

Multisite phosphorylation of a CDK inhibitor sets a threshold for the onset of DNA replication

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SCF ubiquitin ligases target phosphorylated substrates for ubiquitin-dependent proteolysis by means of adapter subunits called F-box proteins. The F-box protein Cdc4 captures phosphorylated forms of the cyclin-dependent kinase inhibitor Sic1 for ubiquitination in late G1 phase, an event necessary for the onset of DNA replication. The WD40 repeat domain of Cdc4 binds with high affinity to a consensus phosphopeptide motif (the Cdc4 phospho-degron, CPD), yet Sic1 itself has many sub-optimal CPD motifs that act in concert to mediate Cdc4 binding. The weak CPD sites in Sic1 establish a phosphorylation threshold that delays degradation *in vivo*, and thereby establishes a minimal G1 phase period needed to ensure proper DNA replication. Multisite phosphorylation may be a more general mechanism to set thresholds in regulated protein–protein interactions.

Numerous regulatory proteins are targeted for degradation in a precisely programmed manner through the covalent conjugation of ubiquitin, which is transferred along a cascade of E1, E2 and E3 enzymes to the substrate¹. Reiterative transfer of ubiquitin generates polyubiquitinated species that are recognized and rapidly degraded by the 26S proteasome. E3 enzymes, or ubiquitin ligases, catalyse the terminal step in ubiquitin transfer, and as such are the crucial determinants of substrate specificity. Substrate recognition depends on often ill-defined sequence elements, referred to as degrons, that are the binding sites for cognate E3 enzymes^{1,2}. The E3–substrate interaction can be regulated at several levels. In some instances, limiting cofactors determine E3 activity, as in the case of the anaphase promoting complex/cyclosome (APC/C), which targets mitotic cyclins and other proteins for degradation during mitosis³. In other cases, E3 recognition depends on post-translational modification of the substrate. In particular, phosphorylation is often used to direct substrates to a recently described class of E3 enzymes termed Skp1–Cdc53/Cul1–F-box protein (SCF) complexes³. SCF complexes target a broad spectrum of substrates through a repertoire of substrate-specific adapter subunits called F-box proteins⁴. The 40-amino-acid F-box motif is a binding site for Skp1, which in turn links F-box proteins to a core ubiquitination complex composed of the scaffold protein Cdc53/Cul1, the RING-H2 domain protein Rbx1 (also known as Roc1 or Hrt1) and, usually, the E2 enzyme Cdc34 (ref. 3). F-box proteins capture phosphorylated substrates by means of carboxy-terminal protein–protein interaction regions, such as WD40 repeat domains or leucine-rich repeat (LRR) domains⁵.

Cell cycle progression requires the precisely ordered elimination of cyclins and cyclin-dependent kinase (CDK) inhibitors by the ubiquitin system³. In yeast, commitment to division, called Start, requires a threshold level of G1 cyclins (Cln1/2/3), which serve to activate Cdc28 (also called Cdk1) in late G1 phase. As cells pass Start, B-type cyclin (Clb5/6)–Cdc28 kinases are activated, which is a necessary step for initiation of DNA replication⁶. The primary function of Cln–Cdc28 activity is to phosphorylate an inhibitor of the Clb–Cdc28 kinases called Sic1, thereby targeting it for

degradation^{6,7}. Phospho-Sic1 is specifically recognized by the F-box protein Cdc4, which recruits Sic1 for ubiquitination by the Cdc34–SCF complex^{4,5,8}. Overexpression of stabilized forms of Sic1 that lack Cdc28 phosphorylation sites cause an arrest at the G1 phase⁹, whereas deletion of *SIC1* causes premature DNA replication and rampant genome instability¹⁰. Cdc4 recruits several other substrates to the SCF core complex in a phosphorylation-dependent manner, including the Cln–Cdc28 inhibitor/cytoskeletal scaffold protein Far1, the replication protein Cdc6 and the transcription factor Gcn4 (ref. 3). In the mammalian cell cycle, SCF complexes target phosphorylated forms of cyclin E1 and the CDK inhibitor p27^{Kip1} (refs 11, 12). The important role of SCF pathways is shown by the G1 phase arrest caused by non-phosphorylatable forms of p27^{Kip1}, and by the genome instability caused by expression of stabilized forms of cyclin E1 (refs 13, 14). In addition to cell cycle control, SCF-dependent proteolysis regulates the NFκB and Wnt/β-catenin signalling pathways, among others¹⁵.

Despite the well documented requirement for substrate level phosphorylation in SCF-dependent ubiquitination, the mechanism by which phosphorylation drives substrate binding is unclear. On the one hand, for some substrates the binding interaction seems to be based on recognition of a phosphopeptide motif^{16–18}, in a manner that is analogous to phosphorylation-dependent recognition by Src homology 2 (SH2), phosphotyrosine-binding (PTB), 14-3-3 and forkhead-associated (FHA) domains¹⁹. On the other hand, degradation of substrates such as Sic1, Cdc6 or Cln2 seems to require phosphorylation on multiple, widely spaced sites^{9,20,21}. Notably, alignment of the numerous genetically relevant phosphorylation sites in such substrates does not reveal an obvious consensus binding motif. To address the mechanism of Cdc4 substrate recognition, we surveyed a number of synthetic phosphopeptides derived from known substrates and identified a high-affinity consensus binding motif, termed the Cdc4 phospho-degron (CPD). The endogenous Cdc4 substrate Sic1 contains nine suboptimal CPD sites, at least six of which must be phosphorylated to allow recognition by Cdc4. This requirement for multisite phosphorylation not only imposes a threshold for Cln–Cdc28 kinase in late G1

phase, but may also confer switch-like characteristics on Sic1 degradation and subsequent entry into S phase.

Phosphorylation requirements in Sic1 recognition

Previous analysis has shown that multiple phosphorylation sites contribute to Sic1 ubiquitination *in vitro* and degradation *in vivo*⁹. To systematically determine the relative contributions of each of the nine CDK consensus sites in Sic1, we mutated each individual site and assessed the mutant proteins for stability *in vivo*, as well as for binding to Cdc4 *in vitro* (Fig. 1). Each of the mutant proteins inhibited Clb5–Cdc28 kinase activity to the same extent as wild-type Sic1, including Sic1^{Op}, which lacks all nine phosphorylation sites (Fig. 1b). Consistent with previous studies^{9,22}, conditional overexpression of *SIC1*^{T45A} from the *GAL1* promoter permanently arrested cells in G1 phase, whereas expression from the wild-type promoter caused a modest G1 delay (data not shown). Repression of the various *GAL1-SIC1* constructs in cells arrested at Start by mating pheromone allowed an estimate of Sic1 half-life *in vivo*, using a curve-fitting algorithm (Fig. 1c). Several phosphorylation sites contributed to Sic1 instability, with a rank order of Thr 45, Ser 76, Thr 5 and Thr 33, followed by less significant contributions from other sites. Analysis of binding of the panel of Sic1 mutants to

Cdc4 indicated that no single site is required for recognition, although loss of either the Thr 45 or Ser 76 sites did diminish *in vitro* binding to some extent (Fig. 1d). We then determined which individual sites, or combinations thereof, are sufficient for degradation. Beginning with the Sic1^{Op} mutant, which lacks all nine CDK sites, we restored increasing numbers of phosphorylation sites and assessed the effects on viability and the Cdc4–Sic1 interaction. Serial reintroduction of the top-ranked five sites failed to restore Sic1 binding to Cdc4 or degradation *in vivo* (Fig. 1e, f). However, introduction of either of two combinations of six CDK phosphorylation sites or seven sites proved sufficient for Cdc4 binding *in vitro* and Sic1 degradation *in vivo* (Fig. 1e, f). Thus, efficient phosphorylation-dependent recognition by Cdc4 minimally requires six of the nine CDK phosphorylation sites in Sic1.

Identification of a Cdc4 phospho-degron

There are at least three possible modes of phospho-Sic1 binding to Cdc4. The first is a phosphorylation-dependent conformational change that exposes a pre-existing cryptic binding epitope on Sic1. The second is direct binding of multiple phosphorylated residues to multiple, distinct binding sites on Cdc4. The third mode is equilibrium binding of multiple phosphorylated residues on Sic1 with a single phospho-recognition site on Cdc4. To investigate these three different possibilities, we examined the ability of various synthetic phosphopeptides to bind to Cdc4 *in vitro* by fluorescence polarization. Phosphopeptides corresponding to sequences centred on Thr 45 of Sic1 or another candidate interaction site in Far1 (ref. 23) could neither bind to Cdc4 in this assay, nor block the interaction between full-length, phosphorylated substrates and Cdc4 (Fig. 2a, Table 1 and data not shown). We then surveyed other phosphopeptides and discovered that a 19-residue peptide centred on Thr 380 of mammalian cyclin E1 (CycE^{19pT380}), a site previously implicated in cyclin E1 degradation^{24,25}, bound to Cdc4 with high affinity. A biotinylated version of CycE^{19pT380}, but not the corresponding unphosphorylated peptide, was able to capture Cdc4 from lysates of insect cells infected with a recombinant baculovirus (see Supplementary Information). Fluorescence polarization measurements performed with CycE^{19pT380} and purified recombinant Cdc4 revealed an equilibrium dissociation constant (*K*_d) of 1.0 ± 0.05 μM and a Hill coefficient of 0.99 for the interaction, indicating a single class of high-affinity binding site on Cdc4 (Fig. 2b). The phosphorylated Thr 380 site in cyclin E1 also mediates Cdc4 recognition within the context of the intact protein, as full-length cyclin E1 is targeted for degradation by SCF^{Cdc4}, both *in vitro* and *in vivo* (see Supplementary Information). Importantly, the CycE^{19pT380} phosphopeptide was able to out-compete both Cdc4 binding and ubiquitination of cyclin E1, Sic1 and Far1 (Fig. 2a and data not shown). Thus, the CycE^{19pT380} peptide binds the same site(s) on Cdc4 as its fully phosphorylated, physiological substrates.

The CPD consensus

To identify the principal phosphopeptide determinants for Cdc4 recognition, a peptide Spots array technique was used²⁶. Each position of the CycE^{19pT380} peptide was systematically altered to each of the 20 natural amino acids in a membrane-array format. Interaction of purified Skp1–Cdc4 complex with peptides on the membrane was detected with an anti-Skp1 antibody (Fig. 2c). Several characteristics of the binding site were revealed by the peptide array analysis. First, phosphorylated threonine (pThr) and a Pro at the +1 position are strictly required, consistent with the specificity of the targeting CDK kinases. Second, binding specificity is contributed by sequences that are amino-terminal to the phosphorylation site as there is a strong selection for Leu, Ile or Pro at the –1 position, whereas only Leu or Ile are preferred at the –2 position. Third, and quite unexpectedly, Arg and Lys residues seem to be disfavoured at the +2 to +5 positions, as is Tyr to a lesser extent. The optimal substrate selectivity of Cdc4 is therefore at odds

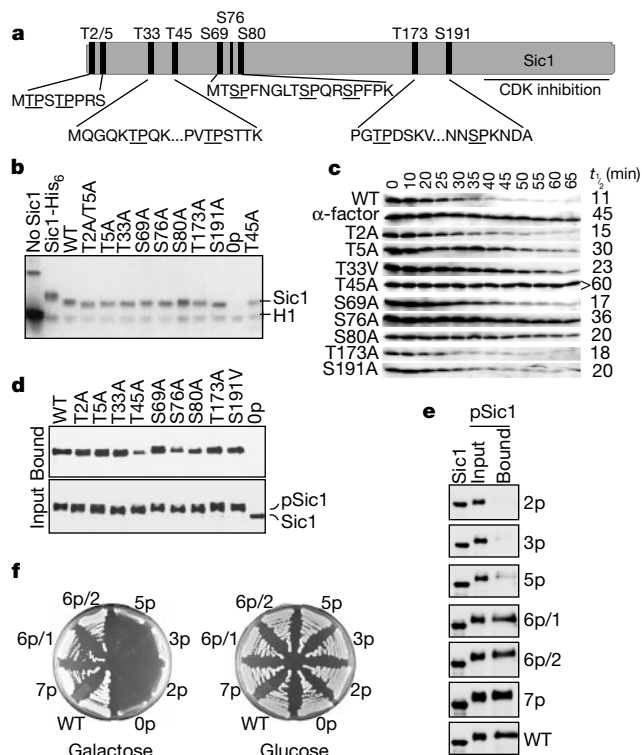


Figure 1 Contribution of CDK phosphorylation sites to Sic1 recognition, ubiquitination and degradation. **a**, Consensus S/T-P CDK phosphorylation sites in Sic1 (underlined). **b**, Inhibition of Clb5–Cdc28 kinase activity against histone H1 by purified Sic1 phosphorylation site mutants. WT, wild type. **c**, Half-life of Sic1 phosphorylation site mutants determined by decay of Sic1 protein on repression of *GAL1-SIC1^{HA}* constructs after release from α -factor arrest. The row labelled α -factor indicates decay rate of wild-type Sic1 before release. **d**, Binding of Sic1 phosphorylation site mutants to purified Cdc4. GST–Sic1 fusion proteins were phosphorylated by recombinant Cln2–Cdc28, captured onto Flag-tagged Cdc4 resin and detected with anti-Sic1 antibody. **e**, **f**, Phosphorylation of a minimum of six sites on Sic1 is required for interaction with Cdc4 *in vitro* (**e**) and degradation of Sic1 *in vivo* (**f**). Sites are indicated as follows: 2p = Thr 45, Ser 76; 3p = Thr 33, Thr 45, Ser 76; 5p = Thr 2, Thr 5, Thr 33, Thr 45, Ser 76; 6p/1 = Thr 2, Thr 5, Thr 33, Thr 45, Ser 69, Ser 76; 6p/2 = Thr 2, Thr 5, Thr 33, Thr 45, Ser 76, Ser 80; 7p = Thr 2, Thr 5, Thr 33, Thr 45, Ser 69, Ser 76, Ser 80. *GAL1-SIC1* strains were incubated for 2 days at 30 °C.

with that of the cognate targeting kinase Cdc28, which strongly prefers to phosphorylate S/T-P sequences followed by C-terminal basic residues²⁷.

The peptide Spots analysis was verified by quantitative fluorescence polarization measurements with various derivatives of the CycE^{19pT380} phosphopeptide (Table 1). The minimal peptide sequence required for binding was delimited to a core recognition sequence, LLpTPP, which bound Cdc4 with a K_d of $0.85 \pm 0.1 \mu\text{M}$. Introduction of basic residues in the +2 to +5 positions caused a decrease in solution binding affinity, most notably at the +2 and +3 positions, in conformity to the sequence preference for phosphorylation directed by CDK enzymes. The detrimental effect of basic residues is mediated by positive electrostatic potential because placement of an acetylated lysine at the +3 position yielded wild-type binding affinity. Finally, the essential pThr residue was strongly preferred over pSer (Table 1), suggesting an additional level of complexity in substrate discrimination. For brevity, we refer to the consensus binding sequence, L/I-L/I/P-P-T-P(RK)₄, as the CPD motif, where (X) refers to disfavoured residues.

Physiological CPD motifs bind over a wide range of affinities

Database searches revealed that the CPD motif is present in

numerous yeast proteins, including a recently characterized SCF^{Cdc4} substrate—the yeast transcriptional activator Gcn4 (ref. 28). Notably, one of the relevant targeting phosphorylation sites on Gcn4, Thr165, is embedded in a sequence that closely matches the CPD consensus, and a phosphopeptide centred on Thr165 of Gcn4 bound to Cdc4 with a K_d of $0.88 \pm 0.1 \mu\text{M}$ (Table 1). When the Gcn4 peptide sequence was permuted in a Spots array, the resulting consensus binding sequence closely matched that derived from cyclin E1 (Fig. 2d). The Gcn4 sequence seems to be exquisitely tuned for Cdc4 binding, as variations in the principal CPD residues at positions -2 and -1 are not tolerated as well as they are in the core cyclin E1 motif. By the same token, residues at the +2 to +5 positions appear less important, presumably because the core region binds more tightly. Another known Cdc4 substrate, Far1, also contained two reasonable matches to the CPD motif, but not within regions previously implicated in Far1 degradation^{23,29}. Sequences centred on Thr63 and Thr306 matched the CPD motif, and indeed a phosphopeptide centred on Thr306 bound weakly to Cdc4 (Table 1). Phosphorylation of this site seems to contribute to activation of Far1 by the MAP kinase Fus3 (ref. 30), raising the possibility that Far1 activation is directly coupled to its recognition by SCF^{Cdc4}. To directly compare the relative affinities of

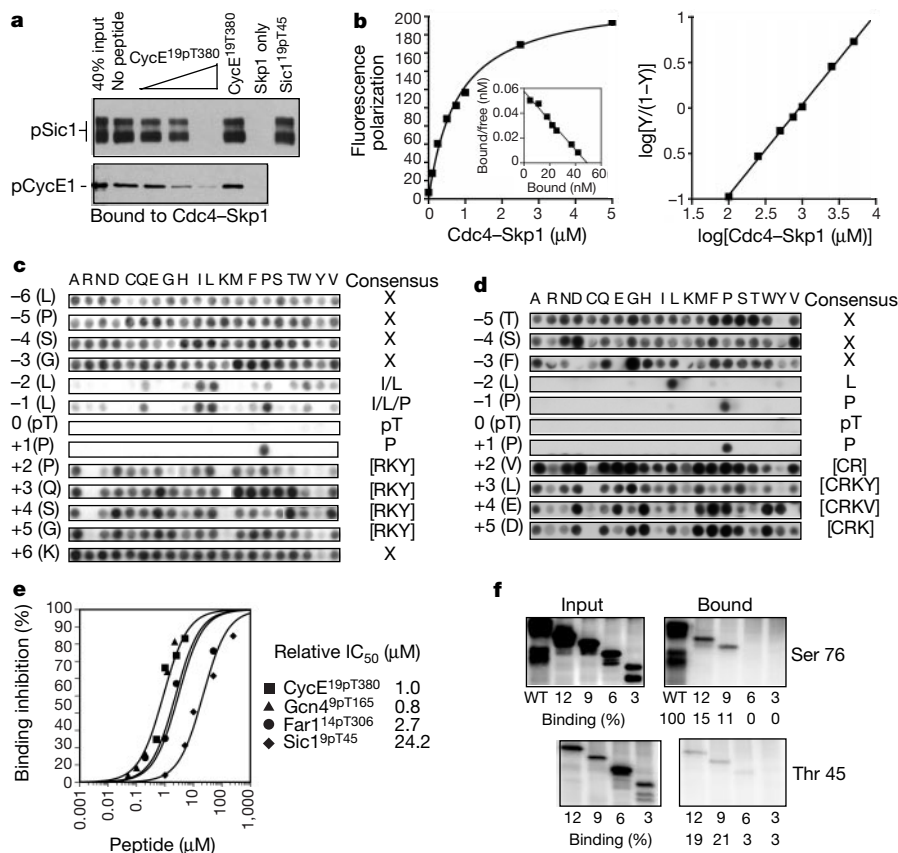


Figure 2 A phosphopeptide derived from cyclin E1 defines a high-affinity binding site on Cdc4. **a**, A CycE^{19pT380} phosphopeptide (ASPLPSGLLP^{19pT380}QSGKKQS), but not a Sic1^{19pT45} phosphopeptide, out-competes binding of phospho-Sic1 and phospho-cyclin E1 to Cdc4. The indicated peptides (3, 17 and 68 μM for CycE^{19pT380}, 68 μM for CycE^{19pT380} and Sic1^{19pT45}) and phosphoproteins were incubated with Flag-tagged Skp1-Cdc4 resin. Bound proteins were detected with anti-Sic1 and anti-cyclin E1 antibodies. **b**, Michaelis-Menton plot (left), Scatchard plot (inset) and Hill plot (right) for the CycE^{19pT380} phosphopeptide interaction with Skp1-Cdc4 as measured by fluorescence polarization. **c**, **d**, Delineation of the CPD consensus with membrane-bound arrays of synthetic peptides based on the CycE^{19pT380} sequence (**c**) or the Gcn4^{11pT165} sequence (**d**). Each residue in the seed sequence (vertical axis) was substituted with every natural amino acid

(horizontal axis). Membranes were incubated with purified Skp1-Cdc4 complex followed by detection with anti-Skp1 antibody. **e**, Peptide half-maximal inhibitory concentration (IC₅₀) curves for competition of fluorescein-CycE^{9pT380} peptide away from Cdc4 by CycE^{19pT380}, Gcn4^{9pT165}, Far1^{14pT306} or Sic1^{19pT45}. **f**, Concatamerization of low-affinity CPD motifs generates a high-affinity interaction with Cdc4. Varying numbers of suboptimal CPD sites corresponding to sequences derived from either a mixed Sic1 S76/S80/T55 site (GLTSPQRSFPFKSSPPRS) or the Sic1 T45 site (VTPSKPVTPSKPVTPSR) were produced as GST fusion proteins, phosphorylated by Cln2/Cdc28 in the presence of [³²P]- γ -ATP, captured onto Flag-tagged Cdc4 resin and detected by autoradiography. Capture efficiencies were normalized to that of phospho-Sic1.

several CPD-containing peptides, we used a competition binding assay with a fluorescein-labelled CycE^{9pT380} peptide. Under these conditions, it was possible to detect weak binding of a Sic1^{9pT45} peptide, demonstrating that weak CPD sites can mediate a direct interaction with Cdc4. Other natural CPD sequences bound with various affinities in the rank order $\text{Gcn4}^{4pT165} \geq \text{cyclin E}^{19pT380} > \text{Far1}^{21pT306} \gg \text{Sic1}^{9pT45}$ (Fig. 2e).

Concatamers of weak CPDs bind Cdc4 with high affinity

To test the possibility that the multiple, weak phospho-dependent interaction motifs in Sic1 might act in concert to form a high-affinity phospho-Sic1–Cdc4 complex, we tested artificial substrates that carried increasing numbers of weak CPD sites derived from Sic1. Glutathione S-transferase (GST) fusion proteins were constructed with 3, 6, 9 or 12 CDK phosphorylation sites, with the reiterated sequences corresponding to either an artificial fusion of the Ser 76, Ser 80 and Thr 5 (substituted to Ser) sites of Sic1, or to a variant form of the Thr 45 site with a disfavoured Lys residue substituted at the +3 position. The purified fusion proteins were phosphorylated with Cln2–Cdc28 kinase in the presence of [^{32}P]- γ -ATP and assayed for Cdc4 binding. Although fusion proteins tagged with three or six weak CPD sites proved incapable of interacting with Cdc4, those with 9 or 12 sites were efficiently captured by Cdc4 (Fig. 2f). All non-phosphorylated versions of the GST–CPD fusions failed to interact with Cdc4 (data not shown). It is unlikely that the two different polymers of weak CPD sites fold into a defined tertiary structure; indeed, circular dichroism studies suggest that in its unphosphorylated form Sic1 lacks any detectable higher-order structure (see Supplementary Information). On the basis of these results, we conclude that the Sic1–Cdc4 interaction is directly mediated by a series of low-affinity CPD sites.

CPD interactions with the WD40 domain

Deletion analysis of Cdc4 demonstrated that the WD40 repeat domain is sufficient for high-affinity interaction with the $\text{CycE}^{19pT380}$ peptide (Fig. 3a). In all known cases, direct phospho-dependent interactions are mediated by Arg and Lys residues³¹. To

identify potential CPD binding sites, we modelled the Cdc4 WD40 repeat sequence onto the β -transducin WD40 structure³², and mapped eight Arg residues conserved between Cdc4 and its homologues from *Candida albicans* and *Schizosaccharomyces pombe* onto the predicted β -propeller structure (Fig. 3b and Supplementary Information). Each Arg residue was individually mutated to Ala, and the resulting mutants were tested for their ability to rescue Cdc4 function in a plasmid shuffle experiment. Three substitutions abrogated Cdc4 function *in vivo* (R467A, R485A and R534A), whereas the other five mutations did not alter growth rate (Fig. 3c). Recombinant R467A, R485A and R534A mutant proteins were severely compromised for binding to both the $\text{CycE}^{19pT380}$ peptide and phosphorylated Sic1 protein, whereas the neutral mutation, R443A, did not alter binding (Fig. 3d, e). Thus, consistent with the single class of high-affinity binding site identified by fluorescence polarization, we have found a single, localized interaction region in Cdc4. We speculate that the triad of essential Arg residues may comprise a phospho-Thr binding pocket.

A single, optimal CPD efficiently targets Sic1

To test whether an optimal CPD can function in a heterologous context, we inserted the cyclin E1 peptide motif or derivatives thereof into the Sic1^{Op} mutant. When the $\text{CycE}^{19pT380}$ sequence was placed at the Thr 45 site of Sic1^{Op} ($\text{Sic1}^{T45::\text{CycE}19T380}$), it was able to confer recognition and ubiquitination by SCF^{Cdc4} *in vitro* (Fig. 4a, b), and elimination of Sic1^{Op} *in vivo* (Fig. 4c). Mutation of the

Table 1 Affinities of phosphopeptides for Cdc4

Peptide name	Peptide sequence	K_d (μM)
Effect of peptide length		
$\text{CycE}^{19pT380}$	ASPLPSGLLP ⁺ TPPQSGKKQS	1.0 ± 0.08
$\text{CycE}^{16pT380}$	ASPLPSGLLP ⁺ TPPQSGK	0.9 ± 0.1
CycE^{9pT380}	GLLP ⁺ TPPQSG	1.0 ± 0.05
CycE^{5pT380}	LLP ⁺ TPP	0.85 ± 0.1
Effect of pT		
CycE^{9pT380}	GLLP ⁺ TPPQSG	1.0 ± 0.05
$\text{CycE}^{9pT380S}$	GLLP ⁺ SPPQSG	6.0 ± 0.9
$\text{CycE}^{9pT380Y}$	GLLP ⁺ YPPQSG	ND
CycE^{9pT380}	GLLP ⁺ TPPQSG	ND
Cost of basic residues		
CycE^{9pT380}	GLLP ⁺ TPPQSG	1.0 ± 0.05
$\text{CycE}^{9pT380/L378K}$	GKLP ⁺ TPPQSG	12 ± 2
$\text{CycE}^{9pT380/L379K}$	GLK ⁺ TPPQSG	6.0 ± 1.2
$\text{CycE}^{9pT380/P381A}$	GLLP ⁺ TAPQSG	ND
$\text{CycE}^{9pT380/P381K}$	GLLP ⁺ TPKQSG	6.3 ± 1.2
$\text{CycE}^{9pT380/Q382K}$	GLLP ⁺ TPPKSG	5.1 ± 1.4
$\text{CycE}^{9pT380/Q382K(Ac)}$	GLLP ⁺ TPPK(Ac)SG	1.0 ± 0.1
$\text{CycE}^{9pT380/S384K}$	GLLP ⁺ TPPKQK	4.3 ± 1.3
$\text{CycE}^{9pT380/G385K}$	GLLP ⁺ TPPKSK	2.4 ± 0.8
Other phosphopeptides		
CycE^{9pT380}	GLLP ⁺ TPPQSG	1.0 ± 0.05
Gcn4^{4pT165}	FLP ⁺ TPVLED	0.88 ± 0.1
Far1^{20pT63}	PKPLNL ⁺ SKLP ⁺ SPPPSLKKTA	ND
Sic1^{9pT45}	VPV ⁺ TPSTTK	ND
$\text{Far1}^{14pT306}$	TGEFPQ ⁺ FpTPQEQLI	$\sim 25 \pm 6$
p27^{9pT187}	VEQ ⁺ pTPKKPG	ND

Results are the average of at least three individual sets of fluorescence polarization readings. Errors are s.e.m. of all measurements. Values for which saturation binding could not be achieved are indicated as approximate (–). ND indicates no binding detected by fluorescence polarization up to a Skp1–Cdc4 concentration of 10 μM . pT, phosphorylated threonine.

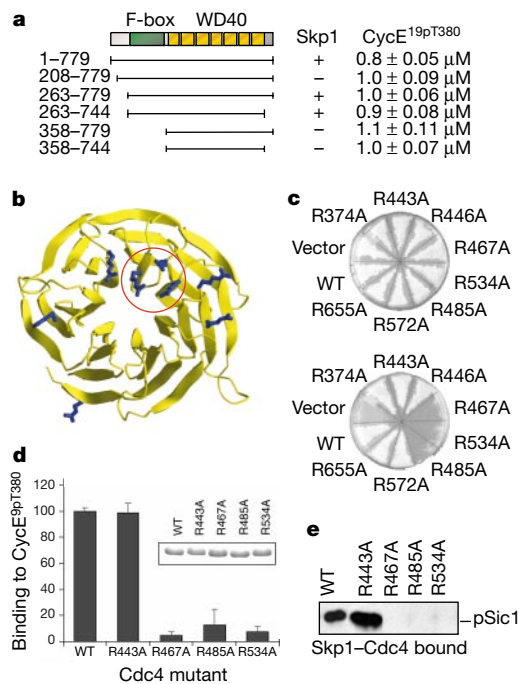


Figure 3 Identification of a CPD binding site on Cdc4. **a**, Equilibrium binding constants for the Cdc4– $\text{CycE}^{19pT380}$ interaction determined by fluorescence polarization for a series of Cdc4 deletion mutants. Skp1 binding to Cdc4 was determined by anti-Skp1 immunoblot. **b**, Ribbons diagram of α -carbon chain of the β -propeller structure of the WD40 repeat region of β -transducin. Conserved Arg residues in Cdc4 are superimposed at predicted positions. The cluster of three Arg residues critical for Cdc4 function and peptide binding is circled in red. An alignment of Cdc4 homologues is provided in Supplementary Information. **c**, Mutation of any one of three conserved Arg residues (R467A, R485A, R534A) abolishes Cdc4 function *in vivo*. *CDC4* alleles on a *TRP1 ARS CEN* plasmid were transformed into a *cdc4Δ* strain containing a *CDC4 URA3 CEN* plasmid and plated on either Trp[–]Ura[–] (top) or 5-FOA medium (bottom) for 2 days at 30 °C. **d**, **e**, Interaction of Cdc4 WD40 repeat region mutants with CycE^{9pT380} peptide measured by fluorescence polarization (**d**) and phospho-Sic1 detected by anti-Sic1 antibody (**e**). Equal levels of soluble protein were tested (inset).

phospho-Ser/Thr followed by a Pro residue. The second is a strong preference for Leu and Ile residues in the -2 position and Pro, Leu or Ile residues in the -1 position. The final requirement is a bias against basic residues in the +2 to +5 positions. Given the minimal experimentally determined CPD, LLpTPP, it is apparent why

inspection of many known phosphorylation sites implicated in targeting various substrates to Cdc4 has failed to yield an obvious consensus sequence. In particular, the nine CDK sites in Sic1 are all non-optimal CPD motifs in that a basic residue is present in the +2 to +5 positions, or a Thr phosphorylation site is replaced with a lower-affinity Ser site, or the -1 and -2 positions lack the preferred hydrophobic residues. Similarly, the eight CDK phosphorylation sites that influence Cdc6 recognition by Cdc4 lack one or more features of an ideal CPD^{20,39,40}. The apparent low affinity of each individual site in Sic1 for Cdc4 explains the requirement for multisite phosphorylation. Given the evidence suggesting a single coincident CPD and substrate-binding site on Cdc4, the strong interaction between phospho-Sic1 and Cdc4 is most easily explained by a high, local concentration of low-affinity binding sites, which effectively drive the equilibrium towards complex formation. The apparent ability of SCF complexes to multimerize might also potentiate high-affinity interactions *in vivo*⁴¹.

There can be no absolute mechanistic requirement for multiple phosphorylation sites in substrate recognition by Cdc4, as it is capable of efficiently capturing substrates that bear a single, high-affinity CPD motif. Despite this observation, it seems probable that most physiological substrates of Cdc4 are recognized by multiple, low-affinity CPD sites. Although Gcn4 degradation depends in part on the strong CPD site centred on Thr 165 (ref. 28), it has recently been shown that multiple CDK sites contribute to Gcn4 degradation⁴²; notably, all of these other sites occur within sub-optimal matches to the CPD. Two other Cdc4 substrates, Far1 and Ash1, also seem to require phosphorylation on multiple sites for efficient Cdc4 recognition (M.T., X.T. and Q. Liu, unpublished data). Similarly, the LRR-containing F-box protein Gr1 targets Cln2 for degradation in a manner that is dependent on multisite phosphorylation^{5,21,43}. The possible generalization of this theme awaits careful investigation of binding interactions between metazoan F-box proteins and their substrates.

Multisite phosphorylation and biological thresholds

Identification of the CPD sequence has uncovered an unexpected mechanism in phosphorylation-dependent protein interactions, in which kinase specificity and phosphopeptide recognition can be antagonistic, as opposed to cooperative, parameters. A dynamic balance between phosphorylation and recognition by the ubiquitination machinery could provide flexibility in substrate degradation to allow fine-tuning of irreversible regulatory switches, such as those that occur in cell cycle transitions. In late G1 phase, a threshold level of Cln1/2-Cdc28 activity is required to activate events associated with Start, including elimination of Sic1 (refs 7, 9). If, on average, phosphorylation on a single site led to the immediate degradation of Sic1, then small fluctuations in CDK activity would erode the ability of the pool of Sic1 to restrain Clb-Cdc28 activity, potentially leading to an uncoordinated onset of DNA replication. Although a single, optimal CPD motif in Sic1 results in recognition by Cdc4, and consequent ubiquitination and degradation, it does not allow precise control of Start, but rather causes precocious S-phase onset and chromosomal instability. That is, the single CPD motif fails to set an appropriate threshold for S phase because it is recognized too efficiently. In contrast, degradation of wild-type Sic1 demands phosphorylation of at least six out of nine CDK sites, thereby imposing a much higher kinase threshold. In this sense, Sic1 integrates Cln-Cdc28 activity, and potentially other G1 phase kinases^{33,44}, to act as a timing device for G1 phase.

Multisite phosphorylation is a common feature of many protein kinase substrates, and yet its biological role has remained enigmatic in most instances. Phosphorylation on multiple residues may help enable events such as multisite docking interactions, integration of different kinase pathways, substrate dephosphorylation, subcellular localization, and protein activity⁴⁵. Perhaps most importantly though, multisite phosphorylation occurs within a cellular

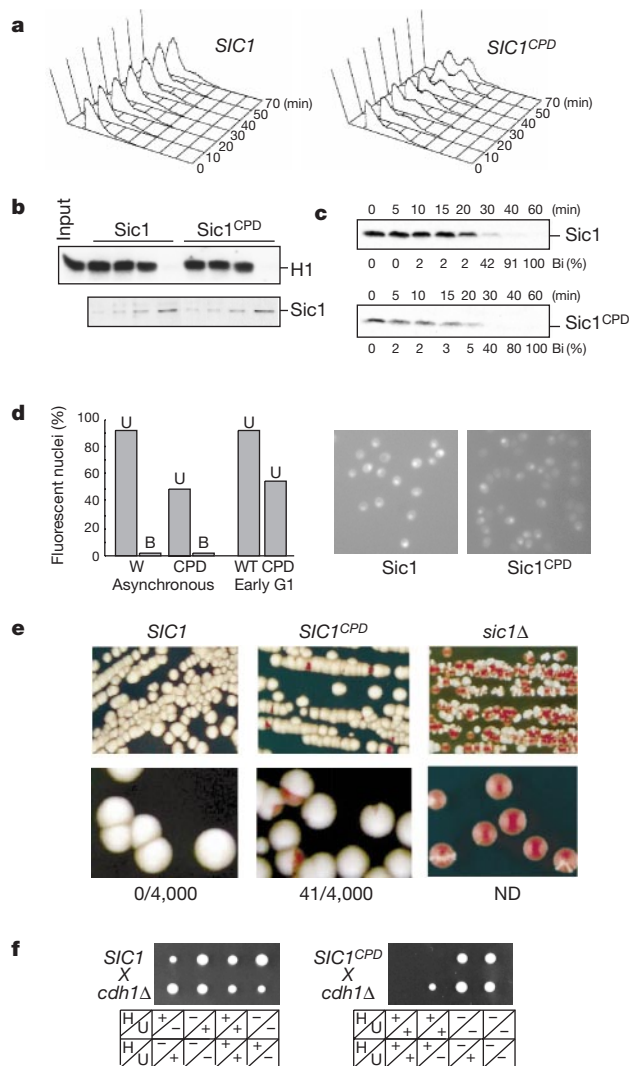


Figure 5 Premature DNA replication and genome instability caused by a single, optimal CPD motif. **a**, Strains bearing integrated wild-type *SIC1* or *SIC1*^{2p-S76LLTPP} (*SIC1*^{CPD}) alleles were synchronized in G1 phase with α -factor, released into raffinose medium at 25 °C and assessed for total DNA content by FACS analysis. **b**, CDK inhibitory activity of Sic1 and Sic1^{CPD} measured against recombinant Clb5-Cdc28 complexes. At the highest concentration shown, both wild-type Sic1 and Sic1^{CPD} are in slight stoichiometric excess of Clb5. **c**, Premature onset of Sic1^{CPD} protein instability *in vivo*. Strains bearing a *cdc28-13* allele and either wild-type *SIC1* or the *SIC1*^{CPD} allele were incubated at 37 °C for 2 h, released at 25 °C and immunoblotted for total Sic1 protein. Bi, per cent of budded cells. **d**, Analysis of Sic1- and Sic1^{CPD}-GFP fusion protein levels in individual cells. Asynchronous cultures or early-G1-phase populations obtained by centrifugal elutriation were assessed for nuclear fluorescence in unbudded (U) and budded (B) cells. Representative fields are shown for G1-phase populations. **e**, Genome instability caused by the *SIC1*^{CPD} allele. Strains carried a marker chromosome that confers an Ade⁺ phenotype (white colonies); red sectors indicate a chromosome loss event. Primary chromosome loss events were determined by scoring 4,000 individual colonies for red sectors that formed approximately 50% of the colony mass. ND, not determined. **f**, Synthetic lethal interaction between *cdh1Δ* and the *SIC1*^{CPD} allele. H and U indicate deduced His and Ura prototrophy. Of 66 tetrads from the *SIC1*^{CPD}-*URA3* cross, 46 His⁺ Ura⁺ spore clones did not form colonies, whereas 19 formed small colonies that could not be propagated. Of 31 tetrads from the *SIC1*-*URA3* cross, two His⁺ Ura⁺ spore clones did not form colonies, whereas 20 formed colonies that could be stably propagated.

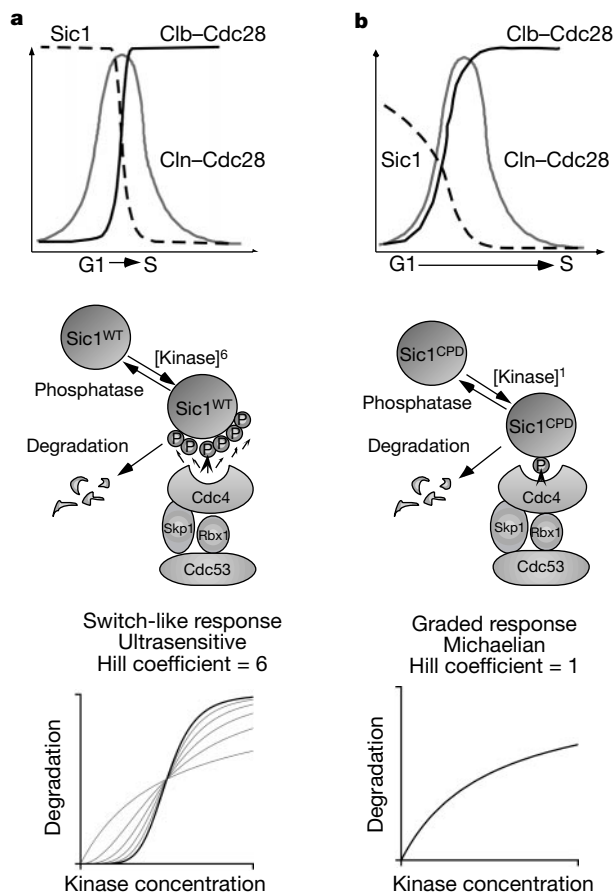


Figure 6 Ultrasensitivity at the G1/S transition. **a**, Multisite phosphorylation of Sic1 sets a threshold for Cln-Cdc28 activity and converts the increase in Cln-Cdc28 into a switch-like response for degradation of Sic1 and onset of Clb-Cdc28 activity. The requirement for six distributive phosphorylation events in Sic1 targeting creates an ultrasensitive response, as modelled by degradation = $([kinase]^{nH}) / (K_m + [kinase]^{nH})$, with a Hill coefficient (nH) of six. K_m , Michaelis constant; [kinase], kinase concentration. The theoretical Hill plot shows the predicted response for 1, 2, 3, 4, 5 and 6 phosphorylation sites. **b**, A single, optimal CPD site suffices for efficient targeting of Sic1 to Cdc4, but only with Michaelian kinetics (Hill coefficient = 1) and a hyperbolic response curve, which results in premature onset of degradation and uncoordinated activation of Clb5/6-Cdc28.

environment in which kinases and phosphatases act in dynamic equilibrium, a situation that lends itself to the generation of ultrasensitive biological responses³⁵. The requirement for phosphorylation of Sic1 on at least six sites by definition introduces a highly cooperative step in the degradation pathway (Fig. 6). That is, the forward reaction rate to form the sextuply phosphorylated Sic1 varies as the sixth order of CDK kinase concentration, provided that phosphorylation occurs in a multi-hit distributive manner, as seems to be the case⁴⁶. Additional factors may well enforce the switch-like elimination of Sic1, including inhibition of the phosphatase back-reaction upon Cdc4 binding, operation of CDK kinases at near saturation, and feed-forward action of liberated Clb5/6-Cdc28 on Sic1 (ref. 35). Quantitative measurements of Sic1 degradation rate as a function of relevant kinase, phosphatase, SCF and proteasomal activities should permit mathematical modelling of this crucial step in cell cycle progression, and allow further elucidation of the basis for switch-like biological responses. □

Methods

Yeast strain construction, culture growth, FACS analysis, fluorescence microscopy and plasmid mutagenesis were carried out by standard methods^{38,47}. Strains, plasmids and oligonucleotides used are listed in Tables 1, 2 and 3 of the Supplementary Information. We

carried out recombinant protein production, peptide synthesis, binding assays, kinase assays and ubiquitination reactions essentially as described^{5,48}. Determination of the equilibrium-binding constant was by fluorescence polarization on a Beacon 2000 Fluorescence Polarization System (Pan Vera) at 22 °C using 5 nM fluorescein-labelled probes. Peptide arrays were constructed according to the Spots-synthesis method²⁶ and probed with 1 μM purified Cdc4-Skp1 complex. Complete details of all methods are provided in the Supplementary Information.

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