta G$ values between 14-3-3 isoforms or E6 phospho-PBM pairs, calculated based on their observed order of binding affinities (from weakest to strongest). Individual $K_D$ values from Supplementary Fig. 1 were first converted into $\Delta G$ values (at T=295 K; excluding cases when $K_D > 300$ μM) and average $\Delta \Delta G$ values ($\Delta \Delta G_{\text{av}}$) were calculated between the indicated motifs/isoforms.

**Atomic structure reveals the 14-3-3ζ / 18E6 PBM interface**

To get structural insight into the 14-3-3ζ interactions with 18E6 PBM, we designed, as previously reported $^{44, 45}$, a protein-peptide chimera, which we co-expressed with PKA and purified from *E. coli* cells by using an engineered cleavable His$_6$ tag (Supplementary Fig. 2A-C). The 14-3-3ζ core was modified to block phosphorylation of the semiconserved Ser58 located in the dimer interface $^{46}$ and to facilitate crystallization $^{47}$ and was tethered to
the 18E6 phosphorylatable peptide RRRETQV-COOH. This approach was justified because of the disordered nature of the E6 PBMs\textsuperscript{48,49}.

The 14-3-3\textgreek{z}-18E6 PBM chimera was stoichiometrically phosphorylated by co-expressed PKA, which gave a distinct downward shift on native PAGE (Supplementary Fig. 2D), and readily formed diffraction-quality crystals (all statistics is presented in Table 1). Structure determination was carried out at a 1.9 Å resolution with a single 14-3-3 dimer in the asymmetric unit (Supplementary Fig. 3A). The 18E6 phosphopeptide of this dimer was bound \textit{in-trans} in the amphipathic 14-3-3 grooves of the crystallographic symmetry neighbor molecule, resulting in swapped phosphopeptides (Supplementary Fig. 3A). Such arrangement, previously observed for 14-3-3\textgreek{sigma} chimeras with other phosphopeptides\textsuperscript{44,50} provided us with high-resolution insight into the 14-3-3/18E6 PBM interaction (Fig. 3A-C). All residues of the 18E6 phosphopeptides could be traced including the conformation of the C-terminal carboxyl-group. The phosphopeptide is stabilized by multiple polar interactions involving those between phospho-Thr\textsubscript{156} (position -2) and conserved residues of the basic pocket in the amphipathic groove of 14-3-3 (Arg56, Arg129, Lys49 and Tyr128, 14-3-3\textgreek{z} numbering), the phosphopeptide backbone interactions to Asn224 and Asn173, and a remarkable polar contact between the carboxyl-group of the peptide and the side chains of Lys120 and one of the two alternative conformations of Ser45 of 14-3-3\textgreek{z} (Fig. 3C). The carboxyl-group orientation is further stabilized by \textit{in-cis} H-bond with the side chain of the preceding residue Gln157 (position -1) (Fig. 3C). The side chain of the C-terminal Val\textsubscript{158} (position 0) of the peptide makes a hydrophobic contact with Val46 of 14-3-3\textgreek{z}. A second remarkable hydrophobic contact is observed between the side chain of Val176 of 14-3-3\textgreek{z} and the methyl-group of phospho-Thr\textsubscript{156} (position -2), which hints at a more stable binding of pThr than would be in the case of pSer.

Importantly, the conformation of 18E6 phosphopeptide bound to 14-3-3\textgreek{z} within the chimera is practically identical (RMSD = 0.17 Å upon superimposition of the six core Ca atoms of the peptides) to the 14-3-3\textgreek{sigma} bound conformation of a synthetic 16E6 phosphopeptide reported very recently at a much lower resolution (Fig. 3D)\textsuperscript{24}. In both cases, the carboxyl-group gives a polar contact to the Lys120 side chain and the hydrophobic side chain of the last peptide residue (Val158 in 18E6 and Leu151 in 16E6) faces the side chain of Val46 of 14-3-3. This validates the chimeric approach\textsuperscript{44} as a strategy to obtain high-resolution crystal structures of 14-3-3/phosphopeptide complexes. Arg residues that are likely to be involved in kinase recognition also contribute to 14-3-3 binding. The Arg residue occupying position -4 forms an \textit{in-cis} interaction with the phosphoryl group, and the Arg residue occupying position -5 forms in both 16E6 and 18E6 a \textpi stacking interaction with Arg60 of 14-3-3 (Fig. 3D). Most of interface contacts of the 18E6/14-3-3\textgreek{z} and 16E6/14-3-3\textgreek{sigma} complexes are similar, suggesting that the two current crystal structures can also serve as templates to build accurate homology models of other 14-3-3/E6 complexes and, more generally, any motif III complexes with a pS/pTXX-COOH consensus.

However, some noteworthy differences appear in a subset of the crystallographic conformers of 14-3-3/16E6 and 14-3-3/18E6 complexes. On the one hand, in 1 of the 4 conformers observed in the asymmetric unit of the 14-3-3\textgreek{sigma}/16E6 crystal, the side chains of Arg -7 (Gln in 18E6) and Glu -3 form an additional \textit{in-cis} salt bridge (Fig. 3D). On the other hand, Arg -6 of 18E6 (Thr in 16E6) mediates a bipartite interaction with 14-3-3 in most of the observed conformers. It simultaneously interacts with the carbonyl of Asp223 and participates in a water-mediated interaction with Asn224 (Fig. 3C and D).
Fig. 3. Molecular interface between 14-3-3ζ and phosphorylated 18E6 PBM at a 1.9Å resolution. A. An overall view on the 14-3-3ζ dimer (subunits are in tints of grey) with two bound 18E6 phosphopeptides (cyan sticks). B. A magnified view on one of the amphipathic grooves of 14-3-3ζ showing the conformation of the 18E6 phosphopeptide and the corresponding 2F₀-Fₐ electron density maps contoured at 1σ. Positions are numbered according to the PBM convention. C. Polar contacts (dashed lines) and hydrophobic interactions (semitransparent spheres) stabilizing the bound 18E6 peptide conformation. D. An overlay of the two 14-3-3 bound phosphopeptides from 16E6 (6TWZ.pdb) and 18E6 (this work) showing the similarity of the conformation. # denotes the C-terminus (-COOH). w – the water molecule, π – π-stacking interaction. Important positions are numbered according to the PBM convention. E. Averaged ΔΔG values between 14-3-3 isoforms or 35E6 phospho-PBM pairs, calculated based on their observed order of binding affinities (from weakest to strongest). Individual Kₒ values from Supplementary Fig. 1 were first converted into ΔG values (at T=295 K; excluding cases when Kₒ > 300 μM) and average ΔΔG values (ΔΔGav) were calculated between the indicated motifs/isoforms.
Rescue of the weakest E6-14-3-3 interaction by rational design

We hypothesized that the remarkable affinity differences of the four studied E6 phospho-PBMs ($\Delta G_{av} = -10.9 \pm 0.7$ kJ/mol between the weakest and the strongest 14-3-3-binder) might result from two opposed mechanisms: interface peptide-14-3-3 contacts favoring complex formation versus intra-peptide contacts within the unbound peptide disfavoring it. On the one hand, position -6 is an Arg in the two strongest 14-3-3-binders (18E6 and 33E6) versus a Thr in the weakest ones (35E6 and 16E6). Accordingly, the crystal structures have shown that Arg -6 can mediate more interactions with the generic 14-3-3 interface (Fig. 3D). On the other hand, all E6 phospho-PBMs have a delicate charge distribution, with an acidic C-terminus (that is also involved in PDZ-domain binding) and a basic N-terminal segment (that is also involved in kinase recognition). These local charged segments may form transient in-cis interactions within the unbound phosphopeptide, so-called "charge clamps" $^{51}$, potentially unfavorable for 14-3-3 binding. We speculated that Glu -1 in p35E6, the weakest 14-3-3 binder, might participate in such a charge clamp, thereby disfavoring the binding.

To test these mechanisms, we synthesized three variants of p35E6. The first variant contained a T-6R substitution, which in principle could allow a more stable bound conformation, but may also stabilize charge-clamps in the free form of the motif. The second variant contained an E-1A substitution, which in principle could destabilize in-cis charge-clamps. A third variant contained both substitutions. All substitutions turned out to reinforce the binding affinities of 35E6 without altering the apparent preferences of the different 14-3-3 isoforms (Fig. 2A and Fig. 3E). Taken individually, T-6R moderately increased binding ($\Delta G_{av} = -1.1 \pm 0.5$ kJ/mol, 1.5-fold $K_D$ ratio), while E-1A strongly reinforced it ($\Delta G_{av} = -5.1 \pm 0.2$ kJ/mol, 11-fold $K_D$ ratio). When combined, the two substitutions synergistically increased binding ($\Delta G_{av} = -8.7 \pm 0.4$ kJ/mol, 35-fold $K_D$ ratio), thereby turning p35E6 from the weakest 14-3-3 binder into the second strongest one, just below p33E6.

Fusicoccin partially destabilizes the 14-3-3/E6 PBM interaction

Fusicoccin (FSC) is commonly used as a stabilizer of 14-3-3 complexes, when its binding in the distinct pocket in the 14-3-3/phosphopeptide interface is allowed by phosphopeptide side chains of the amino acids in downstream positions relative to the phospho-residue $^{42, 52, 53, 54}$. Binding of FSC is known to stabilize the 14-3-3 interaction with few internal motif I phosphopeptides (e.g., Gab2 peptide, 5EXA.pdb $^{55}$ and CFTR peptide, 5D3F.pdb $^{42}$) and numerous peptides with the short motif III consensus, pS/pTX-COOH $^{31, 56, 57}$. Most of the internal motif phosphopeptides, when bound to 14-3-3, apparently leave insufficient space for FSC and therefore hinder formation of the ternary complexes with FSC. Many motif III peptides with a consensus pS/pTX-COOH enable synergistic binding of the phosphopeptide and FSC, reinforcing the 14-3-3/phosphopeptide complex $^{31, 52, 53, 57, 58, 59}$. At the same time, very little is known about the effect of FSC on binding of the longer motif III peptides to 14-3-3, such as in the case of E6 PBMs.

We investigated whether binding of phosphorylated E6 PBM to 14-3-3 might be affected by this small molecule and analyzed its effect on the affinity of the E6 PBMs to 14-3-3. We used FP experiments to measure equilibrium binding affinity constants of complexes.
between the four HPV-E6 phosphopeptides and 14-3-3 isoforms ζ and γ, in the presence of 100 µM FSC (Supplementary Fig. 1B and Fig. 4A). The addition of FSC consistently decreased by 1.5 to 2 fold the affinities of all interactions ($\Delta \Delta G_{av} = -1.3 \pm 0.5$ and $-1.8 \pm 0.4$ kJ/mol for ζ and γ, respectively) without altering the apparent preferences of the different peptides (Fig. 4A and B). Therefore, fusicoccin moderately destabilized all studied 14-3-3/E6 interactions, in contrast to its much broadly reported, stabilizing effect on 14-3-3 complexes 31, 42, 52. Nevertheless, the binding of E6 peptides and FSC was not mutually exclusive.

|                      | 14-3-3ζ | 14-3-3ζ + FSC | 14-3-3γ | 14-3-3γ + FSC |
|----------------------|---------|---------------|---------|---------------|
| pHPV33E6             | 2.72 ± 0.39 | 3.94 ± 0.40   | 0.94 ± 0.25 | 2.15 ± 0.26   |
| pHPV18E6             | 22.4 ± 1.3   | 37.3 ± 5.2    | 11.1 ± 0.6  | 19.3 ± 2.0    |
| pHPV16E6             | 73.6 ± 19.1  | 161 ± 23      | 37.2 ± 3.0  | 91.7 ± 13.2   |
| pHPV35E6             | 191 ± 18     | 294 ± 15      | 125 ± 6     | 228 ± 61      |

$\Delta \Delta G_{av} = -1.28 \pm 0.45$ (kJ/mol) for 14-3-3ζ + FSC < 14-3-3ζ

$\Delta \Delta G_{av} = -1.77 \pm 0.42$ (kJ/mol) for 14-3-3γ + FSC < 14-3-3γ

**Fig. 4.** The destabilizing effect of FSC on the 14-3-3ζ/18E6 PBM interaction. A. Affinities of four selected HPV-E6 phospho-PBMs towards human 14-3-3ζ and 14-3-3γ in the absence and in the presence of FSC as determined by fluorescence polarization using FITC-labeled HSPB6 phosphopeptide as a tracer. Apparent $K_D$ values determined from competitive FP experiments are presented. The binding curves are shown in Supplementary Fig. 1. B. Averaged $\Delta \Delta G$ values between 14-3-3-E6 phospho-PBM pairs in the absence or in the presence of FSC, calculated based on their observed order of binding affinities (from
weakest to strongest). Individual $K_D$ values from Supplementary Fig. 1 were first converted into $\Delta G$ values (at $T=295$ K; excluding cases when $K_D > 300$ μM) and average $\Delta \Delta G$ values ($\Delta \Delta G_{\text{av}}$) were calculated between the indicated motifs/isofoms. C. An overall view on the ternary complex between 14-3-3ζ (subunits are shown by surface using two tints of grey), 18E6 phosphopeptide (cyan sticks) and FSC (pink sticks). FSC was soaked into the 14-3-3ζ-18E6 chimera crystals. 2F$_{\text{o}}$-F$_{\text{c}}$ electron density maps contoured at 1σ show are shown for the phosphopeptide and FSC only. D. A closeup view showing polar contacts (dashed lines) and hydrophobic interactions (semitransparent spheres) positioning the 18E6 phosphopeptide (cyan sticks) and FSC (thin pink sticks) in the amphipathic groove of the 14-3-3ζ subunit (semitransparent light grey ribbon). 2F$_{\text{o}}$-F$_{\text{c}}$ electron density maps contoured at 1σ are shown for the peptide and FSC. # denotes the C-terminus (-COOH). The GGGG linker is omitted for clarity. E. Chemical formula of FSC showing the positions of the functional groups discussed in the text. F. The effect of FSC binding. Conformational changes upon FSC binding are shown by red arrows reflecting the closure of the C-terminal α-helix of 14-3-3 and the rotation and translocation of the C-terminal residues in the 18E6 peptide, accompanied by a significant rise of the local B-factors (are shown as a gradient from blue to red as indicated). The amplitudes of the conformational changes in 14-3-3 and 18E6 peptide are indicated in Å by dashed arrows.

**Structural basis for the destabilizing effect of fusicoccin**

To get structural insights into the peculiar destabilizing effect of FSC on the 14-3-3/E6 PBM interaction, we soaked the 14-3-3ζ/18E6 PBM chimera crystals with FSC and solved the crystal structure of the 14-3-3ζ/18E6 PBM/FSC ternary complex (Fig. 4C-F and Table 1). FSC soaking did not disrupt the crystal lattice of the 14-3-3ζ/18E6 PBM chimera crystals (Supplementary Fig. 3) and the overall assembly (Fig. 4C and D); however, significant local structural rearrangements could be observed. FSC (Fig. 4E) binding results in a ~4 Å closure of the last α-helix of 14-3-3ζ (Fig. 4F), similar to what has already been described for other 14-3-3 complexes containing FSC. Crystal lattice obtained by FSC soaking preserved the characteristic phosphopeptide swap stabilizing the contacts with the neighboring chimera molecules, and FSC was bound in each 14-3-3ζ subunit, side-to-side with the C-terminal end of the 18E6 PBM (Fig. 4C and D, Supplementary Fig. 3). All residues and atoms within the analyzed area could be observed in the electron density thanks to the high resolution (Table 1).

FSC occupies its well-defined cavity where it is positioned by hydrophobic interactions with Phe117, Ile166, Ile217 and Leu216, polar contacts with residues Asn42 and Asp213, and a remarkable H-bond involving its 3-methoxy oxygen and the side chain of Lys120 of 14-3-3ζ (Fig. 4D). The latter contact breaks the Lys120 interaction with the carboxyl-group of the 18E6 PBM formed in the absence of FSC, displacing the carboxyl to another position, where it establishes a new contact with the 12-hydroxy group of FSC (Fig. 4D and E). 14-3-3ζ Lys49 also switches its position, and loses a contact to the backbone carbonyl of pThr156 (Fig. 3C) to establish instead a contact with the 12-hydroxy group of FSC. The side chain of the C-terminal Val158 of the 18E6 PBM shifts 3.5 Å towards the phosphate moiety of Thr156, breaking the hydrophobic contact with Val46 of 14-3-3ζ and significantly dispersing the local electron density (Fig. 4F). As a result, while most of the peptide conformation remained unchanged, the B-factors of the last 18E6 PBM residue in the refined FSC-bound structure increased significantly (Fig. 4F). Therefore, FSC binding...
causes an apparent strain in the conformation of the C-terminus of 18E6 PBM (Figs 4 and 5). In addition, the presence of FSC significantly reshuffles the water network that surrounds the E6 carboxy-terminal extremity, which directly contacts the bound FSC (Fig. 5). The simultaneous binding of the 18E6 PBM and FSC in the amphipathic groove of 14-3-3 and the mild destabilizing influence of FSC indicate that this ternary complex can be used as a starting point to design both stabilizers and inhibitors of 14-3-3/E6 interactions.

Fig. 5. Comparison of the water-mediated polar contacts formed in the 14-3-3ζ/18E6 interface in the absence (A) or in the presence of fusicoccin (FSC) (B). The main residues involved in the interactions are shown by sticks with color coding: 14-3-3 residues are shown in light grey, 18E6 residues are in cyan, phospho-group of Thr156 is shown by orange sticks. FSC is shown by thin magenta sticks, water molecules affected by FSC binding are shown by lime green, those similar in two structures are black. The C-terminal 18E6 residue (V158) is denoted by #. Note the significant redistribution of water molecules upon FSC binding.

DISCUSSION

E6 oncoproteins of all high-risk alpha HPV types contain a conserved C-terminal PDZ-Binding Motif which can turn into a potential 14-3-3-binding motif upon phosphorylation of
the conserved Thr/Ser residue at position -2 (Fig. 1). We demonstrated here that the phosphorylated forms of four selected alpha HPV-E6 PBM s detectably bound to all seven human 14-3-3 isoforms. Unexpectedly, the binding affinities of these 28 distinct 14-3-3/E6 complexes showed large variations, spanning a 10-fold $K_D$ range for different 14-3-3 isoforms binding to a given PBM, and a 100-fold $K_D$ range for different E6 phospho-PBMs binding to a given 14-3-3 isoform. Furthermore, affinity variations followed trends that were all but random. The 14-3-3 binding profiles of all E6 phosphopeptides turned out to be remarkably parallel, and so were the E6-binding profiles of all 14-3-3 isoforms (Fig. 2B and C).

When considering the present data together with previously published literature, 14-3-3 interactions with phosphorylated motifs from different origins have a strikingly wide affinity range, spanning from low nanomolar to low millimolar detectable dissociation constants (Fig. 6). Despite variable interaction modes (e.g. monovalent vs. divalent), the 14-3-3 family-wide affinity trends are surprisingly parallel, with 14-3-3$\gamma$ and 14-3-3$\eta$ consistently being the strongest binders and 14-3-3$\sigma$ and 14-3-3$\epsilon$ being the weakest binders. Independently of the nature of the target motif, the average maximal $K_D$ ratio between the strongest-binding and the weakest-binding 14-3-3 is around 10-fold in our present work and also in the literature (Fig. 6). Conversely, phosphopeptides, even very similar in sequence, can sample much wider ranges of binding affinities for a given 14-3-3 isoform. For instance, for 14-3-3$\gamma$, the $K_D$ ratio between the strongest and the weakest binding phosphopeptide is almost 625-fold in the present work, and 39,000-fold when taking into account other reports $^{42, 43, 60, 61}$ (Fig. 6). However, the discussed weak interactions do not necessarily lead to the absence of complex formation in cellulo, as 14-3-3 proteins are among the most abundant proteins in human cells $^{62, 63}$. Indeed, according to Protein Abundance Database $^{63}$, 14-3-3$\epsilon$ is the 48th most abundant human protein (2479 ppm) and 14-3-3$\zeta$ is the 72nd (1680 ppm), whereas all seven 14-3-3 isoforms are within the 820 most abundant proteins (i.e., the top 4.1%) out of 19949 proteins in the integrated whole human body dataset. Given such high abundance, strong 14-3-3 interactions might lead to extremely tight complex formation, while weak interactions might lead to transient, more dynamic 14-3-3 complexes.

The observed strong and parallel affinity variations are noteworthy if one considers the high degree of conservation of the amphipathic grooves of the seven 14-3-3 isoforms and of the amphipathic groove-binding residues of the four E6 PBM s. Indeed, affinity differences may stem not only from structural particularities of the bound complexes but also from intrinsic properties of the unbound partners. Conformations unfavorable for complex formation may exist in various proportions for each 14-3-3 isoform and each E6 phosphopeptide, thereby influencing at various degrees the formation of the complexes and their resulting binding affinities. On the one hand, full-length 14-3-3 proteins feature the flexible C-terminal tails that are the most variable elements among isoforms; at least for some isoforms they have been shown to sample conformations also occupying the amphipathic 14-3-3 grooves and thereby regulating their affinities for target peptides $^{64, 65}$. Conformational dynamics of the 14-3-3 dimers along the open-closed state trajectory $^{26, 44}$ may also differ among 14-3-3 isoforms, which can probably affect the effectiveness of adopting the peptide-bound conformation. On the other hand, it has been shown that a phosphoryl-group in a disordered segment can form local "charge clamps" in-cis with a neighboring Arg/Lys residue, thereby decreasing its apparent availability for binding partners $^{51}$. The four E6 PBM s studied herein display distinctive charged, polar or non-polar residues at several
positions (-1; -6; -7; -8; -9) that may influence their exploratory conformational spaces. By replacing the Glu -1 by Ala in the weakest 14-3-3 binder, p35E6, we obtained a significantly stronger 14-3-3 binder, suggesting that we successfully disrupted a charge clamp that is unfavorable to binding of the p35E6 motif. Nonetheless, fine differences of contact networks at the interfaces of E6/14-3-3 complexes may also influence affinity. In this line, comparison of the present p18E6/14-3-3ζ structure to the previously solved p16E6/14-3-3γ structure \(^{24}\) suggested that an Arg residue at position -6 established favorable interface contacts (Fig. 3C and D). Accordingly, replacing Thr -6 of p35E6 with Arg -6 as found in p18E6 and p33E6, effectively reinforced the binding event. Finally, combining both Glu→Ala and Thr→Arg substitutions at positions -1 and -6, respectively, resulted in a p35E6 variant that bound 14-3-3 proteins almost as strongly as the strongest studied E6 binder, p33E6 (Fig. 2A and Fig. 3E). Interestingly, the RSK1 phospho-PBMs (RRVRKLPSTpT-L-COOH and RRVRKLPStTTL-COOH), featuring non-charged residues Thr and Leu in the downstream position relative to the phosphothreonine, and therefore being unable to form the equivalent charge-clamp that is suggested in the case of p35E6 (SKPTTRRepTEV-COOH), show remarkably high affinities to all 14-3-3 isoforms \(\text{per se}\) (Fig. 2A and Supplementary Fig. 1). Our data prove that charge clamps can indeed strongly contribute to the binding preferences of different phosphopeptide motifs. This principle needs to be further examined for all kinds of phosphorylation-regulated interactions as charge-clamp formation might be an often ignored biochemical property of practically all phosphorylation sites.

**Fig. 6.** Compilation of 14-3-3 family-wide affinity measurements of exemplary 14-3-3-binding motifs. A. Affinity maps of 14-3-3 interactions based on experimentally determined dissociation constants against the 14-3-3ome, as obtained in the current work and in refs \(^{42, 43, 60, 61}\). B. When normalized to the strongest partner-binder 14-3-3γ, all studied peptides are following very similar affinity trends between the different 14-3-3 isoforms. Between the strongest and the weakest partner-binding 14-3-3 isoform, a 12-fold dissociation constant decrease was found on average. C. When normalizing to the generally strongest studied 14-3-3-binding motif, the isoform-specific affinity variations for each peptide are compensated to clearly show the average differences between different binding motifs. Between the strongest and the weakest analyzed 14-3-3-binding motif, a 34,000-fold \(K_0\) decrease was found on average for all 14-3-3 isoforms. Only counting the very similar HPV-E6 peptides sequence-wise, this affinity difference can still be larger than 100-fold. Therefore, the motif-to-motif affinity differences are expected to be much higher than the 14-3-3 isoform-to-isoform affinity differences. Binding motifs that are analyzed in other studies are highlighted with a grey background \(^{42, 43, 60, 61}\). The color scale is either based on affinity values or in \(K_0\) ratios. √ denotes affinities weaker than the limit of quantitation of the fluorescence polarization assay.
When phosphorylated, all the E6 PBMs studied here ideally match motif III consensus recognized by 14-3-3 with two C-terminal residues downstream of the phosphorylated threonine (pS/pTXX-COOH) (Fig. 1B). Structures of motif III complexes with 14-3-3 have been less documented than the internal motif I/II complexes. Moreover, the pS/pTXX-COOH subtype of motif III sites, studied here, is underrepresented in the Protein Data Bank, where mostly motif III sites with only one residue after a phospho-residue have been co-crystallized with 14-3-3 (Supplementary Table 1). To the best of our knowledge, the only two reported structures with pS/pTXX-COOH consensus peptides are those of 14-3-3γ with the influenza virus protein NS1 (4O46.pdb) and our recently reported 14-3-3σ complex with the synthetic papillomavirus 16E6 phosphopeptide (6TWZ.pdb 24), however, both of them have rather low resolution (2.8-2.9 Å) (Supplementary Table 1). Together with the henipavirus protein W binding to 14-3-3σ via its C-terminal motif III peptide RRMPSN-COOH 66, the adeno-associated virus protein Rep68 binding to 14-3-3 proteins using its C-terminal motif III peptide RGHpSL-COOH 67 and the hepatitis B virus protein X binding to 14-3-3ζ via an internal motif I peptide RPLpSGP 68, these examples illustrate that viral proteins commonly use their elements, efficiently mimicking the host 14-3-3-binding peptides, to hijack the cellular functions controlled by 14-3-3 proteins. One of the possible mechanisms is the 14-3-3-mediated stabilization of viral proteins to evade dephosphorylation and degradation, which may prolong the half-life and increase chances for successful replication and further infections 21, 68.

The phosho-PBM conformation revealed by our crystal structures, ideally congruous to the amphipathic groove of 14-3-3, leaves vacant the cavity that is known to be druggable and occupied by a fungal toxin called fusicoccin (FSC) 33 54 56. Benefitting from the well-diffracting crystals of the 14-3-3ζ/18E6 PBM chimera, we proceeded with their soaking with FSC, which resulted in a 1.85 Å crystal structure of the ternary 14-3-3ζ/18E6 PBM/FSC complex. The direct comparison of the 14-3-3ζ/18E6 complex in the absence and in the presence of FSC revealed a significant conformational change in the last α-helix of 14-3-3ζ, a rearrangement of bound water molecules and a remarkable strain in the C-terminal part of the 18E6 PBM peptide (Figs 4 and 5). This strain forced the translocation of the carboxyl-group and increased the displacement factors for the last C-terminal residue, suggesting destabilization of the 14-3-3ζ/18E6 PBM complex by FSC. This partial destabilization, and thus the less-documented inhibitory action of FSC, was confirmed by in vitro binding assays using fluorescence polarization (Fig. 4A and B). Noteworthy, the structural data on the effect of FSC on the C-terminal end of the E6 PBM is very consistent with the preservation of the affinity differences for various 14-3-3/phospho-PBMs complexes in the presence of FSC (Fig. 4A). This supports the notion that those affinity differences are dictated by positions beyond the last C-terminal residue, including those involved in formation of charge clamps.

We could find only one earlier reported example when FSC decreased the binding affinity of a 14-3-3 to a motif III peptide (the interleukin 9 receptor alpha chain (IL-R9α) peptide, RSWpTF-COOH 31), but in that case no experimental structural information was available. In addition, it was reported that binding of the shortest motif III peptide from the cyclin-dependent kinase inhibitor (p27Kip1) RRQpT-COOH to 14-3-3 is not affected by FSC, most likely due to the absence of direct contacts between FSC and the phosphopeptide, which both bind in the amphipathic groove of 14-3-3 independently 31. The structural basis for the inhibitory action of FSC has very recently been reported for several internal motif I 14-3-3-
binding peptides 58. Therefore, complementing the range of reports on the stabilizing effect of FSC 31, 42, 54, 55, 56, our results provide the first structural evidence that FSC can be a negative regulator of 14-3-3 interactions with typical motif III peptides.

Thus, our structural and in vitro binding data with FSC confirm the druggability of the 14-3-3-E6 interaction and suggest that appropriate modification and optimization of the small molecule may provide promising opportunities for selective modulation of viral protein-14-3-3 interactions in the future.

METHODS

Cloning, protein expression and purification and peptide synthesis

Previously described chimeras contained the C-terminally truncated human 14-3-3ζ (Uniprot ID P31947; residues 1-231, 14-3-3ζΔC) bearing on its N-terminus a His6-tag cleavable by 3C protease and phosphorylatable peptides tethered to the 14-3-3ζ C-terminus by a GSGS linker 44. The novel chimera was designed taking into account the following modifications. First, it contained the C-terminally truncated human 14-3-3ζ sequence (Uniprot ID P63104; residues 1-229, 14-3-3ζΔC) connected to the PKA-phosphorylatable 18E6 heptapeptide around Thr156. Second, the 14-3-3ζ core was modified to block Ser58 phosphorylation (S58A) 69, 70. Third, to improve crystallizability, the 14-3-3ζ sequence was mutated by introducing the 73EKK75→AAA and 157KKE159→AAA amino acid replacements in the highest-scoring clusters 1 and 2 predicted by the surface entropy reduction approach 47, 71. Finally, the linker was changed to GGGS to exclude its unspecific phosphorylation (Supplementary Fig. 2A).

cDNA of the 14-3-3ζ-18E6 chimera was codon-optimized for expression in Escherichia coli and synthesized by IDT Technologies (Coralville, Iowa, USA). The 14-3-3ζΔC gene was flanked by NdeI and AgeI restriction endonuclease sites to enable alteration of the 14-3-3ζ or E6 PBM peptide sequences. The entire 14-3-3ζ-GGGG-18E6 PBM construct was inserted into a pET28-his-3C vector 72 using NdeI and XhoI restriction endonuclease sites. The resulting vector was amplified in DH5α cells and verified using DNA sequencing in Evrogen (Moscow, Russia, www.evrogen.ru).

The assembled vector (Kanamycin resistance) was transformed into chemically competent E. coli BL21(DE3) cells for expression either in the absence or in the presence of the His6-tagged catalytically active subunit of mouse PKA 72. Protein expression was induced by the addition of isopropyl-β-thiogalactoside (IPTG) to a final concentration of 0.5 mM and continued for 16 h at 25 °C. The overexpressed protein was purified using subtractive immobilized metal-affinity chromatography (IMAC) and gel-filtration essentially as described earlier for 14-3-3ζ chimera 44 (Supplementary Fig. 2B and C). The purified phosphorylated 14-3-3ζ-18E6 chimera revealed the characteristic downward shift on native PAGE compared to the unphosphorylated counterpart (Supplementary Fig. 2D). Given the absence of PKA phosphorylation sites in the modified 14-3-3ζ core and the linker, this strongly indicated 18E6 phosphorylation by co-expressed PKA. The chimera was fully soluble and stable at concentrations above 20 mg/ml required for crystallization. Protein concentration was determined at 280 nm on a Nanophotometer NP80 (Implen, Germany) using extinction coefficient equal to 0.93 (mg/ml)^-1 cm^-1.

For affinity measurements, full-length human 14-3-3 constructs with a rigid N-terminal MBP fusion were used. The coding sequences of the full-length 14-3-3 epsilon, gamma and zeta
were received from Prof. Lawrence Banks. cDNAs encoding other full-length 14-3-3 isoforms β, τ, η and σ were obtained as codon-optimized for *E. coli* expression synthetic genes from IDT Technologies (Coralville, Iowa, USA). All 14-3-3 isoforms were fused via a three-alanine linker to the C-terminus of a mutant MBP carrying the following amino acid substitutions: D83A, K84A, K240A, E360A, K363A and D364A, as previously described. All resulting clones were verified by sequencing. The MBP-fused proteins were expressed in *E. coli* BL21 with IPTG induction. Proteins were affinity purified on an amylose column and were further purified by ion-exchange chromatography (HiTrap Q HP, GE Healthcare). Protein concentrations were determined by UV spectroscopy. The double-purified samples were supplemented with glycerol and TCEP before aliquoting and freezing in liquid nitrogen.

HPV peptides (35E6: biotin-ttds-SKPTRRETev; 16E6: biotin-ttds-SSRTRRETQL; 18E6: biotin-ttds-RLQRRRETQV; 33E6: biotin-ttds-SRSRRRETAL; p35E6: biotin-ttds-SKPTRREPTEV; p35E6 E-1A: biotin-ttds-SKPTRREPTEV; p35E6 T-6R: biotin-ttds-SKPRREpTEV; p35E6 E-1A T-6R: biotin-ttds-SKPRREpTEV; p16E6: biotin-ttds-SSRRREpTQL; p18E6: biotin-ttds-RLQRRREPQV; p33E6: biotin-ttds-SRSREpTAL) and RSK1 peptides (SKPRREpTEV; p16E6: biotin-ttds-RRVRKLPSTpTTL and RSK1 -1P: biotin-ttds-RRVRKLPSpTTL) were chemically synthesized in-house on an ABI 443 A synthesizer with Fmoc strategy. The fluorescently labeled HSPB6 (WLRRApSAPLPGLK) peptide (fpB6) was prepared by FITC labeling of the chemically synthesized peptide as described previously.

**Fluorescence polarization (FP) assay**

Fluorescence polarization was measured with a PHERAstar (BMG Labtech, Offenburg, Germany) microplate reader by using 485 ± 20 nm and 528 ± 20 nm band-pass filters (for excitation and emission, respectively). In direct FP measurements, a dilution series of the 14-3-3 protein was prepared in 96-well plates (96 well skirted pcr plate, 4ti-0740, 4ttitude, Wotton, UK) in a 20 mM HEPES pH 7.5 buffer containing 150 mM NaCl, 0.5 mM TCEP, 0.01% Tween 20, 50 nM fluorescently-labeled fpB6 peptide and 100 μM fusicoccin (FSC), if indicated. The volume of the dilution series was 40 μl, which was later divided into three technical replicates of 10 μl upon transferring to 384-well micro-plates (low binding microplate, 384 well, E18063G5, Greiner Bio-One, Kremsmünster, Austria). In total, polarization of the probe was measured at 8 different protein concentrations (whereas one contained no protein and corresponded to the free peptide). In competitive FP measurements, the same buffer was supplemented with the protein to achieve a complex formation of 60-80%, based on the titration. Then, this mixture was used for creating a dilution series of the unlabeled competitor (i.e. the studied peptides) and the measurement was carried out identically as in the direct experiment. Analysis of FP experiments were carried out using ProFit, an in-house developed, Python-based fitting program. The dissociation constant of the direct and competitive FP experiment was obtained by fitting the measured data with quadratic and competitive equation, respectively. ΔG values were calculated using the formula \( \Delta G = -RT \ln(K_D) \), at 295 K. \( \Delta G_{av} \) values were obtained by calculating the average and the standard deviation of all obtained individual \( \Delta G \) values (between different motifs or different proteins), excluding cases when \( K_D > 300 \mu M \).

**Crystallization and structure determination**
Crystallization conditions were screened using commercially available and in-house developed kits (Qiagen, Hampton Research, Emerald Biosystems) by the sitting-drop vapor-diffusion method in 96-well MRC 2-drop plates (SWISSCI, Neuheim, Switzerland), using a Mosquito robot (TTP Labtech, Cambridge, UK) at 4 °C. The optimized condition of the crystals consisted of 19% polyethylene glycol 4000, 0.1M cacodylate buffered at pH 5.5. For soaking, crystals were transferred to a mother-liquor solution containing (saturated, partially precipitated) 5 mM fusicoccin and crystals were harvested after an 18h incubation period. All crystals were flash-cooled in a cryoprotectant solution containing 20% glycerol and stored in liquid nitrogen.

X-ray diffraction data were collected at the Synchrotron Swiss Light Source (SLS) (Switzerland) on the X06DA (PXIII) beamline and processed with the program XDS. The crystal structure was solved by molecular replacement with a high-resolution crystal structure of 14-3-3ζ (PDB ID 2O02) using Phaser and structure refinement was carried out with PHENIX. TLS refinement was applied during the refinement. The crystallographic parameters and the statistics of data collection and refinement are shown in Table 1.

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Author contributions

G.G. purified proteins, carried out FP experiments, performed crystallographic studies, analyzed the data and edited the paper. K.V.T. cloned, purified and characterized proteins. C.K. cloned and purified proteins. P.E. synthesized the peptides. G.T. performed data analysis and data interpretation and edited the paper. N.N.S. contributed to protein purification and crystallographic experiments, supervised the research, analyzed the data and wrote the paper.

Conflict of interests

The authors declare no conflict of interest.

Data availability

The refined model and the structure factor amplitudes have been deposited in the PDB with the accession codes 6ZFD and 6ZFG. All other data supporting the findings of this study are available from the corresponding authors upon reasonable request.

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Table 1. Crystallographic statistics.

|                      | 14-3-3ζ-18E6 chimera | 14-3-3ζ-18E6 chimera + FSC |
|----------------------|-----------------------|----------------------------|
| **Data collection**  |                       |                            |
| Wavelength           | 1.00                  | 1.00                       |
| Resolution range     | 39.26 - 1.9 (1.95 - 1.9) | 38.05 - 1.85 (1.9 - 1.85) |
| Space group          | P 2 1 2 1 2 1         | P 2 1 2 1 2 1              |
| Unit cell (a, b, c, α, β, γ) | 72.35, 78.53, 90.3, 90, 90, 90 | 73.23, 76.1, 88.95, 90, 90, 90 |
| Total reflections    | 547783 (37726)        | 557315 (41375)             |
| Unique reflections   | 41285 (2986)          | 41927 (3038)               |
| Multiplicity         | 13.3 (12.6)           | 13.3 (13.6)                |
| Completeness (%)     | 100 (100)             | 97.1 (96.5)                |
| Mean I/σ(I)          | 13.19 (1.39)          | 12.36 (1.40)               |
| R-meas               | 16.6 (205)            | 16.3 (216)                 |
| CC1/2                | 99.9 (54.7)           | 99.8 (58.4)                |
| **Refinement**       |                       |                            |
| R-work               | 0.1764                | 0.1908                     |
| R-free               | 0.2071                | 0.2192                     |
| Number of non-hydrogen atoms | 4468                   | 4589                       |
| macromolecules       | 4034                  | 4004                       |
| ligands              | 24                    | 102                        |
| solvent              | 410                   | 483                        |
| Protein residues     | 481                   | 482                        |
| RMS(bonds)           | 0.006                 | 0.006                      |
| RMS(angles)          | 0.83                  | 0.71                       |
| Ramachandran favored (%) | 99.36                  | 98.51                      |
| Ramachandran allowed (%) | 0.64                  | 1.49                       |
| Ramachandran outliers (%) | 0                    | 0                          |
| Rotamer outliers (%) | 1.65                  | 2.13                       |
| Clashscore           | 6.68                  | 3.17                       |
| Average B-factor     | 33.39                 | 29.6                       |
| macromolecules       | 32.42                 | 28.49                      |
| ligands              | 62.93                 | 32.67                      |
| solvent              | 41.2                  | 38.11                      |
| Number of TLS groups | 15                    | 11                         |
| PDB ID               | 6ZFD                  | 6ZFG                       |