Understanding the interaction of 14-3-3 proteins with hDMX and hDM2: a structural and biophysical study

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p53 plays a critical role in regulating diverse biological processes: DNA repair, cell cycle arrest, apoptosis and senescence. The p53 pathway has therefore served as the focus of multiple drug-discovery efforts. p53 is negatively regulated by hDMX and hDM2; prior studies have identified 14-3-3 proteins as hDMX and hDM2 client proteins. 14-3-3 proteins are adaptor proteins that modulate localization, degradation and interactions of their targets in response to phosphorylation. Thus, 14-3-3 proteins may indirectly modulate the interaction between hDMX or hDM2 and p53 and represent potential targets for modulation of the p53 pathway. In this manuscript, we report on the biophysical and structural characterization of peptide/protein interactions that are representative of the interaction between 14-3-3 and hDMX or hDM2. The data establish that proximal phosphosites spaced ~20–25 residues apart in both hDMX and hDM2 cooperate to facilitate high-affinity 14-3-3 binding and provide structural insight that can be utilized in future stabilizer/inhibitor discovery efforts.

Introduction

The transcription factor p53 is often described as ‘the guardian of the genome’ because of its central role in regulating diverse biological processes: DNA repair, cell cycle arrest, apoptosis and senescence [1]. Thus, the p53 pathway has seen significant attention in developing new anticancer treatments [2]. One such target has been the interaction between p53 and hDM2 or hDMX (Fig. 1A) [3–5]; p53 is negatively regulated by hDMX and hDM2 [6–10]; they bind to the transactivation domain of p53 sterically blocking its interaction with DNA [9–12]; p53 is additionally targeted for nuclear export and proteasomal degradation through hDM2 which bears an E3 ligase domain [13–15]. Although hDMX lacks such a domain, the E3 ligase activity of hDM2 is enhanced through formation of hDMX/hDM2 heterodimers [16–18]. This continuous targeting of p53 for degradation keeps p53 levels low under normal conditions. It is only under stressed conditions (e.g. DNA

Abbreviations
Ahx, aminohexanoic acid; AUC, analytical ultracentrifugation; Boc, tert-butoxycarbonyl; CD, circular dichroism; DCM, dichloromethane; DIPEA, N,N-Diisopropylethylamine; DMF, dimethylformamide; DMSO, dimethyl sulfoxide; DTT, dithiothreitol; FA, fluorescence anisotropy; FAM, (5,6)-carboxyfluorescein; Fmoc, fluorenylmethoxycarbonyl protecting group; HBS, HEPES-buffered saline; HCTU, (2-(6-Chloro-1H-benzotriazole-1-yl)-1,1,3,3-tetramethylammonium hexafluorophosphate); hDM2, human double minute 2 protein; hDMX, human double minute 4 protein (hDM4); HRMS, high resolution mass spectrometry; IPTG, isopropyl β-d-1-thiogalactopyranoside; ITC, isothermal titration calorimetry; LB, lysogeny broth; p53, tumour suppressor protein 53; SPPS, Fmoc-solid-phase synthesis; SPR, surface plasmon resonance; TCEP, Tris(2-carboxyethyl)phosphine; TFA, trifluoroacetic acid.
damage, hypoxia, oncogene activation and ribosomal stress) that p53 is rapidly activated through multiple post-translational modifications; subsequent stabilization and accumulation of p53 upregulates gene expression leading to cell cycle arrest and/or apoptosis. This tightly controlled autoregulatory loop is thus of vital importance; knockout studies of the oncogenes MDMX and MDM2 (the murine homologues of hDMX and hDM2) have shown that these proteins are essential in mice studies [19]. However, hDM2 and hDMX are themselves subject to regulatory control and have additional independent roles in cell cycle regulation [20–22]. For instance, in addition to modulating p53 binding, post-translational modification of hDMX and hDM2 such as phosphorylation influences their localization and degradation [21–25].

Several kinases have been reported to phosphorylate hDMX and/or hDM2 whilst cellular studies have shown that phosphorylated hDMX and hDM2 interact with 14-3-3 [26]. First identified in 1967 [27], 14-3-3 proteins are eukaryotic adapter proteins that control diverse physiological functions through interaction with a large network of proteins, for example, they are involved in signal transduction, protein trafficking, apoptosis and cell cycle regulation [28]. Given these functions, they have been shown to play an important role in inflammatory conditions, neurodegenerative diseases, cancer and cystic fibrosis [29–33]. And have consequently received attention as potential drug-discovery targets [28,34]. The 14-3-3 family of proteins consist of seven isoforms in humans: β, γ, ε, η, ζ, σ and τ/θ, all of which share a high degree of sequence conservation [35] (see Fig. S1). Even though sequence conservation between the isoforms can indicate redundancy in protein function, this is not the case for 14-3-3 proteins as different affinities towards a target...
protein are clear between the isoforms [36,37]. The 14-3-3 proteins exist as homo or heterodimers where each monomer consists of nine α-helices forming an amphipathic binding groove. Four α-helices are directly responsible for dimer formation, while the rest of the α-helices form the ‘W’ shape of the protein dimer [38,39]. The N terminal domain of 14-3-3 proteins modulates homo/heterodimerization while the C terminal domain binds client proteins [38]. Activity and modulation of many cellular pathways by 14-3-3 is controlled by phosphorylation events. Target proteins are typically recognized by 14-3-3 through well-defined sequences that can be generalized into three categories: (I) RXX pS/pT XP, (II) RX(Y/F)X pS/pT XP and (III) XX pS/pT X COOH (pS and pT being phosphorylated Ser and Thr, respectively, X being any amino acid). [40] In modes I and II, proline is always located at position +2 in relation to the central pSer/pThr and arginine is at position −3 in mode I. Mode III represents C terminal sequences where pSer/pThr are the penultimate amino acid of the partner protein.

Phosphorylated residues on hDMX or hDM2 are considered relevant to recognition by 14-3-3 proteins, as well as their role in cell signalling is outlined in Fig. 1B [26,41–45]. Relevant sites on hDMX: Ser342 and Ser367 are targeted by several kinases (e.g. AMP [45], Akt [41], Chk2 [26,44], or Chk1 [42]). hDMX phosphorylation in response to stress or DNA damage with consequent enhancement of hDMX mediated hDMX degradation has been reported [44,46], together with a role for Ser367 [43]. Phosphorylation of hDMX on Ser367 by Chk2 was shown to promote 14-3-3 binding, hDMX nuclear import and degradation by hDM2 [26]. Similarly, Chk1 was shown to phosphorylate hDMX at Ser367, to promote 14-3-3/hDMX interaction and cytoplasmic retention of hDMX to counter hDMX-enhanced p53 ubiquitination, leading to its stabilization and activation [42]. However, the potential relevance of both Ser342 and Ser367 to interaction with 14-3-3 has been noted [44,47]. In contrast, Akt has been shown to phosphorylate hDMX on Ser376, promoting 14-3-3 binding, stabilizing hDMX and consequently enhancing hDM2 stability leading to suppression of p53 transcriptional activity [41]. Adenosine monophosphate (AMP) mediated phosphorylation of hDMX on Ser342 has been shown to induce hDMX/14-3-3 interaction resulting in inhibition of p53 ubiquitylation and consequent stabilization [45]. hDM2 phosphorylation in the NLS/ELS region, by PKB/Akt, can lead to its stabilization [48,49], although the effects are complex; phosphorylation on Ser166 and Ser188 are promoted by Akt kinase [50], and result in positive regulation of hDM2 by inhibiting its self-ubiquitination and nuclear translocation [48,51,52]. Phosphorylation at Ser166 and Ser188 does not appear to result in binding of hDM2 to 14-3-3 [53] whilst such an interaction was observed when phosphorylated at Ser166 and Ser186 by Pim kinase, which simultaneously suppresses phosphorylation of Ser188 [53]. Finally, the interaction between hDM2 phosphorylated in the ring domain and 14-3-3σ was observed to accelerate the self ubiquitination of hDM2, with a resulting downstream stabilization of p53 [54]. Thus cellular studies have linked hDMX and hDM2 to 14-3-3 proteins; 14-3-3 proteins may indirectly modulate the interaction between hDMX or hDM2 and p53 and represent potential targets for pharmacological modulation of p53 [21,23] but the molecular bases of these interactions remain unexplored. In this manuscript, we report on the biophysical and structural characterization of peptide/protein interactions that are representative of the interaction between 14-3-3 and hDMX or hDM2. The data reveal the interactions to be mode I [55,56]; whilst such 14-3-3 interactions are not stabilized by typical 14-3-3 interaction stabilizers, for example, Fusicoccin and Cotelynin A [57]. These data provide structural insights that could be utilized in future stabilizer/inhibitor discovery efforts. More significantly, our results reveal that in both cases proximal phosphosites spaced ~20–25 residues apart in both hDMX and hDM2 co-operate to facilitate high-affinity 14-3-3 binding and do so through 1:1 stoichiometry.

**Results**

There are multiple serine and threonine residues in the hDMX and hDM2 proteins that could be phosphorylated (protein sequences with highlighted Ser and Thr residues on hDMX and hDM2, as well as their alignment [35] can be found in Figs S2 and S3). Therefore, 14-3-3Pred [58] was used to identify sequences from hDM2 and hDMX that might serve as 14-3-3 binding sequences. 14-3-3Pred is a webservice that predicts 14-3-3 binding sites in client proteins based on published examples of 14-3-3 interactions with specific amino acids surrounding Ser/Thr. Scores are assigned to each Ser/Thr residue on the protein target whose sequences in FASTA format are queried, using an ensemble approach comprising three different base classification systems: position-specific scoring matrix, support vector machines and artificial neural network. End score values are calculated as the average of three independent methods with a score > 0.5 indicating a high probability of the sequence binding to 14-3-3. The webservice also provides information on the
phosphorylation state for each Ser/Thr residue. The phosphorylated sites on hDMX and hDM2 shown to be relevant for the interaction of 14-3-3 in previous cell-based studies [26, 42–45, 50, 53] were predicted to have a high probability of 14-3-3 binding (Table 1), except pSer342 on hDMX, which showed a low probability of binding (score 0.19) whereas Ser151 on hDMX – a previously unreported site of phosphorylation – was predicted to be a potential 14-3-3 binding site (score 0.75).

Using the 14-3-3Pred results server, several key regions of hDMX and hDM2 were thus identified. Peptides containing these binding regions were designed bearing five/six residues flanking the central phosphorylated residue on either side and prepared by solid-phase peptide synthesis (SPPS). Given the proximity of pSer342 and pSer367 from hDMX, which showed a low probability of binding (score 0.19) whereas Ser151 on hDMX, which showed a high probability of 14-3-3 binding (Table 1), except pSer342 on hDMX, we also prepared doubly phosphorylated sequences for both to probe for avidity effects. Peptides were synthesized using a combination of manual or automated Fmoc SPPS on Rink amide resin (Rink amide ProTide for doubly phosphorylated sequences) and modified at the N terminus by acetylation for isostructural titration calorimetry (ITC), surface plasmon resonance (SPR) and crystallography studies or with 5,6-carboxyfluorescein minus by acetylation for isothermal titration calorimetry studies or with 5,6-carboxyfluorescein. Generally, all peptides exhibited tightest binding towards the 14-3-3 isoforms, followed by γ, β, τ, ζ, σ, respectively. hDMX and hDM2 peptides mimicking one phosphorylation site fit well to a 1 : 1 binding isotherm. In this previously described model, each 14-3-3 dimeric protein can bind two peptides, one per 14-3-3 protomer [38]. K_δ values for all 14-3-3 isoforms are reported in Table 2, but for simplicity, only the 14-

### Table 1. 14-3-3 binding hDM2 and hDMX Sequences studied

| Sequencea | 14-3-3-Pred Score |
|-----------|-------------------|
| hDMX144-158pThr151 | X-STRKRTpT EDDIPTL-NH2 | 0.75 |
| hDMX155-165pSer342 | X-SKLTSHpS TSDITAI-NH2 | 0.19 |
| hDMX166-176pSer367 | X-DCRTPipS APVVRPK-NH2 | 1.28 |
| hDMX162-172pSer342/pSer367 | X-SKLTSHpS TSDITAEPENEGNDVPDCRRTipSp pSer342 and hDMX361-374pSer367 showed affinity for 14-3-3 proteins, whereas peptide hDMX144-158pThr151 exhibited no response under the conditions of the assay (see Fig. S5). None of the corresponding unphosphorylated control hDMX sequences exhibited evidence of binding under the conditions of the assay confirming the role of phosphorylation for binding of pSer342 and pSer367 to 14-3-3 (see Fig. S6). hDM2 peptides representing sites pSer151 and pSer186 showed low micromolar affinities to all 14-3-3 isoforms (Fig. S4).

Generally, all peptides exhibited tightest binding towards the 14-3-3η isoforms, followed by γ, β, τ, ζ, σ, and ε, respectively. hDMX and hDM2 peptides mimicking one phosphorylation site fit well to a 1 : 1 binding isotherm. In this previously described model, each 14-3-3 dimeric protein can bind two peptides, one per 14-3-3 protomer [38]. K_δ values for all 14-3-3 isoforms are reported in Table 2, but for simplicity, only the 14-

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*aX = acetyl or FAM-Ahx where FAM is 5,6-carboxyfluoresceine and Ahx is aminohexanoic acid.
3-3η isoform is discussed below. hDMX\textsubscript{361-374pSer367} exhibited higher affinity ($K_d = 98.8 \pm 4.6$ nM) than hDMX\textsubscript{335-349pSer342} ($K_d = 20.0 \pm 1.2$ μM). hDM2 sequences bound with weaker affinity (hDM2\textsubscript{160-171pSer166} $K_d = 10.2 \pm 0.3$ μM and hDM2\textsubscript{180-192pSer186} $K_d = 31.6 \pm 1.9$ μM). As might be expected the highest affinity sequence (hDMX\textsubscript{361-374pSer367}) is a mode I 14-3-3 sequence/binder (see later) with Pro at the +2 position and Arg at the −3 position.

Because of its dimer structure, 14-3-3 is known to bind multiple binding partners simultaneously, either by binding two different phosphorylated residues from the same target or possibly two phosphorylated sites within a multiprotein complex [31,61–63]. Thus, doubly phosphorylated hDMX and hDM2 peptides were used to assess the effects of proximal phosphorylation on affinity. Two state binding in FA was observed for both doubly phosphorylated hDMX and hDM2 peptides.
(Fig. 2C for hDMX peptide and Fig. S4 for hDM2 peptide). Fitting for the observed biphasic-dose–response curves were performed in the same manner as for the short peptides by fitting each binding event separately to a 1 : 1 model (one peptide to one 14-3-3 monomer) so as to obtain stepwise $K_d$ values. Accurate fitting was possible for 14-3-3 η, γ and β isoforms as reported in Table 2 (for 14-3-3η, hDMX$_{335-373}^{pSer342/pSer347}$ $K_d$ = 30.3 ± 6.4 nM and $K_d$ = 10.67 ± 1.2 μM, hDM2$_{60-192}^{Ser166/pSer186}$ $K_d$ = 49.6 ± 12.1 nM and $K_d$ = 17.5 ± 2.5 μM). Approximately 200-fold higher affinity $K_d$ was observed for hDM2$_{160-192}^{pSer166/pSer186}$ compared to hDM2$_{160-192}^{pSer166/pSer186}$ whilst a more modest 3-fold increase in binding affinity for hDMX$_{335-373}^{pSer342/pSer347}$ was observed in comparison to hDMX$_{361-374}^{pSer347}$ (although enhancement was more pronounced for binding to β and γ isoforms).

Isothermal titration calorimetry was used as an orthogonal assay to confirm the binding affinities of hDMX and hDM2 peptides to 14-3-3η and to gain additional information about the stoichiometry of the interactions. Moreover, such experiments provide thermodynamic data and as a label-free method require no immobilization/functionlization of any of the binding partners. Here data were obtained by titration of the peptide into 14-3-3η protein (Fig. 2D,E and Fig. S7). Thermodynamic parameters for interaction of 14-3-3 with hDMX and hDM2 peptides obtained by ITC are summarized in Table 3.

### Table 2. $K_d$ values for hDMX and hDM2 peptides binding to 14-3-3 proteins (in μM) (increasing affinity is denoted by increasingly darker grey shading).

| Peptide            | 14-3-3   | 14-3-3 | 14-3-3   | 14-3-3 | 14-3-3 | 14-3-3 |
|-------------------|----------|--------|----------|--------|--------|--------|
| hDMX$_{335-349}^{pSer342}$ | 20.0 ± 1.2 | 49.4 ± 11.5 | 107.8 ± 6.0 | 95.8 ± 3.6 | 37.8 ± 0.7 | 135.9 ± 3.6 | 53.4 ± 2.9 |
| hDMX$_{335-373}^{pSer342/pSer347}$ | 0.099 ± 0.005 | 0.51 ± 0.02 | 1.2 ± 0.2 | 0.92 ± 0.03 | 0.53 ± 0.04 | 3.6 ± 0.4 | 2.7 ± 0.2 |
| hDMX$_{335-373}^{pSer342/pSer347} K_{d1}$ | 0.030 ± 0.006 | 0.26 ± 0.009 | – | – | – | 0.091 ± 0.015 | – |
| hDMX$_{335-373}^{pSer342/pSer347} K_{d2}$ | 10.7 ± 1.2 | 7.6 ± 0.8 | – | – | – | 18.7 ± 0.7 | – |
| hDM2$_{60-192}^{pSer166/pSer186}$ | 10.2 ± 0.3 | 11.1 ± 0.5 | 29.3 ± 2.4 | 19.6 ± 0.2 | 3.9 ± 0.3 | 32.7 ± 1.4 | 23.0 ± 1.2 |
| hDM2$_{160-192}^{pSer166/pSer186}$ | 31.6 ± 1.9 | 207.9 ± 6.3 | >250 | 165.8 ± 5.6 | 82.9 ± 2.1 | >170 | >260 |
| hDM2$_{160-192}^{pSer166/pSer186} K_{d1}$ | 0.050 ± 0.012 | – | – | – | 0.089 ± 0.010 | – |
| hDM2$_{160-192}^{pSer166/pSer186} K_{d2}$ | 17.5 ± 2.5 | – | – | – | 28.0 ± 0.8 | – |

Isenthalamic titration calorimetry data showed similar trends for binding of hDMX peptides to 14-3-3η to those observed in the FA assays; binding affinities of hDMX$_{361-374}^{pSer347}$, hDMX$_{335-349}^{pSer342}$ and hDMX$_{335-373}^{pSer342/pSer347}$ were $K_d$ = 870 ± 95 nM, $K_d$ = 21 ± 3 μM and $K_d$ = 14 ± 10 nM. An increase in binding affinity when both binding sites of hDM2 are present was also confirmed, with $K_d$ = 151 ± 57 nM. The stoichiometry between doubly phosphorylated hDMX and hDM2 peptides and 14-3-3η was found to be approximately 0.94 indicating a 1 : 1 binding model. If two phosphorylated sites from one peptide were binding simultaneously, the expected stoichiometry would be around 0.5. The contrasting absence of two binding events in the ITC data, when compared to the FA experiments, can be accounted for by the different experimental configurations. In the former, titration of the peptide into protein means that protein begins in excess and becomes saturated with the highest affinity site, whereas for FA, peptide begins in excess and protein (and therefore monomer/dimer ratio) varies in concentration across the experiment.

Finally, association and dissociation rates for the binding of hDMX and hDM2 peptides to 14-3-3η were determined by SPR (Fig. 2E,F and Figs S8 and S9). Similar $K_d$ values were confirmed from dose–response SPR. SPR measured affinities were broadly in agreement with $K_d$ values determined from FA and ITC experiments (Table 2 and 3). For peptides exhibiting

### Table 3. Thermodynamic parameters for hDMX(2) peptides/14-3-3 proteins.

| Peptide            | $K_d$ (ITC) | Δ$G$ (kJ·mol$^{-1}$) | Δ$H$ (kJ·mol$^{-1}$) | −Δ$S$ (kJ·mol$^{-1}$) | $n$ | $K_d$ (SPR)$^a$ | $k_{on}$ (1/M·s$^{-1}$) | $k_{off}$ 1/s |
|-------------------|-------------|-------------------|-------------------|-------------------|-----|---------------|----------------------|----------------|
| hDMX$_{335-349}^{pSer342}$ | 21 ± 3 μM | −26.7 | −21.4 ± 1 | −5.3 | 0.72 | 37 μM | – | – |
| hDMX$_{361-374}^{pSer347}$ | 870 ± 95 nM | −34.6 | −16.1 ± 0.2 | −18.5 | 1.28 | 189 nM | 3.89 × 10$^8$ | 0.1208 |
| hDMX$_{335-373}^{pSer342/pSer347}$ | 14 ± 10 nM | −44.7 | −22.1 ± 0.7 | −22.6 | 0.94 | 8.6 nM | 1.51 × 10$^9$ | 0.0048 |
| hDMX$_{160-192}^{pSer166/pSer186}$ | 151 ± 57 nM | −38.9 | −12.3 ± 0.6 | −26.6 | 0.94 | 156 nM | 1.45 × 10$^9$ | 0.1215 |

$^a$Dose–response curves were fitted by plotting RUs against the peptide concentration.
lower affinities \((hDMX_{335-349}p^{\text{Ser342}}, hDM2_{160-171}p^{\text{Ser166}}\) and \(hDM2_{180-192}p^{\text{Ser186}}\) it was only possible to determine steady-state affinities, but not their kinetic profile (see Supporting Information). The doubly phosphorylated peptides showed the expected increase in affinity, which could be attributed to a much slower \(k_{\text{off}}\) rate in comparison to singly phosphorylated \(hDMX_{361-374}p^{\text{Ser367}}\) and \(hDMX_{335-349}p^{\text{Ser342}}\) peptides.

The 1 : 1 stoichiometry between the doubly phosphorylated peptides and 14-3-3 contrasts with previously reported work [61,62], although it may be that the length and/or accessible conformation of the \(hDMX\) and \(hDM2\) peptides are incompatible with the simultaneous engagement of both 14-3-3 protomers in the dimeric structure. To further explore this behaviour and understand the basis of cooperativity and affinity enhancement we performed additional experiments. To exclude the possible formation of higher-order oligomers (e.g. 14-3-3 tetramers held together by two peptides bridging two dimers), an analytical ultracentrifugation experiment was performed. The 14-3-3\(\eta\) (at the concentration of 14 \(\mu\)M, approximately corresponding to the second binding events observed in FA, Fig. 2C) and \(hDMX_{335-373}p^{\text{Ser342}/p^{\text{Ser367}}}\) peptide in two different concentration ratios (1 : 1 and 1 : 0.5, protein vs. peptide) were analysed. The sedimentation coefficient value remained unchanged for 14-3-3 protein alone and when varying peptide/protein concentrations, indicating that only one dimer of 14-3-3\(\eta\) is involved in binding (see Fig. S10). Secondly, peptide length may also have a significant effect on the affinity and binding mode. This has been demonstrated in a recent study with p53 peptides with 14-3-3, where variation in the peptide length (from 9 to 32 residues) correlated with increased binding affinities [64]. Two long variants of \(hDMX_{335-373}p^{\text{Ser342/pSer367}}\) peptide were synthesized, with only one phosphorylation site: pSer367 or pSer342, extending the peptide length by 26 amino acids. The two peptides were tested in FA assays to decouple the influence of peptide length from the co-operative influence of having two phosphosites (Fig. 3). The \(hDMX_{335-373}p^{\text{Ser342}}\) peptide showed an increase in binding affinity with 14-3-3\(\eta\) \((K_d = 5.4 \pm 1.1 \mu\text{M})\) as expected, in comparison to the short version of the \(hDMX_{335-349}p^{\text{Ser342}}\) peptide \((K_d = 20.0 \pm 1.2 \mu\text{M})\). On the other hand, increasing the peptide length for \(hDMX_{335-373}p^{\text{Ser367}}\) decreased the binding affinity \((K_d = 0.8 \pm 0.1 \mu\text{M})\) and significantly lowered the maximum anisotropy signal. The lower maximum anisotropy signal observed might be due to the fluorophore being moved approximately 20 amino acids further from the binding pSer-367 (see earlier discussion on fluorophore mobility). Whilst there is a moderate increase in binding affinity of \(hDMX_{335-373}p^{\text{Ser342}}\) on increasing the peptide length this is insufficient for high-affinity interaction. Similarly, the slight loss in potency for the longer \(hDMX_{335-373}p^{\text{Ser367}}\) peptide relative to \(hDMX_{361-374}p^{\text{Ser367}}\) is noteworthy in considering the interaction with full-length protein as these data hint at interference from the additional residues, and is therefore likely to be important in relating these results to the behaviour for the full-length proteins. Taken together, the data indicate that proximal phosphorylation of both sites in \(hDMX_{335-373}p^{\text{Ser342/pSer367}}\) is required for high affinity 1 : 1 stoichiometric binding between \(hDMX\) and

\[\text{Fig. 3. Difference in the affinity between short } hDMX_{335-349}^{p^{\text{Ser342}}} \text{ and } hDMX_{335-373}^{p^{\text{Ser342/pSer367}}} \text{ peptides and their longer singly phosphorylated variants } hDMX_{335-373}^{p^{\text{Ser342}}} \text{ and } hDMX_{361-374}^{p^{\text{Ser367}}} \text{ as determined by fluorescence anisotropy titration (tracer 50 nM, 10 mM HEPES, 150 mM NaCl, 0.1% Tween 20, 0.1% BSA pH 7.4, concentration of proteins given as 14-3-3 monomer concentration, error bars represent SD for } n = 3 \text{ replicates).} \]
14-3-3 protomers. Given the absence of 1:2 binding, the presence of multiple sites may increase affinity through statistical rebinding or interaction with a second non-canonical phosphate-binding site on the surface of 14-3-3. The second binding event observed only in fluorescence anisotropy experiments, may represent a further non-specific binding event or an interaction linked to 14-3-3 dimerization [60] and will be investigated in future work.

Structural characterization of the interaction between 14-3-3 and hDMX or hDM2 peptides

The W-shaped structure of a 14-3-3 dimer provides a groove for phosphorylated client proteins to bind. One dimer usually accommodates two copies of phosphorylated sites from one target, but there are examples of two different targets simultaneously binding to one dimer of 14-3-3 proteins [61,62]. Phosphorylated Ser/Thr residues from partner proteins generally bind directly to Lys49 and Arg56 in helix α3, and Arg127 and Tyr128 in helix α5 as they form a basic binding pocket in otherwise acidic surroundings. Structural data were obtained by solving three crystal structures of hDMX361-374 as pSer367 (PDB: 6YR5), hDMX180-192 as pSer186 (PDB: 6YR6) and hDMX335-373 as pSer342/pSer367 (PDB: 6YR7) in complex with 14-3-3ΔC (the C-terminus was truncated for the crystallography purposes). Refinement statistics for all images of each structure with electron density for pep-map. The phosphoserine of hDMX361-374 was observed to adopt an unusual conformation by taking a sharp turn. A closer examination of the groove of monomer A indicated partial ligand occupation and this peptide conformation is stabilized by hydrogen bonds with water molecules.

The structure of a doubly phosphorylated peptide hDMX335-373 as pSer342/pSer367 in complex with 14-3-3ΔC was solved to 2.1 Å resolution (Fig. 4C). Here, 14-3-3ΔC crystallized as a dimer in space group C121 with the asymmetric unit containing one dimer (green) with additional electron density for the peptide (cyan) observed in both monomeric units of 14-3-3ΔC (Fig. 4C and Fig. S13 for additional images). Based on previously published structures of doubly phosphorylated peptides, it was expected to observe pSer342 binding to monomer B and pSer367 binding to monomer A [31,61,65]. The electron density within the binding groove of monomer B could unambiguously be assigned to pSer342, whereby eight amino acids of the peptide could be resolved (HSL(pS)TSDI). The peptide was observed to adopt an unusual conformation by taking a sharp turn. A closer examination of the pSer342 binding site in monomer B of 14-3-3ΔC shows pSer342 binding to a basic pocket as expected, making polar contacts with Arg56, Arg129 and Tyr130. In this orientation, the Ile+4 from the peptide makes hydrophobic contact with Leu218 and Ile219 on the protein (represented as spheres). Peptide residues Thr+1 and Ser+2 make polar contacts with Lys122, Asn175, Glu182 and Trp230, while Asn226 makes polar contacts with the backbone of the peptide. Overall, the C-terminus of the peptide takes a sharp turn after the pSer-Thr residues and this peptide conformation is stabilized by hydrogen bonds between water molecules and Thr+1, Ser+2 and Asp+3 peptide residues. In contrast, the electron density map within the binding groove of monomer A indicated partial ligand occupancy and could be interpreted in different ways, indicating an overlay of multiple conformations (both pSer342 and pSer367 sites, with the dominant electron
density map ascribable to pSer342 binding see Fig. S13 for additional images and discussion).

**Discussion**

We used FA, ITC and SPR to determine the binding affinity of hDMX and hDM2 peptides to 14-3-3 proteins. hDMX335-373pSer342, hDMX316-374pSer367, hDM2160-171pSer166 and hDM2180-192pSer186 mimicking one phosphorylation site fit well to a 1 : 1 binding isotherm and exhibited greatest affinity towards the 14-3-3η isofrom, followed by γ, β, τ, ζ, σ and ε, respectively. The highest affinity peptide (hDMX361-374pSer367) bound strongly ($K_d = 98.8 \pm 4.6 \text{ nM}$), whereas the
remaining sequences bound 14-3-3 less tightly (hDMX335-349 pSer342, $K_d = 20.0 \pm 1.2$ $\mu$m $\{14-3-3$ isoform$\}$. hDMX2160-192 pSer166 $K_d = 10.2 \pm 0.3$ $\mu$m and hDMX2180-192 pSer186 $K_d = 31.6 \pm 1.9$ $\mu$m). Interestingly, different maximum anisotropies were observed for peptide binding across the 14-3-3 isoforms, which may reflect a difference in the 14-3-3 dimerization affinity [60] or mobility of the fluorophore. Binding experiments with doubly phosphorylated peptides were observed to result in two binding events. As expected for a multivalent interaction with 14-3-3 proteins [63], the first binding event in both cases was higher affinity than either monovalent interaction, whereas the second binding event was in the micromolar regime (for 14-3-3 $\eta$, hDMX335-373 pSer342 pSer367 $K_d1 = 30.3 \pm 6.4$ $\mu$m and $K_d2 = 10.67 \pm 1.2$ $\mu$m, hDMX2160-192 pSer166 pSer186; $K_d1 = 49.6 \pm 12.1$ $\mu$m and $K_d2 = 17.5 \pm 2.5$ $\mu$m). This enhancement in affinity for the first binding event was also observed in ITC experiments which lacked the second binding event (but were performed by titration of peptide into protein in contrast to the FA experiments) and SPR experiments which indicated that a slower $k_{off}$ was associated with co-operative binding of doubly phosphorylated peptides. Importantly, for the doubly phosphorylated peptides, binding was negatively co-operative (i.e. less than additive; Table 4).

Taken together the data can be rationalized by assuming that the dimerization affinity of 14-3-3 differs across isoforms and occurs within the concentration regime of the FA experiments (Fig. 5). As the titration progresses, once 1 : 1 stoichiometry is attained, further addition of 14-3-3 results in unbound 14-3-3 monomer and eventually all peptides will be bound in complexes containing one peptide to two 14-3-3 monomers. For the doubly phosphorylated peptides early in the titration peptides bind with high affinity in 1 : 1 stoichiometry – higher affinity likely results from statistical rebinding or a second phosphosite on the surface of the 14-3-3 protein. The second binding event either derives from a non-specific interaction, or arises from 14-3-3 dimerization [60] but is visible in the titration as a distinct binding event due to the strong affinity of the first event. Such behaviour is not observed in ITC experiments because the peptide is titrated into 14-3-3 protein so its concentration does not vary, and, the peptide becomes the saturating component. Considering the structural data for hDMX361-374 pSer367, hDMX2180-192 pSer186 and hDMX335-373 pSer342 pSer367 bind to 14-3-3 directly between K49 and R56 in helix $\alpha3$, and R127 and Y128 in helix $\alpha5$ as is typical for 14-3-3 client proteins. The hDMX361-374 pSer367 peptide is a Mode I sequence (RXXpS/pTPX) which can account for its enhanced affinity in comparison to the other sequences [38,56]. Curiously, for the structure containing the longer hDMX335-373 pSer342 pSer367 peptide, the sequence surrounding pSer342 dominates in the electron density for bound ligand despite being the weaker 14-3-3 binding site of the two individual pSer sequences, this may favour the hypothesis of a second phosphorecognition site over the statistical rebinding model. Moreover, in the context of earlier cellular studies [26,41–45,48–53] the data highlight the complex nature by which phosphorylation regulates hDMX in particular [26,41–45]. pSer367 may be sufficient for 14-3-3 binding and the consequent effects on localization, stability and ultimately p53 function; the presence of both pSer342 and pSer367 confers some increase in 14-3-3 binding affinity (particularly for other isoforms), and it is plausible that accessibility of the binding sites in the full-length protein may play a crucial role that requires phosphorylation at both sites in vivo. In contrast, a much clearer difference in binding affinity for hDMX2160-192 pSer166 pSer186 is observed in contrast to both monophosphorylated peptides and points to a functional requirement for double phosphorylation.

In conclusion, we have performed a biophysical and structural study on the interaction of 14-3-3 with peptides derived from hDMX and hDM2. We have shown that peptides derived from hDMX containing pSer342 or pSer367 and from hDM2 containing pSer166 or pSer186 exhibit affinity for 14-3-3 as demonstrated by FA, ITC and SPR. Structural analyses reveal that the peptides bind to 14-3-3 directly between K49 and R56 in helix $\alpha3$, and R127 and Y128 in helix $\alpha5$ as is typical for 14-3-3 client proteins. For high-affinity binding, proximal phosphosites are beneficial and in the case of hDM2 essential. In contrast to other doubly phosphorylated peptides of this length the interaction involves one

Table 4. Thermodynamic values for binding of hDM2 and hDMX peptides to 14-3-3$\eta$

| Peptide | $\Delta G_{14-3-3\eta}$ (kJ mol$^{-1}$)$^a$ | $\Delta G_{14-3-3\eta}$ (kJ mol$^{-1}$)$^b$ |
|---------|---------------------------------|---------------------------------|
| hDMX335-349 pSer342 | $-21.1$ | $-26.7$ |
| hDMX361-374 pSer367 | $-34.3$ | $-34.6$ |
| hDMX335-373 pSer342 pSer367 | $-37.2$ | $-44.7$ |
| hDMX2160-192 pSer166 | $-22.8$ | n.d |
| hDMX2180-192 pSer186 | $-20.0$ | n.d |
| hDMX2160-192 pSer166 pSer186 | $-36.0$ | 38.9 |

$^a$Calculated from $K_d$ values determined by fluorescence anisotropy titration of FAM-Ahx labelled peptides with 14-3-3$\eta$; $^b$Determined by ITC for Acetyl labelled peptides titration with 14-3-3$\eta$. 

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peptide for each 14-3-3 protomer [61–63]. The additional affinity likely derives from a reduced $k_{\text{off}}$ rate for doubly phosphorylated peptides. Importantly, the results suggest that sequential phosphorylation of proximal sites in both hDMX and hDM2 plays a role in promoting changes in their localization and degradation in a cellular context. Thus, these data provide structural insight on the indirect effects of 14-3-3 on the p53 pathway, which may inform the development of chemical intervention strategies (e.g. use of kinase inhibitors) and provide starting points for structure-based ligand design of 14-3-3/hDMX and 14-3-3/hDM2 modulators, both stabilizers and inhibitors.

Materials and methods

Peptide synthesis

General remarks

Resins and amino acids were purchased from either Sigma–Aldrich (part of Merck Life Science UK Limited, Gillingham, Dorset, UK) or Novabiochem (part of Merck Life Science UK Limited, Gillingham, Dorset, UK). All amino acids were N-Fmoc protected and side chains protected with Boc (His, Lys), t-Bu (Asp, Glu, Ser, Thr), Pbf (Arg), Trt (Asn, Gln). Peptides were synthesized either manually or using a microwave-assisted automated peptide synthesizer (CEM Liberty Blue) on a 0.05 or 0.1 mmol scale. DMF used in peptide synthesis was of ACS grade and from Sigma–Aldrich.

Manual peptide synthesis

Manual peptide synthesis followed this cycle: swelling of a resin (20 min) in cartridge used for solid-phase synthesis, washing (DMF, 3 $\times$ 2 mL $\times$ 2 min), deprotection (Method A), and coupling of the desired amino acid (Method B), where successful coupling and deprotection were determined by a colour test (Method C). Acetylation (Method D) or coupling of a fluorescent dye (Method E) were performed prior to cleavage (Method F).

Method A: Deprotection N-terminal Fmoc-protecting groups were removed by adding 20% piperidine in DMF ($5 \times 2 \text{ mL} \times 2 \text{ min}$) and washed with DMF ($5 \times 2 \text{ mL} \times 2 \text{ min}$) after.

Method B: Manual coupling of amino acid and Ahx.

The desired amino acid or Ahx (5 equiv.), DIPEA (10 equiv.) and HCTU (5 equiv.) were dissolved in DMF...
(2 mL) and added to the resin, followed by agitation for 1 h. Reagents were removed by filtration and the resin was washed with DMF (3 × 2 mL × 2 min).

Method C: Kaiser test.

Successful coupling or deprotection for any residue coupled manually was determined by the Kaiser test. A few resin beads were transferred into a vial and mixed with 2 drops of each of the solutions:
1. Nihydrin (5% w/v) in ethanol,
2. Phenol (80% w/v) in ethanol,
3. 1 mM KCN (aq.) in pyridine (2% v/v).

The solution was heated at 100 °C for 5 min before observing the change in colour. Successful deprotection was observed by colour of the beads changing into blue, when successful coupling gave no change in colour.

Method D: N-terminal acetylation.

Acetic anhydride (10 equiv.) and DIPEA (10 equiv.) were dissolved in DMF (2 mL) and the solution was transferred to the resin. After 2 h, the resin was drained and washed with DMF (3 × 2 mL × 2 min). Successful capping was determined by colour test (Method C).

Method E: N-terminal Fluorescent Dye coupling.

5,6-carboxyfluorescein (5 equiv.), DIPEA (5 equiv.) and HCTU (5 equiv.) were dissolved in DMF (2 mL) and added to the resin, followed by agitation for 1 h. Reagents were filtered and the resin was washed with DMF (3 × 2 mL × 2 min) ahead of cleavage and deprotection.

Method F: Cleavage and deprotection of peptides of the resin.

After elongation and acetylation or fluorescent dye coupling were complete, the resin was washed with CH3Cl (2 × 2 mL × 2 min), Et2O (5 × 2 mL × 2 min) and dried under vacuum. Peptides were cleaved and side-chain deprotected using ‘Reagent K’ TFA : EDT : Thioanisole : Phenol : H2O 82 : 3 : 5 : 5 : 5 (2 mL × 3 h). The peptide was precipitated in ice-cold Et2O (10 mL) and placed in a centrifuge (1107 g × 5 min). The supernatant was removed and the precipitate resuspended in ice-cold Et2O and placed in a centrifuge again (3 x). The precipitate was dried under a stream of nitrogen overnight, before being dissolved in H2O and lyophilized.

Automated peptide synthesis method

Peptides prepared using automated peptide synthesizer followed cycles described below. Resin loading cycle cleans the reaction vessel, washes with DMF : DCM (1 : 1), transfers resin to the reaction vessel, washes with DMF : DCM (1 : 1), and drains the vessel at the end of a cycle. Deproteinization and coupling cycle consists of: washing with DMF (4 mL), adding 20% piperidine in DMF (6 mL), microwave deproteinization cycle (30 s), washing with DMF (4 + 4 + 4 + 4 mL), the addition of amino acid (2.5 mL, 5 eq or 3 eq for phosphorylated amino acids), coupling reagent (1 mL, 5 eq) and activator base (0.5 mL, 5 eq), coupling microwave cycle (5 min), washing with DMF (2 mL) and draining. HCTU and DIPEA were used during automated peptide synthesis as well. As a rule, all amino acids were coupled using 75 °C coupling and deprotection cycles up to Ser(OP(OBzl)OH)-OH or Thr(OP(OBzl)OH)-OH, where conventional coupling and deprotection method was used (coupling at the rt for 2 h, deprotection at rt for 15 min) as well for every amino acid following pSer/pThr. After the final residue was coupled, the resin was ejected from the reaction vessel. Alx coupling, deprotection, acetylation or fluorescent dye coupling, and cleavage were performed manually using the methods described above.

Peptide purification

Crude peptides were dissolved in H2O or DMSO and purified by UV- or MS-directed HPLC. Jupiter Proteo (250 × 21.2 mm) or a Kinetex EVO C18 (250 × 21.2 mm) preparative column (reversed-phase) was used with an increasing gradient of acetonitrile in water with 0.1% formic acid, over 30 min at the flow rate of 10 mL·min⁻¹. Fractions containing peptides were combined, concentrated and lyophilized. The purity of peptides was assessed by analytical HPLC and HRMS.

Protein expression and purification

The pl-ProEx HTb-His-14-3-3 constructs were expressed in BL21(DE3) cells. A single colony from a freshly made agar plate (8 g LB broth mixed with 8 g agar in 400 mL, autoclaved and poured into Petri dishes to use for the transformation of plasmids) was picked and mixed with 5 mL of LB media with ampicillin to inoculate a starter culture overnight at 37 °C. The cells were grown in 2 L of TB media (48 g peptone, 24 g yeast, 4.6 g KH2PO4, 24 g KHPO4, 5 mL glycerol in 2 L of dH2O, autoclaved for 20 min at 121 °C) at 37 °C until the OD reached 0.4–0.6. Expression was induced by adding 0.4 mM IPTG and agitating overnight at 18 °C. The expression culture was spun down (7871 g, 20 min, 4 °C), resuspended in 200 mL of lysis buffer (50 mL Tris, 300 mM NaCl, 12.5 mM imidazole, 2 mM β-mercaptoethanol) with 5 mM MgCl2 and DNAse (1 : 1000). The cells were lysed by French press or sonication and the solid fragments were removed by centrifugation (49 192 g, 30 min, 4 °C). The cleared lysate was loaded on a Ni2+–NTA column, washed with 50 mL Tris, 300 mM NaCl, 12.5 mM imidazole, 2 mM β-mercaptoethanol, 0.1% triton X-100, and the protein was eluted with 50 mL Tris, 300 mM NaCl, 250 mM imidazole and 2 mM β-mercaptoethanol. Imidazole was removed by overnight dialysis using the Tris buffer (50 mL Tris, 300 mM NaCl, 2 mL β-mercaptoethanol), full-length proteins were concentrated by centrifugation, rebuffered in HEPES buffer (25 mM HEPES pH 7.5, 100 mM NaCl, 10 mM MgCl2 and 0.5 mM TCEF) and stored in −80 °C.
ΔC proteins were dialyzed with TEV protease to remove the expression tag and purified again on Ni²⁺-NTA column, followed by size-exclusion chromatography (HiLoad 16/600 Superdex 75 pg column) in HEPES buffer (20 mM HEPES, 150 mM NaCl, 2 mM DTT). ΔC proteins were concentrated and rebuffed for ~80 °C storage. All proteins were analyzed by ESI QTOF-MS.

Fluorescence anisotropy

Direct titration assay

All assays were performed in 384 well plates (each experiment was run in triplicates) and data were collected by Perkin Elmer EnVision 2013 plate reader with excitation at 480 nm (30 nm bandwidth), polarized dichroic mirror at 505 nm and emission at 535 nm (40 nm bandwidth, S and P polarised). Experiments were carried out in HBS buffer (10 mM HEPES, 150 mM NaCl, pH 7.4) + 0.1% Tween 20 + 0.1% BSA). A ½ fold dilution series was performed in the titration, with plates read after 30 min, 4 h and 20–24 h. These gave consistent data and values when fitted. Collected data were processed in Microsoft Excel using the equations below. Total intensity I and anisotropy r were calculated using Equations 1 and 2 for each well. Average anisotropy was plotted against protein concentration using OriginPro and a logistic curve was fitted to give r_min and r_max. Using Equation 3 anisotropy was converted into fraction bound and multiplied by peptide concentration to be fitted in Origin using Equation 5 to obtain K_d values.

\[
I = 2PG + S \\
r = \frac{S - PG}{I}
\]  

(1)  

(2)  

\[
L_b = \frac{(r - r_{\text{min}})}{\lambda(r_{\text{max}} - r) + r - r_{\text{min}}}
\]  

(3)  

\[
y = r_{\text{min}} + \frac{r_{\text{max}} - r_{\text{min}}}{1 + 10^{(x - \log k_d)}}
\]  

(4)  

\[
y = \frac{((K + X + FL) - \sqrt{(K + X + FL)^2 - 4xFL})}{2}
\]  

(5)  

r = anisotropy, I = total intensity, P = perpendicular intensity, S = parallel intensity, L_b = fraction ligand intensity, \( \lambda = I_{\text{bound}}/I_{\text{free}} \approx 1 \), FL = fluorescent ligand concentration, K = K_d.

Surface plasmon resonance

Experiments were performed at 25 °C using a Biacore T200 instrument (GE Healthcare, Buckinghamshire, UK). For immobilization, the running buffer was HBS buffer (HEPES, NaCl, Tween-20, pH 7.4). 14-3-3q was immobilized on an NTA sensor chip (Series S Sensor Chip GE Healthcare) using NTA reagent and Biacore Amine Coupling Kit. Briefly, 14-3-3q was diluted to 0.2 mg·mL⁻¹ and 200 µL was injected over a chip surface that had been activated with an injection of Ni²⁺, followed by 140 µL of 1 : 1 NHS/EDC. A 30 µL of 0.5 M ethanolamine was then injected to cap the excess free amine groups. Immobilization levels of the 14-3-3q were found to be around 2000 response units (RU) for kinetic measurements. hDMX and hDM2 peptides were serially diluted 11 times from concentrations 10 times the K_d values measured in FA, on a 96-well plate. Then, 20 µL of these solutions were injected over a 14-3-3q immobilized surface for 1 min at the 20 mL·min⁻¹ flow rate, followed by a 4 min regeneration period with HBS buffer. Kinetics and the binding affinity were calculated using the BIACORE evaluation software (Buckinghamshire, UK). Standard dose–response curves were fitted by plotting RUs against the peptide concentration.

Isothermal titration calorimetry

Isothermal titration calorimetry experiments were carried out on a MicroCal iTC200 in HBS buffer at 25°C (25 mM HEPES pH 7.5, 100 mM NaCl, 10 mM MgCl₂, 0.5 mM TCEP). Peptides (0.2–1 M) in the syringe were titrated into a cell containing 14-3-3q (0.02 or 0.1 M). Data were fitted using MICROCAL software (Northampton, MA, USA) to give binding constant (K_d), enthalpy (ΔH), entropy (ΔS) and binding stoichiometry (n).

Protein crystallography

A solution of 14 mg·mL⁻¹ 14-3-3-σΔC with hDMX_361-374_pSer342/pSer367 peptide in 1 : 2 molar ratio was incubated overnight in crystallization buffer (20 mM HEPES pH 7.5, 2 mM MgCl₂ and 2 mM BME). Crystals were obtained at 4 °C in a hanging drop vapour diffusion set up with 0.2 mM sodium sulfate, 0.1 M Bis-Tris propane, 20% w/v PEG 3350 at pH 8.5 as precipitation buffer. Protein/peptide solution and precipitation buffer were mixed in a 1 : 1 ratio in a total volume of 2 µL. Diffraction data were recorded at the DESY, Petra III P11, Hamburg to a resolution of 2.2 Å. hDMX_335-374_pSer342/pSer367 crystals were obtained at 4 °C after overnight incubation of 12 mg·mL⁻¹ 14-3-3-σΔC with hDMX_335-374_pSer342/pSer367 peptide (1 : 2 ratio) in sodium citrate dihydrate, bis-tris propane, pH 8, 20% PEG 3350. hDMX_335-374_pSer342/pSer367 crystals were obtained at 4 °C after overnight incubation of 12 mg·mL⁻¹ 14-3-3-σΔC with hDMX_335-374_pSer342/pSer367 peptide (1 : 2 ratio) in 0.075 M HEPES, pH 7.5, 0.75 M NaAc 3H₂O and 25% glycerol. Diffraction data for hDMX_335-374_pSer342/pSer367 and hDMX_335-374_pSer342/pSer367 crystals were recorded at the Diamond Light source, UK with a resolution of 2.1 Å for hDMX_335-374_pSer342/pSer367 and a resolution of 1.75 Å for hDMX_335-374_pSer342/pSer367. The CCP4 software package [66] was used for

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structure determination by molecular replacement with 4DAT (PDB accession code) as a template for the 14-3-3p structure. Further rounds of manual model building and refinement were performed using COOT [67] and REFMAC [68] respectively. Data collection and refinement statistics for each structure are provided in Table 5.

### Analytical ultracentrifugation

AUC-SV experiments were performed using a Beckman Coulter Optima XL-I ultracentrifuge. Samples were prepared in HBS buffer (10 mM HEPES, 150 mM NaCl, pH 7.4) and loaded into 12 mm aluminium centrepieces with sapphire windows. Three samples were run: (I) 14 μM 14-3-3pSer342, (II) 14 μM 14-3-3pSer367, and (III) 14 μM 14-3-3 + 14 μM hDMX335-374pSer342/pSer367, with buffer as a reference, and recorded at 206 070 g in the An50-Ti rotor, at the 25 °C. The diffusion-deconvoluted sedimentation coefficient distributions c(s) were calculated using the SEDFIT program (Bethesda, MD, USA).

### Circular dichroism spectroscopy

Spectra were recorded on a Chirascan circular dichroism spectropolarimeter (Applied Photophysics, Leatherhead, UK), using 1 mm cells, scan speed of 5 nm·min⁻¹, 2 nm bandwidth and 180–260 nm range. The experiments were performed in 50 mM sodium phosphate buffer, pH 7.5 at 20 °C. The spectra were averaged over three repeats with

| Table 5. Data collection and refinement statistics. Values in parenthesis correlate to high-resolution shell if not indicated otherwise. |
|---------------------------------------------------------------|
| **PDB ID** | hDMX351-374pSer367 | hDMX345-374pSer342/pSer367 | hDMX345-374pSer342/pSer367 |
| Source | DESY, PETRA III | DLS | DLS |
| Beamline | P11 | I03 | I03 |
| Wavelength | 1.033208 | 0.9763 | 0.9763 |
| Resolution (Å) | 74.53–2.25 | 45.77–2.11 | 73.26–1.75 |
| Space group | C 1 2 1 | P 1 | P 1 |
| Cell constants a, b, c (Å) | 97.3 (74.53–2.25) | 97.6 (71.50–2.25) | 97.6 (71.50–2.25) |
| % Data completeness (in resolution range) | 95.15° 113.14° 93.84° | 90.00° 101.97° 90.00° | 90.00° 101.97° 90.00° |
| Total reflections | 438 829 (26 508) | 83 977 (7971) | 445 180 (21 904) |
| Unique reflections | 75 667 (4474) | 42 037 (3995) | 127 733 (12 631) |
| Redundancy | 5.8 (5.9) | 2.0 (2.0) | 3.5 (3.5) |
| Rmerge | 0.11 | 0.04 | 0.07 |
| < (θ (θ) | 5.3 (at 2.25 Å) | 0.68 (at 2.10 Å) | 1.33 (at 1.75 Å) |
| Refinement statistics | | | |
| Resolution range (Å) | 74.53–2.25 (2.33–2.25) | 45.77–2.11 (2.19–2.11) | 56.90–1.75 (1.81–1.75) |
| Wilson B-factor (Å²) | 27.3 | 35.8 | 25.3 |
| Anisotropy | 0.338 | 0.397 | 0.241 |
| Bulk solvent ksol(Å²), Bsol(Å²) | 0.30, 31.7 | 0.36, 40.0 | 0.41, 47.8 |
| L-test for twinning | No twinning to report. | No twinning to report. | No twinning to report. |
| Estimated twinning fraction | No twinning to report. | No twinning to report. | No twinning to report. |
| Fo, Fc correlation | 0.93 | 0.94 | 0.94 |
| Total number of atoms | 7709 | 7376 | 15129 |
| Average B, all atoms (Å²) | 34.0 | 54.0 | 40.0 |
| R factor (%) | 19.71 (28.03) | 21.56 (34.09) | 19.89 (28.06) |
| Rfree (%) | 21.94 (30.14) | 25.00 (34.57) | 22.38 (29.45) |
| Rfree test set | 2993 reflections (4.85%) | 2108 reflections (5.02%) | 6318 reflections (5.03%) |
| No. of protein non-H atoms | 7709 | 3848 | 8038 |
| No. of macromolecules non-H atoms | 7333 | 3650 | 7296 |
| No. of ligand non-H atoms | 5 | 45 | 101 |
| No. of water molecules | 371 | 179 | 693 |
| R.m.s.d bond lengths (Å) | 0.017 | 0.004 | 0.006 |
| R.m.s.d bond angles (°) | 1.95 | 0.86 | 0.93 |
| Ramachandran plot | | | |
| Favoured region (%) | 97.68 | 98.23 | 98.45 |
| Outliers (%) | 0.22 | 0.00 | 0.00 |

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a buffer baseline subtracted. Protein concentrations of approximately 0.2 mg mL$^{-1}$ were used for all proteins.

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**Conflict of interest**

CO is co-founder and shareholder of Ambagon Therapeutics. The authors declare no competing financial interests.

**Data availability statement**

The data that supports the findings of this study are available in [Figs 1–4] and the Supporting Information of this article. The structural data that support these findings are openly available in the wwPDB at https://doi.org/10.2210/pdb6YR5/pdb, https://doi.org/10.2210/pdb6YR6/pdb and https://doi.org/10.2210/pdb6YR7/pdb.

**Author contributions**

SS, CO, SLW and AJW conceived and designed the research program, SS designed studies and performed the research with support from MW, CT and CO on crystallographic analyses. The manuscript was written by SS and AJW and edited into its final form by SLW and AJW with contributions from all authors.

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**Supporting information**

Additional supporting information may be found online in the Supporting Information section at the end of the article.

**Fig. S1.** Sequence alignment of 14-3-3 proteins.

**Fig. S2.** hDMX and hDM2 sequences.

**Fig. S3.** Sequence alignment for hDM2 and hDMX proteins.

**Fig. S4.** Fluorescence anisotropy assays for the hDM2 peptides.

**Fig. S5.** Fluorescence anisotropy assays for the hDMX_{P}Thr^{151} peptide with all isoforms of 14-3-3.
Fig. S6. Fluorescence anisotropy assays for unphosphorylated hDMX and hDM2 peptides with 14-3-3-η.

Fig. S7. ITC data for the interaction of hDMX and hDM2 peptides with 14-3-3-η.

Fig. S8. Dose–response SPR experiments for the binding of hDMX and hDM2 peptides to immobilized 14-3-3-η.

Fig. S9. Data Fitting for dose–response SPR experiments for the binding of hDMX and hDM2 peptides to immobilized 14-3-3-η.

Fig. S10. AUC data for 14-3-3-η and hDMX335-374 pSer342/pSer367.

Fig. S11. Additional images for the hDMX361-374 pSer367/1433σ structure (PDB: 6YR5).

Fig. S12. Additional images for the hDM2180-192 pSer186/1433σ structure (PDB: 6YR6).

Fig. S13. hDMX335-374 pSer342/pSer367/1433σ structure (PDB ID: 6YR7).

Table S1. High-resolution mass spectrometry data for peptides.