Proposed allosteric inhibitors bind to the ATP site of CK2α

Paul Brear¹, Darby Ball², Katherine Stott¹, Sheena D’Arcy² and Marko Hyvönen¹*

¹ Department of Biochemistry, University of Cambridge, CB2 1GA, UK
² Department of Chemistry and Biochemistry, The University of Texas at Dallas, 75080, USA
*Correspondence to Paul Brear (pdb47@cam.ac.uk) and Marko Hyvönen (mh256@cam.ac.uk)

**Keywords:** CK2α, inhibitor, ATP-competitive, kinase, biophysics

Abstract
CK2α is a ubiquitous, well-studied kinase that is a target for small-molecule inhibition, for treatment of cancers. While many different classes of ATP-competitive inhibitors have been described for CK2α, they tend to suffer from significant off-target activity and new approaches are needed. A series of inhibitors of CK2α has recently been described as allosteric, acting at a previously unidentified binding site. Given the similarity of these inhibitors to known ATP-competitive inhibitors, we have investigated these further. In our thorough structural and biophysical analyses, we have found no evidence that these inhibitors bind to the proposed allosteric site. Rather, we report crystal structures, competitive ITC and NMR, HDX mass spectrometry and chemoinformatic analyses that all point to these compounds binding in the ATP pocket. Comparison of our results and experimental approach with the data presented in the original report suggest the primary reason for the disparity is non-specific inhibition by aggregation.

Introduction
Potent inhibitors of kinases have existed for some time and the majority of these bind in the active site of the kinase, competing with the co-factor ATP¹². Because of the significant conservation of the ATP binding site, kinase inhibitors often suffer from poor selectivity³–⁷. This poor selectivity is a disadvantage for several reasons. Firstly, promiscuous inhibitors tend to manifest off-target toxicity if used therapeutically and, to combat this, end up being used at suboptimal concentrations⁸⁹. Secondly, they are poor chemical tools as they cannot provide unambiguous answers to target validation questions. In order to develop more selective inhibitors of kinases, a promising strategy is to target sites outside the highly conserved ATP pockets. This leads to increased selectivity as these sites are structurally not constrained by the shared property of binding to ATP and therefore tend to be less conserved among different kinases¹⁰–¹³. When developing inhibitors that bind outside the primary ATP site, it is vital that the new sites are thoroughly validated biochemically, biophysically, and structurally to confirm the inhibitor binding mode unambiguously. Often inhibitors are identified as being allosteric if they are shown to be non-competitive in biochemical assays. However, this data can be
misinterpreted, as non-specific inhibitors also often show similar non-competitive behaviour. This problem can be further compounded by moderate/weak affinity of the hits being investigated, as they must be used at higher concentrations, which makes secondary binding more likely. Therefore, apparent allosteric behaviour must be very carefully validated to ensure that vital time and resources are not wasted on optimising these inhibitors further.

CK2α has a long history as a target for drug discovery, due to its ubiquitous role in multiple diseases such as cancer and fibrosis. CK2α is an unusual kinase in that it is constitutively active and does not require phosphorylation for activation. It is also a highly promiscuous kinase with hundreds of recorded cellular substrates. Inhibition of CK2α has been achieved mostly through an ATP-competitive mechanism and a number of high affinity inhibitors of distinctly different chemotypes are known and characterized structurally. Despite significant effort, most inhibitors have poor selectivity, although claims are often made to the contrary. For example, the clinical trial candidate CX4945 inhibits at least ten other kinases with nanomolar affinity (see Table S1 in Brear et al.).

We have recently demonstrated a new approach to CK2α inhibition. By using a cryptic so-called αD pocket just below the active site as an anchoring point, we have been able to develop highly selective inhibitors of CK2α. The most potent of these is the most specific CK2α inhibitor reported to date, CAM4066. The fragment-based development of these inhibitors was supported by a significant amount of structural data to identify the optimal αD site anchor moiety and subsequent growth of the fragment to the ATP site. In addition to active site targeting, another approach is the inhibition of CK2α binding to its scaffolding partner CK2β. This method has the potential to generate specific inhibitors that affect only a subset of CK2 substrates and has led to some initial encouraging success that requires further development.

Recently, a series of CK2α inhibitors (Series A, SI Fig. 1), proposed to bind in a novel site (Site A, SI Fig. 2) just outside of the highly conserved ATP site, has been published. The authors used enzymatic assays, native mass spectrometry and competitive ligand-based NMR studies to characterize the mode of inhibition. They interpret their data as supporting an allosteric mechanism. While their data were presented as supporting non-ATP-competitive inhibition, there were several reasons why we speculated that the mechanism of inhibition by these molecules was not necessarily allosteric. In particular: the enzyme assays showed mixed mode inhibition; ATP analogues were replaced in NMR studies; high affinity ligands showed unexpected signal in ligand-based NMR experiments; and, mutations close to the ATP site affected inhibitory potential. Our motivation to investigate these inhibitors in more detail arose from the somewhat conflicting data presented, as well as the
chemical properties of these new inhibitors, which are not unlike known ATP competitive kinase inhibitors.

We have used several orthogonal biophysical and structural methods for the characterization of the binding mode of these new CK2α inhibitors and we find no evidence for interaction with the proposed allosteric site (Site A). We show, using crystal structures, competitive Isothermal titration calorimetry (ITC) and NMR, hydrogen-deuterium exchange mass spectrometry (HDX) and computational analyses of the ligand structures, that these molecules are clearly binding to the ATP site of the kinase. It thus seems likely that most of the inhibitory effect, if not all, comes from traditional type I inhibition of the kinase through orthosteric competition with ATP.

Results
Chemoinformatic analysis
The structures of the compounds presented in the original papers and their comparison to other validated kinase and CK2α inhibitors raised concerns over the claim that these compounds had a novel binding mode. The conserved scaffold of the compounds shares significant similarity to many established and well validated kinase inhibitors that bind in the ATP site (Fig. 1)\textsuperscript{28,37,46–55,38,56–59,39–45}. This scaffold consists of a central thiazole ring that is connected directly to a phenyl group on one side and to a benzoic acid moiety through an amine on the other side (Fig. 1A). We have used two representative compounds from Series A for further investigation, which we denote 1 and 2 (these were referred to as 6 and 27 respectively in the original papers\textsuperscript{35,36}). Compound 1 was chosen as much of the validation of Series A and Site A relied upon this compound\textsuperscript{35}. Compound 2 was chosen as it was the highest affinity inhibitor reported in Series A\textsuperscript{36}. A search of the Protein Data Bank (PDB, http://www.pdb.org) for ligands that bind to serine/threonine kinases (EC=2.7.11.1) and contain a benzoic acid group, which is a critical part of the inhibitors in question (Fig. 1B), identified 49 molecules. Of these, 46 bind in the ATP site and the remaining three bind in interface sites that do not exist in CK2α (SI Table 1). Narrowing the search by inclusion of the amine that links the benzoic acid to the thiazole ring identified 15 ligands (SI Table 2). Of these ligands, all 15 bind in the ATP site. When a similar search was performed on CK2α, 14 ligands were identified, all of which bound in the ATP site (SI Table 3). Four representative structures of these compounds are shown in Fig. 1C. All of these inhibitors display the same binding mode where the binding to CK2α is dominated by the interaction between the carboxylic acid and the ζ-amino group of Lys68 (Fig. 1C). Further refining the search to include only compounds that also bind in the hinge region, similar in structure to the thiazole-amine group of 1, identified eight inhibitors (SI Table 4). These compounds, including CX4945, an extensively validated ATP-competitive CK2α inhibitor currently in clinical trials, interact with the hinge and Lys68\textsuperscript{46}. Overlaying the 2D structure of 1 with CX4945 shows that 1 could replicate the binding mode
of CX4945 (SI Fig. 3) and bind in the ATP site with its carboxylic acid interacting with Lys68 and amine hydrogen bonding with the hinge (Fig. 1C). Indeed, modelling of 1 into the ATP site of CK2α replicates the crystallographic binding mode of CX4945 (SI Fig. 3). The carboxylic acid is interacting with Lys68 and an ATP site water as predicted and the hydrophobic core of the molecule is sandwiched between Met163 and Val66. The amine is not predicted to interact with the hinge. However, the phenyl group is predicted to bind in a similar position to the phenyl chloride of CX4945.

Further evidence in support of our hypothesis that 1 and 2 may bind in the ATP site comes from the analysis of a fragment-based X-ray crystallographic screen against CK2α27 previously performed at our lab. One purpose of fragment-based X-ray crystallographic screening is to use the fragments to probe the target protein and identify binding hotspots60. From the screen of 354 fragments, 21 were identified as binding to CK2α. All 21 bound in the ATP site and interacted with Lys6827. A total of 19 of the CK2α-binding fragments were structurally related to 1 and contained a carboxylic acid moiety that mediated this interaction (SI Fig. 4). The preference of CK2α for binding ligands with a carboxylic acid in this position is further supported by the serendipitous binding of acetate. Acetate ions can be utilised as a very low molecular weight probe to identify binding sites for carboxylic acids60. In CK2α structures deposited to the PDB, at least nine binding sites for acetate can be identified (SI Fig. 5). Among these sites are the Lys68 hotspot in the ATP site, the CK2β interface and the substrate-binding pocket. Notably, none of the nine sites overlaps with Site A, the proposed binding site of 1 and 2. When such a conserved binding mode is observed for an ATP site-binding pharmacophore, it seems unlikely that inhibitors with significant structural similarity would not also bind to the ATP site. These observations led us to initiate additional studies to investigate the binding mode of these inhibitors using two compounds (1 and 2, Fig. 1A) from the original publications.
Figure 1. Conserved pharmacophore binding in the ATP site of protein kinases (A) The structures of 1 and 2, denoted 6 and 27 respectively in the original studies. (B) A summary of the analysis of the established ligands that bind to Ser/Thr protein kinases (EC number 2.7.11.1) or CK2α in the ATP site. (C) The structures of the four CK2α inhibitors and their protein-bound crystal structures which contain a benzoic acid group that binds in the ATP site.\cite{6,48,56,59}
X-ray crystallographic analysis

The original publication contained no structural data on the inhibitor binding mode. In fact, the authors state that 'cocrystallisation attempts aimed at elucidating the non-ATP competitive binding mode of 2-aminothiazole derivatives with CK2α were not successful'\textsuperscript{35,36}. We sought to rectify this and have determined structures of both 1 and 2 in complex with CK2α (Fig. 2, SI Table 5). These structures were obtained by soaking the compounds into CK2α crystals at 10 mM for 16 hours. The crystals diffracted at less than 2 Å resolution and unambiguous positive difference electron density corresponding to both 1 and 2 can be clearly seen in the ATP site before the ligands were included (Fig. 2A and 2B, SI Fig. 6). As we predicted, the carboxylic acid groups of both compounds interact directly with Lys68 like the carboxylic acid of CX4945 (Fig. 2C and 2D). The remainder of the molecules however, each have their own binding nuances. The amine of 1 interacts very weakly with the hinge region backbone nitrogen of Val116 via a bridging water molecule whereas the OH group of 2 interacts with His160, via a bridging water, which pulls the amine of 2 further away from the hinge region and prevents interactions with the hinge (Fig. 2C and 2D). The 5-membered rings from 1 and 2 stack on top of Met163 which leads to the phenyl and naphthyl ring of 1 and 2 stacking on top of His160 at the entrance to the ATP site. The observed binding mode agrees well with the binding mode of established ATP site inhibitors of CK2α, such as CX4945. No density corresponding to either 1 or 2 was observed in the proposed allosteric binding site (Site A, SI Fig. 6), despite the high concentration of the ligand (>1,000-fold excess over $K_d$) used for soaking. The binding mode of 1 in the ATP site of CK2α observed in the crystal structure agrees well with that predicted by the modelling study (SI Fig. 3).
Figure 2. Crystal structures of compounds 1 and 2 bound to CK2α (A) The Fo-Fc map (green, displayed at 3σ) in the ATP site and Site A before the ligand has been placed. Refined structure of 2 (green) has been superimposed on the ATP site (PDB: 6YPJ) and the predicted binding mode of 2 (transparent purple) is shown on Site A. (B) The Fo-Fc map (green, contoured at 3σ) for the ATP site and Site A before the ligand has been placed. Refined structure of 1 (blue) has been superimposed on the ATP site (PDB: 6YPG) and the predicted binding mode of 2 (transparent purple, from supplementary material of reference 34) is shown on Site A. (C) 2 (green) bound to CK2α, (PDB: 6YPJ) with key interacting residues and H-bonding highlighted. (D) 1 bound to CK2α (PDB: 6YPG), with key interacting residues and H-bonding highlighted. The structure of CX4945 (PDB:3PE1, yellow) has been superimposed on the structure of 2 and 1 in panels C and D. Electron densities of the final, refined structures are shown in SI Fig. 7.

To reduce the impact that different crystal forms and packing may have on the accessibility of binding sites and bias in the resulting complex structure, we have soaked the inhibitors into two different crystal forms of CK2α. One is obtained using an engineered form of CK2α (referred to as CK2α KKK/AAA) in which lysines 74-76 are mutated to alanines to facilitate crystallization. These lysines are in the vicinity of Site A and could affect binding of ligands to this site. However, these lysines do not affect the ATP site and we show later that both 1 and 2 bind to CK2α KKK/AAA and to wild type CK2α with the same affinity (see ITC measurements). The second crystal form (CK2α SF) has no mutations in the lysine-rich loop but has a substrate peptide fused to the N-terminus of the protein, aimed at facilitating crystallization. In both crystal forms, we observe electron density for 1 and 2 only in the ATP site, and not in the proposed allosteric site (Site A, SI Fig. 6).

The two crystal forms of CK2α have been extensively used for the determination of CK2α:ligand complexes (66 structures currently in the PDB). More than eight different ligand binding sites can be
accessed by soaking into these crystal forms (SI Fig. 8), including the proposed allosteric site for 1 and 2, Site A. We have observed both ADP and GDP bound to Site A in crystal structures of wild type CK2α (PDB:6YPN) and in the KKK/AAA mutant (PDB:6YPK) (Fig. 3). The adenosine and guanosine rings sit in the hydrophobic pocket identified in the previous modelling work\textsuperscript{35} and the diphosphate interacts with Arg80 that has been identified as an acetate-binding hotspot (SI Fig. 5). This confirms that Site A is accessible to ligands in these crystal forms and soaking with 1 or 2 would be expected to result in clear electron density in the site, should they bind there. Indeed, ADP can be seen simultaneously in both the ATP site and in Site A (Fig. 3D, PDB: 6YPN). The binding conformation of ADP in Site A is very different to the conformation in the ATP site (SI Fig. 9). These observations demonstrate that binding in Site A does not in itself prevent binding in the ATP site and that if 1 or 2 did bind in both sites, we would expect to see density corresponding to it. Furthermore, 1 and 2 would be expected to bind predominantly in the highest affinity site and this site would dominate the inhibition. These crystal structures prompted us to perform further experiments to unambiguously establish the binding site in solution and verify the likely mode of inhibition for these compounds.
Figure 3. Crystal structures of ADP and GDP binding to CK2α Site A. (A) GDP binding in Site A. The Fo-Fc map (contoured at 3σ) centered on the proposed allosteric site. No density was observed in the ATP site for GDP therefore the density in the ATP site was not shown for clarity. The CK2α KKK/AAA construct was used for this structure (PDB: 6YPK). B) The structure of GDP bound to CK2α, with the extensive H-bonding network shown by dotted lines (PDB: 6YPK). C) The proposed binding mode of 1 (PDB: 6YPJ) superimposed on the structure of GDP bound to CK2α (6PYK). D) The structure of ADP bound simultaneously in the ATP site and in Site A of WT CK2α (PDB: 6YPN).

Hydrogen-deuterium exchange mass spectrometry

Hydrogen-deuterium exchange (HDX) allows the localization of interaction sites and conformational changes in proteins. We have analysed the binding of four different classes of ligands to CK2α (SI Fig. 10 and 11) using HDX. These ligands were: ADP with Mg²⁺ and adenosine-5’-(β,γ-imido)triphosphate
(AMPPNP)-Mg^{2+}, both of which bind across the entire ATP site interacting with the hinge region and Lys68 (SI Fig. 10C and 10D); 3, which is a fragment that binds only on the right hand side of the ATP site, interacting with Lys68^{26} (SI Fig. 10E); 4, which is an αD site binder (SI Fig. 10G)^{26} and 6 which links the αD site and the ATP site (SI Fig. 10F)^{27}. We recovered 192 shared peptides from each of these HDX experiments that redundantly span almost the entire sequence of CK2α (SI Fig. 12). This allows for unbiased detection of ligand interaction fingerprints across the whole of the protein. Binding can be identified by observing changes in deuterium uptake between CK2α alone and a concurrently run ligand-bound sample (SI Table 6). We show all changes greater than ±0.25 Da that have a p-value less than 0.01 in a Welch’s t-test (Fig. 4 and SI Fig. 13). Extended data for all peptides and timepoints are shown in the Supplementary Information (SI Fig. 13 and a separate spreadsheet). For all the ligands, a decrease in deuterium uptake is observed at late timepoints particularly for peptides containing residues 54-111 (sample peptides shown in SI Fig. 13). Since this change occurs with all the ligands, it appears to result from general stabilization of the region termed the ‘N-lobe’, on ligand binding, rather than an effect attributable to a specific site.

Addition of ADP and AMPPNP-Mg^{2+} causes clear changes in deuterium uptake in the ATP site (Fig. 4). The largest change is a decrease in deuterium uptake for peptides covering residues 39-55 and 121-131. Residues 39-55 correspond to the P-loop that interacts directly with the phosphates of ADP, while residues 121-131 are the αD loop that does not interact directly with ADP or AMPPNP-Mg^{2+}. Protection occurs in the latter due to the interaction of the adenosine ring with the hinge region that directly links to and stabilizes the αD loop. These observations are similar to those previously observed when binding of ATP analogues to other kinases has been studied^{61,62}. The observed decreases in uptake are larger for AMPPNP-Mg^{2+} than ADP-Mg^{2+} because AMPPNP binds with a higher affinity than ADP. To further probe interactions in a more localized area of the ATP site, we used 3 that binds only at the right hand side of the ATP site (SI Fig. 10 E), coordinated to catalytic Lys68 (PDB: 5CSP)^{36} (Fig. 4). With 3, we see a decrease in deuterium uptake in the P-loop and no change in the αD loop. The pattern of protection within the P-loop is also distinct, indicative of different binding contacts being used by 3 compared to the ATP analogues. We can thus distinguish binding in the entire ATP site, including interaction with the hinge, from binding on just one side where Lys68 is located.

To ensure we could detect different binding sites with our HDX experiments, we used another well characterized ligand (4) that binds to the αD pocket only (SI Fig. 10G)^{26}. This compound shows a markedly different HDX fingerprint compared to the ATP analogues and 3 (Fig. 4). The largest differences are seen in the αD loop, with binding of 4 resulting in increased uptake compared to unbound CK2α. Increased uptake in the αD loop is opposite to the decrease seen upon ADP and AMPPNP-Mg^{2+} binding. This indicates that the binding of 4 in the αD pocket puts the loop in a more
flexible or open conformation amenable to exchange, as is seen in the crystal structure of CK2α in complex with 4 (PDB: 5ORJ). Finally, we characterised the HDX fingerprint for 6 that occupies both the αD pocket and the ATP site. We observed a similar increase in uptake in the αD loop as with 4 (Fig. 4). Compounds 4 and 6 also cause decreased uptake in the P-loop, with 6 showing more protection than 4, similar to the protection we observed with 3.

All these fingerprints are consistent with the known binding modalities of these ligands to CK2α and can be used to identify the binding modes of new compounds. We performed the same experiments to characterise the binding of compound 1 to CK2α (Fig. 4). Addition of 1 to CK2α causes decreased uptake in the P-loop with a pattern and magnitude of change similar to the addition of AMPPNP-Mg²⁺. However, no changes are observed in the αD pocket, as was observed with the addition of 3. These results are consistent with the binding mode seen in the crystal structures where 1 makes significant hydrogen bonding just to Lys68 with only weak water-mediated interaction to the hinge region. These data do not allow us to unequivocally eliminate binding at the proposed allosteric site. However, if the allosteric binding mode were correct, two clear differences would be expected. Firstly, a change in the uptake by residues 67-88, part of Site A, would be expected due to the main interaction being with the acidic group of 1 (SI Fig. 10). We do not observe changes in uptake in this region despite it being redundantly covered by eight peptides (SI Fig. 13). Secondly, no change in the uptake by the P-loop would be expected since the modelling does not predict significant interactions with that part of the protein. Interactions with the P-loop, such as that seen for 1, are one of the major features of ATP site binders, consistent with our structural characterization of 1 and CAM4066.
Figure 4. HDX analysis of ligand binding to CK2α. Heatmaps show the change in deuterium uptake upon binding of different ligands to apo CK2α. Each panel, with the ligand listed on the top, shows the heatmaps for 10 s and 100 s time points for peptides corresponding to the P-loop and αD-loop, in comparison to unliganded CK2α. All depicted changes are greater than ±0.25 Da and have a p-value less than 0.01 in a Welch’s t-test. The exact start and end residues for each peptide are shown on the left of the panels. Next to each heatmap are two graphs showing deuterium uptake for one peptide from each P-loop (top) and αD loop (bottom) for apo CK2α (black) and CK2α with ligand (grey). The Y-axis range is 80% of max D-uptake assuming the N-terminal residue undergoes 100% back-exchange.
Data have not been corrected for back-exchange. Error bars are ±2σ derived from three technical replicates. A structural diagram in between the uptake plots has the P-loop and αD loop coloured purple and green, respectively, and the bound ligand in surface rendering in yellow. Full HDX data are shown in SI Fig. 13 and available in the supplementary spreadsheet.

**Ligand-observed NMR analyses**

To further explore the binding of the new compounds to CK2α, we have used various ligand-based NMR methods, similar to the experiments in the original characterization of these molecules. We were particularly concerned about the evidence for the simultaneous binding of **CX4945** and 1 due to the presence of strong signals for **CX4945** in the STD NMR spectra. In order to observe a signal in an saturation transfer difference (STD) spectrum, the saturation of protein resonances has to be transferred to the bound ligand, which then dissociates to join the pool of bulk ligand in solution before it can be detected. This means that the size of the signal for a given ligand is proportional to the amount of the ligand that binds to and is released by the protein during the saturation step of the STD experiment. Therefore, ligands with relatively fast association (k\(_{\text{on}}\)) and dissociation (k\(_{\text{off}}\)) rates give the largest signal\(^{63,64}\).

We have determined the off rate of **CX4945**, that binds to CK2α with low nanomolar to picomolar affinity\(^{46}\), from CK2α to be 0.0039 s\(^{-1}\) by SPR (SI Fig. 14). When ligands have k\(_{\text{off}}\) rates <0.1 s\(^{-1}\), the saturation cannot be transferred effectively to bulk ligand in solution resulting in no observable STD effect\(^{64}\). The experimentally determined off-rate for **CX4945** means that less than 1% of **CX4945** would have dissociated after the 2-second saturation time used in the NMR pulse sequence. This would mean that the concentration of saturated ligand in solution would be less than 0.1 μM. This difference would not be observable by NMR.

To resolve these issues, we have repeated the STD NMR studies reported in the original publication\(^{35}\) and used two additional ligand-based NMR experiments, Carr-Purcell-Meiboom-Gill (CPMG) and Water-LOGSY\(^{63}\), for further validation. Our STD experiment shows that 1 binds to CK2α, as revealed by detectable transferred \(^1\)H intensities (Fig. 5A(i)). However, when the same experiment is run in the presence of **CX4945**, a significant reduction of the STD signals from 1 are observed, indicating direct competition between the two ligands (Fig. 5A(ii)). In the original paper, strong **CX4945**-derived signals are also clearly visible. The signals - which are comparable in magnitude to the much weaker binding ligand 1 - could instead arise from non-specific binding of **CX4945** to lower affinity sites on CK2α (as was shown by native mass spectrometry\(^{35}\)), precipitation of **CX4945** leading to false positive signals in the experiment or binding of **CX4945** to soluble aggregates of 1.
Given the disparity in the STD NMR experiments between our results and those in the original report, we have also used CPMG and Water-LOGSY experiments. These exploit different NMR phenomena (relaxation rates, transfer of saturation) to observe the binding of the ligand to the protein\textsuperscript{63}. Using three different methods to observe the binding of the ligand reduces the chance of experimental anomalies interfering with the outcome of the experiment. The Water-LOGSY experiment shows clear binding of 1 to CK2\textalpha{} and complete inhibition of binding in the presence of \textbf{CX4945} (Fig. 5B). Similarly, in the CPMG experiment, an increase in the signal from the aromatic protons of 1 was observed in the presence of \textbf{CX4945} indicating a reduction in binding of 1 to CK2\textalpha{} (Fig. 5C). In conclusion, all three ligand-observe NMR experiments show binding of 1 to CK2\textalpha{} and in all three experiments binding is abolished by \textbf{CX4945}, suggesting direct competition between these two ligands as they bind to the same site.
Figure 5. Ligand-based NMR experiments to detect binding of 1 to CK2α and competition by CX4945. All panels are overlays of spectra recorded in the presence (black) and absence (green) of CK2α. (A) Saturation Transfer Difference (STD) $^1$H spectra of 1 (i) in the absence of CX4945, and (ii) in the presence of 50 μM CX4945. (B) WaterLOGSY spectra of 1 (i) in the absence of CX4945, and (ii) in the presence of 50 μM CX4945 (ii). (C) Relaxation-edited (CPMG) $^1$H spectra of 1 (i) in the absence of CX4945, and (ii) in the presence of 50 μM CX4945 (ii). (NB: the signals at ca. 7.7 ppm are in fact two overlapping doublets and therefore small changes to either chemical shift results in an apparent change to the multiplet structure.)
To further confirm the validity of our results, we repeated the CPMG NMR study using the lower affinity ligands 3, 5 and CAM4066 as competitors (SI Fig. 15). 3 is a small, low affinity ligand in the ATP site and as such it is less likely to have an allosteric effect that might prevent binding at the adjacent Site A – an argument that could explain inhibition of binding to 1 in the presence of CX4945. Therefore, the reduction in the binding of 1 in the presence of 3 (SI Fig. 14B) is a more convincing proof that 1 is binding in the ATP site. CAM4066 links the ATP site and the αD site and, as predicted, reduces the binding of 1 to CK2α (SI Fig. 15C). To rule out binding of 1 to the αD site, we have also used 5, an optimized fragment with high affinity for the αD site. This should not prevent binding in the ATP site. Indeed, ligands similar to 5 have been shown to bind simultaneously with ATP site fragments by X-ray crystallography (PDB:6EHK)26. As predicted, and unlike with CX4945, 3 and CAM4066, the αD site ligand 5 has no effect on the binding of 1 to CK2α (SI Fig. 15D). Similar CPMG and STD NMR experiments were performed with the CK2α KKK/AAA mutant with the same outcomes (SI Fig. 16 and 17). These results further demonstrate that 1 binds in the ATP site.

Isothermal titration calorimetry
As a final confirmation, we have used isothermal titration calorimetry to determine affinities of 1 and 2 to CK2α and to evaluate the effect of competing ligands and CK2α mutations. This process has been successfully used to verify the binding site of novel CK2 inhibitors that bind in the αD site26. The affinities of compounds 1 and 2 for CK2α were determined to be 22 μM and 2.2 μM respectively (Fig. 6A and C). When the same experiments were performed in the presence of saturating concentrations of CX4945, no binding of either compound to CK2α was observed (Fig. 6B and 6D). This observation is in line with all our previous data and suggests that compounds 1 and 2 bind in the same site as CX4945.

We also performed the experiment with 2 in the presence of CAM4066, which competes with ATP, but uses only a small benzoic acid moiety to interact with the ATP site, deriving most of its binding energy from the αD pocket. CAM4066 abolished the binding of 2 to CK2α (SI Fig. 18). In the modelling of the binding of 1 and 2 reported previously35,36, Lys74 was proposed to form a vital part of the binding site and to interact with the carboxylic acid of 1 and 2. Therefore, mutations to this residue should reduce the affinity of 1 and 2 for CK2α. To test this, we repeated the ITC experiments using the construct CK2α KKK/AAA, in which three adjacent lysine residues (Lys74-76) have been mutated to alanine. These mutations had a negligible effect on the affinity of 1 or 2 for CK2α as we determined near identical affinities of 27 μM and 3.3 μM, respectively (SI Fig. 18). The binding of 1 and 2 to CK2α KKK/AAA was also fully inhibited by the presence of CX4945 and CAM4066 respectively (SI Fig. 18). These results are further evidence that the ATP site, not the proposed allosteric site, is the primary binding site for 1 and 2.
**Figure 6. ITC measurement of binding of 1 and 2 to CK2α.** (A) Titration of 1 into 10 μM CK2α. (B) Titration of 2 into 10 μM CK2α in the presence of CX4945. (C) Titration of 2 into 20 μM CK2α. (D) Titration of 2 into 20 μM CK2α in the presence of 100 μM CX4945.

**Discussion**

We present compelling and consistent evidence that 1 and 2 bind in the ATP site of CK2α, not in Site A, as was previously proposed.\(^{35,36}\) We see no evidence for binding in Site A. This is as expected, based on our analysis of the structural data of similar kinase inhibitors. In the original study of Series A,\(^{36}\) five main pieces of evidence were used to verify a novel binding mode. This evidence included: competitive native mass spectroscopy studies, ligand-based NMR, thermal shift analysis and biochemical assays.

While we were preparing this manuscript for publication, the Niefind group published their analysis of the same series of compounds, which also showed binding to the ATP site by X-ray crystallography.\(^{65}\) While they have not done other biophysical analyses on the binding, they do show by enzymatic assay that these inhibitors are ATP competitive, consistent with our data.

One of the cornerstones of the theory of allosteric inhibition was the observation that that CX4945 binds CK2α simultaneously with 1, and that 1 competes with inositol hexaphosphate (IP₆, phytic acid), shown by mass spectrometry. The data presented indicate that at least three molecules of 1 bind to CK2α. In the presence of CX4945 this is reduced to two binding sites, but no data were presented to confirm the identity of these sites in which 1 was binding when it bound simultaneously with CX4945 (for example mutational studies combined with native mass spectrometry). Some of these sites are expected to be non-specific, low affinity sites and the mass spectrometric data does not confirm which is the primary binding site of 1. The first evidence presented for the location of the binding site of these compounds is the observation that in the presence of 1 the complex between IP₆ and CK2α is not observed in the native mass spectra. This led to the theory that 1 binds in the proposed site as a crystal structure was published by Lee et al (PDB:3W8L)\(^{37}\) of IP₆ apparently binding to CK2α in this site.
However, the binding site of IP₆ presented in that crystal structure is more controversial than is discussed in the publication. In this structure, IP₆ is found at a crystal contact where half of the site is composed of residues from a symmetry related CK2α molecule in the crystal lattice (SI Fig. 19). Therefore, IP₆ could in theory bind to both these sites. From this crystal structure the binding site could be located around Lys74 or His276. However, as the site is only fully formed in a crystal lattice (SI Fig. 19), IP₆ may not in actual fact bind to CK2α at either of these sites in solution. IP₆ is also seen to bind weakly at a site centered on His236 in several other crystal structures (PDB: 5OQU, 5ORJ, 5MPJ, 5CVH)⁶⁶. The native mass spectrometric competition studies with CX4945 or IP₆ do not therefore unambiguously confirm the binding mode of 1.

Another cornerstone of the allosteric site binding was the observation of the binding by ligand-based NMR and simultaneous detection of CX4945 binding, interpreted as lack of competition. As discussed before, a high affinity ligand like CX4945, with slow dissociation rate should not yield detectable signal in these experiments. The binding of CX4945 observed in those STD experiments is however more likely to be due to an artefact in the experimental setup. This could be through a number of mechanisms including aggregation of the ligand or binding to non-specific lower affinity binding sites outside the ATP site.⁶⁶ These lower affinity sites would have higher on/off rates than the ATP site and therefore show a larger signal in the STD spectrum. Non-specific and low affinity sites can have a confusing influence on competition studies. This is a particular problem with the NMR studies as ligand-observed NMR is optimized for studying low affinity sites and requires ligands with rapid on/off rates in order for the signal (and therefore binding) to be observed. Therefore, blocking the primary (high affinity) site may not remove the binding signal arising from other sites. The competition studies in the original papers were done at 500 μM ligand concentration, well beyond the observed affinities or Kᵢs of 1 and 2 and in a range where non-specific binding would be more likely to occur.³⁵,³⁶ The binding signals observed for CX4945 in the STD spectra cast some doubt on the conclusions drawn about the simultaneous binding of 1 and CX4945. This is because the signals observed for CX4945 cannot be due to binding in the ATP site as the dissociation rate for this site is too slow. The major difference between our experimental system and the system reported in the literature is the DMSO content. In brief, in the procedure reported here stocks of 1 and 2 were made in d₆-DMSO and diluted 20-fold in the buffer. This was done to ensure the ligands were fully soluble in the experiment and led to a final d₆-DMSO content of 5% (which was also used to provide the field-frequency lock signal). When we first attempted to run the experiment without DMSO, we observed precipitation of the ligands. As aggregation can lead to false positives in STD experiments it was decided to use DMSO in all our experiments.⁶⁶ However, in the original experiment D₂O was used for the lock and no DMSO was used to solubilise the ligand. The difference in the solvent conditions may mean that 1 and 2 were
closer to the limits of their solubility in the original experiments and thus more likely to form aggregates. Aggregates are common causes of false positives in biochemical assays and are often missed as they may not be easily visible to the naked eye. These aggregates may lead to the signals observed in the STD for CX4945 and 1.

The biochemical assays performed previously showed that the compounds exhibited mixed mode non-competitive behavior and led to the understandable conclusion that they do not bind in the ATP site. However, this is contradictory to the binding mode we see in the crystal structures. A proposed explanation for the non-competitive inhibition observed in the assay comes from the thermal stability data also presented in the original paper. The data show that the thermal stability of the proteins decreases in the presence of 1 and 2. This thermal de-stabilization could be due to a number of different mechanisms caused by specific and non-specific effects. These mechanisms include; breaking up of protein complexes by specific binding to the protein; binding to the unfolded or partially unfolded protein; the formation of aggregates that bind the protein and unfold it and specific binding that traps the protein in a less stable conformation.

It is impossible that the first mechanism is operating in this example as CK2α is present as a monomer. The other mechanisms however may explain the conflicting results presented. All proteins exist in an equilibrium between the native and unfolded form of the protein. For the majority of ligands, the ligand binds to the native form of the protein which stabilizes the folded form leading to a positive thermal shift. If the ligand binds to the unfolded or partially unfolded form of the protein and stabilize the unfolded form this could lead to a destabilization of the protein. This would in turn lead to a decrease in the $T_m$. Binding to the unfolded protein can occur by two mechanisms, the first mechanism is by the formation of colloidal aggregates of the small molecules, onto which the protein would then be sequestered. The aggregates are hydrophobic and preferentially bind the unfolded or partially unfolded form of the protein where the hydrophobic core is exposed. Therefore, a negative shift in the $T_m$ is observed. This is one of the most common forms of non-specific inhibition that plague biochemical assays and would be seen in assays as noncompetitive inhibition. The second mechanism occurs when the ligand monomers bind to a partially unfolded or fully unfolded form of the protein stabilizing that form. It may be possible for the binding to the partially unfolded form to be a specific form of inhibition.

Modelling experiments proposed Site A as the binding site for 1 and 2 (SI Fig. 1). We have accumulated extensive evidence that these compounds bind in the ATP site. However, Site A as a general ligand binding site has been verified by our structures of ADP and GDP bound in this location. This suggests that Site A could be utilized for the development of novel inhibitors of CK2α and it also
demonstrates that this site is accessible for ligand binding in our crystal forms. If this site were to be successfully targeted by inhibitors, it would require extensive structural and biochemical validation to confirm that compounds are really binding in the new site and not in one of the several known small molecule binding sites in CK2α.

Conclusions
We believe that the data we have presented here leads to the conclusion that the compounds represented by 1 and 2 bind to the ATP site and will inhibit CK2α via an orthosteric, ATP-competitive mechanism. This diverse set of data pointing to 1 and 2 binding at the ATP site includes multiple crystal structures, competitive ITC studies, competitive NMR studies, HDX mass spectrometry and computational analyses of the ligand structures. We also believe that the evidence presented previously in support of an allosteric mode of inhibition for these compounds may have been distorted by non-specific binding of the compounds to CK2α, which in turn affected the interpretation of competition data with CX4945.

Experimental Section
Modelling
Modelling of binding of 1 and 2 to CK2α was done using the GOLD software package (CCDC) using the structure from PDB entry 3PE1 as the target. The site of interest was defined as a 20 Angstrom sphere around the ligand CX4945, which includes both the ATP site and the proposed allosteric site. Ligand files were generated with grade (Global Phasing). The conserved water interacting with Lys68 was included but no other waters. The ChemPLP scoring system (CCDC) was used to rank the results. Coordinates of the modelling results are included as supplementary files.

Expression and purification
Expression and purification of CK2α (wild type, KKK/AAA and SF) was done as described before56. For the surface plasmon resonance experiments a new construct carrying two His-tags, one at the N-terminus and one at the C-terminus was created. CK2α was amplified by PCR using primers “AGCTAGAATTCGGAGAAGGCGAGGGTAGTGCAGGCCCCCGTGCAAGCAGGGCCAGAGT” and “ATACTCAAAGCTTAGTTAGGTAGTGGTGAGGGAACCCTGGAAGTACAGGTGGTCTTTCCTCCACACACGATG
GAAATAGGGGTG”. The resulting fragment was digested with EcoRI and HindIII restriction enzymes and ligated into a pHAT4 vector digested with the same enzymes. The resulting fragment was digested with the protein that the resulting construct (pHAT4_CK2-TEV-L-His) encodes for is shown below with His-tags (bold) and two TEV cleavage sites (underlined).

Expression of this protein was done as with other CK2α construct, using the BL21(DE3) strain of E. coli. Single colonies of the cells were grown in 6 l 2xTY with 100 μg/mL ampicillin at 37 °C until OD600 of 0.6 after which 0.4 mM IPTG was used to induce expression. After three hours of culturing at 37 °C the
cell were harvested by centrifugation at 4,000 g for 20 minutes. The cells were resuspended in 20 mM Tris pH 8.0, 500 mM NaCl and lysed using an Emulsiflex C5 homogeniser. DNaseI was added to aid with digestion of *E. coli* genomic DNA and protease inhibitor tablets were added to minimise proteolytic degradation. Following centrifugation for 30 min at 15,000 g, the clarified supernatant was loaded onto 4 ml Ni²⁺-NTA agarose column and washed with the lysis buffer, followed by 20 mM Tris pH 8.0, 500 mM NaCl, 20 mM imidazole and the protein eluted with 20 mM Tris pH 8.0, 500 mM NaCl, 300 mM imidazole. Peak fractions, as analysed by SDS-PAGE, were pooled together and dialysed overnight against 4 l of 20 mM Tris pH 8.0, 500 mM NaCl at +4°C. The dialysed sample was diluted to ~65 mM salt concentration in two batches and immediately loaded onto HiTrapQ HP column to minimise aggregation in low ionic strength. CK2α was eluted using a linear salt gradient. Peak fractions of each of the two chromatographic steps were pooled, concentrated to ca. 5 ml in volume and purified with size exclusion chromatography on a Superdex 75 16/60 HiLoad column using 20 mM Tris pH 8.0, 500 mM NaCl buffer. Peak fractions were pooled, concentrated to ca. 10 mg/ml using Vivaspin 20 5000 MWCO centrifugal concentrators and flash frozen in small aliquots.

Source of compounds
The synthesis of Compound 1 and Compound 2 was described previously by Bestgen et al. Purity was determined as >95% by LC-UV. Compound 1 was also purchased from enamine (ID: Z2239047232, no longer available). Purity was determined as >95% by LC/MS. CX4945 was purchased from Cambridge bioscience (HY-50855B-5mg). Purity was determined as >99% by HPLC. ATP was purchased from Sigma Aldrich (Cat: A7699). The purity was determined as >99% by HPLC. GDP was purchased from Sigma Aldrich (G7127). The purity was determined as >96% by HPLC.

X-ray crystallography
Crystallisation, soaking of ligands and structure determination was done as described before. X-ray diffraction data were collected at the Diamond Light Source and data from automated data processing with autoProc were used for the structure determination. Fo-Fc electron density maps for all the ligands, calculated prior to ligand placement in the model, are shown in Supplementary Figure 5 and the final densities, as calculated by PDB AutoDep software, are shown in Supplementary Figure 6. All coordinates have been deposited to the Protein Data Bank under accession numbers 6YPG, 6YPH, 6YPJ, 6YPK, 6YPN. We will release the atomic coordinates and experimental data upon article publication. Data collection and refinement statistics are shown in Supplementary Table 5.

Hydrogen-deuterium exchange mass spectrometry
Stocks of 25 μM CK2α with and without ligand were prepared in 20 mM Tris-HCl, 500 mM NaCl, pH 8.0 at 25°C. In some experiments a small percentage of organic solvent was added to increase solubility. Variations and ligand concentrations are noted in Supplementary Table 6. Stocks were equilibrated at 4°C for 10 minutes and then diluted 1:24 with the same buffer containing H₂O for controls (pH_{read}=8.0 at 25°C) or D₂O (final D₂O of 96% (v/v) with pH_{read}=7.6 at 25°C). After 10 s, 10² s, 10³ s, 10⁴ s, and 10⁵ s, samples were mixed 1:1 with 0.8 % formic acid, 1.6 M guanidinium chloride, pH 2.3 to give a final pH of 2.6 at 0 °C, and flash frozen in liquid nitrogen for storage at –80°C. Samples were thawed and LC-MS performed using a Waters HDX manager and SYNAPT G2-Si Q-Tof. Three technical replicates of each sample were analyzed in a random order. Samples were digested online by *Sus scrofa* Pepsin A (Waters Enzymate BEH) at 15°C and peptides trapped on a C18 pre-column (ACQUITY UPLC Peptide BEH C18 VanGuard Pre-column) for 3 min at 100 μL/min and 1°C. The LC buffer was 0.1% formic acid. Peptides were separated over a C18 column (Waters acquity UPLC BEH) and eluted with a linear 3-40% (v/v) acetonitrile gradient for 7 min at 40 μL/min and 1°C. Raw data of control samples were processed by PLGS (Waters Protein Lynx Global Server 3.0.2) using a database containing *Sus scrofa* Pepsin A and *Homo sapiens* CK2α. In PLGS, the minimum fragment ion matches...
per peptide was 3, and methionine oxidation was allowed. The low and elevated energy thresholds were 250 and 50 counts respectively, and overall intensity threshold was 750 counts. Labelled data were analyzed in DynamX 3.0 with peptide filters of 0.3 products per amino acid and 1 consecutive product. All mass spectrometry data were acquired using positive ion mode in either HDMS or HDMSE modes. HDMSE mode was used to collect both low (6V) and high (ramping 22-44 V) energy data-independent peptide fragmentation data for peptide identification. HDMS mode was used to collect low energy ion data for all deuterated samples. All samples were acquired in resolution mode. Capillary voltage was set to 2.8 kV for the sample sprayer. Desolvation gas was set to 650 L/hour at 175°C. The source temperature was set to 80°C. Cone and nebulizer gas was flowed at 90 L/hour and 6.5 bar, respectively. The sampling cone and source offset were both set to 30 V. Data were acquired at a scan time of 0.4 s with a range of 100-2000 m/z. Mass correction was done using [Glu1]-fibrinopeptide B as a reference mass. To allow access to the HDX data of this study, the HDX data summary table (Supplementary Table 6) and the HDX data table are available online per consensus guidelines.

Ligand-observe NMR experiments

In the control experiment a stock solution of 2.5% DMSO-d6, 7 μM CK2α WT in 0.5mM Tris, 200 mM NaCl, pH 8.0 was prepared. This was split into two samples. Ligand in DMSO-d6 or DMSO-d6 alone was added to each such that one sample contained 5% DMSO-d6 with a total volume of 180 μL, and the second sample contained 5% DMSO-d6 and 500 μM 1. In a typical competition experiment a stock solution of 50-500μM competing ligand, 2.5% DMSO-d6, 7 μM CK2α WT in 0.2mM Tris, 100 mM NaCl, pH 8.0 was prepared. This was again split into two samples. One sample contained 5% DMSO-d6 and 50-500 μM competing ligand with a total volume of 180 μL. The second sample contained 5% DMSO-d6, 500 μM 1 and 50-500 μM competing ligand.

NMR data were recorded at 300 K with a Bruker AVANCE spectrometer operating at 500 MHz and equipped with a room-temperature probe. CPMG-filtered 1D 1H experiments were run with 600 loops of length 1 ms. Water suppression was achieved with low-power presaturation and DPFGSE WATERGATE W5. Saturation transfer difference (STD) experiments were acquired with 2 s of alternating on- and off-resonance saturation (20 ms Gaussian pulse, +0.8 and –5 ppm), a $T_1\rho$ filter to suppress protein signals, and DPFGSE WATERGATE W5. WATER-LOGSY experiments were recorded using a 20 ms Gaussian to invert the water resonance, a mixing time of 1.2 s, a $T_2$ filter to suppress protein signals, and DPFGSE WATERGATE W5.

Isothermal calorimetry

ITC All ITC experiments were performed at 25 °C using a MicroCal iTC200 instrument (GE Healthcare). CK2αWT (20 mg/mL, 20 mM Tris pH 8.0, 500 mM NaCl) was diluted in Tris buffer (200 mM Tris, 300 mM NaCl, 10% DMSO) and concentrated to 20-50 μM. Compounds in 100x stock solutions were diluted into the buffer ensuring that the DMSO concentrations were carefully matched. In a typical experiment CK2αWT (10 μM) was loaded into the sample cell and 0.4-2.0 mM of the ligand was titrated in eighteen 2 μL injections of 2 s duration at 150 s intervals, stirring at 750 rpm. Heats of dilution were determined in identical experiments, but without protein in the cell. The data fitting was performed with a single site binding model using the Origin software package.

Surface Plasmon Resonance

An Ni-NTA sensor chip (GE Healthcare) was activated with 0.5 mM NiCl2, 0.2 μM CK2α with both N- and C-terminal His-tags was immobilised onto the activated Ni-NTA sensor chip in 20 mM Tris pH 8.0, 500 mM NaCl. Binding of CX4945 was determined in 0.01 M HEPES pH 7.4, 0.15 M NaCl, 0.05% v/v TWEEN 20, 0.5% DMSO at five different concentrations (15 nM, 10 nM, 5 nM, 1 nM and 0 nM). The
chip surface was regenerated with an initial wash of 500 mM EDTA followed by 4 M guanidinium chloride, 250 mM EDTA, 50 mM NaOH. The reference channel was activated with 0.5 mM NiCl₂. All data were processed using the manufacturer’s BiAevaluation software.

Ancillary Information
Additional figures are available free of charge via the Internet at http://pubs.acs.org

Structural coordinates in Protein Data Bank:
6YPG (CK2α in complex with 2), 6YPH (CK2α in complex with 2), 6YPJ (CK2α in complex with 1), 6YPK (CK2α in complex with GDP), 6YPN (CK2α in complex with ATP).

Author Information
Corresponding Authors
Correspondence to Paul Brear (pdb47@cam.ac.uk) and Marko Hyvönen (mh256@cam.ac.uk)

Acknowledgements:
We would like to thank Dr Glyn Williams and Teodors Pantelejevs for their critical comments on the manuscript. We are grateful for Diamond Light Source for access to beamlines i04, i04-1 and i24 (proposals: mx9537, mx14043, mx18548). We thank the X-ray Crystallographic and Biophysics Facilities for access to instrumentation. We thank Apollo Therapeutics that funds our CK2α inhibitor development project for allowing the use of the data on CX4945 binding kinetics for this publication. The work in the D’Arcy group was funded by the NIH National Institute of General Medical Sciences (GM133751) and start-up funds to S. D’Arcy.

Abbreviations used
AMPPNP, Adenylyl-imidodiphosphate; CPMG, Carr-Purcell-Meiboom-Gill; HDX, Hydrogen-Deuterium exchange; IP₆, inositol hexaphosphate; ITC, Isothermal titration calorimetry; STD, Saturation transfer difference;

References


Niefind, K.; Bischoff, N.; Golub, A. G.; Bdzhol, V. G.; Balanda, A. O.; Prykhod’ko, A. O.; Yarmoluk, S. M. Structural Hypervariability of the Two Human Protein Kinase CK2 Catalytic


