High resolution crystal structure of the human PDK1 catalytic domain defines the regulatory phosphopeptide docking site

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3-phosphoinositide dependent protein kinase-1 (PDK1) plays a key role in regulating signalling pathways by activating AGC kinases such as PKB/Akt and S6K. Here we describe the 2.0 Å crystal structure of the PDK1 kinase domain in complex with ATP. The structure defines the hydrophobic pocket termed the ‘PIF-pocket’, which plays a key role in mediating the interaction and phosphorylation of other substrates such as S6K1. Phosphorylation of S6K1 at its C-terminal PIF-pocket-interacting motif promotes the binding of S6K1 with PDK1. In the PDK1 structure, this pocket is occupied by a crystallographic contact with another molecule of PDK1. Interestingly, close to the PIF-pocket in PDK1, there is an ordered sulfate ion, interacting tightly with four surrounding side chains. The roles of these residues were investigated through a combination of site-directed mutagenesis and kinetic studies, the results of which confirm that this region of PDK1 represents a phosphate-dependent docking site. We discuss the possibility that an analogous phosphate-binding regulatory motif may participate in the activation of other AGC kinases. Furthermore, the structure of PDK1 provides a scaffold for the design of specific PDK1 inhibitors.

Keywords: AGC protein kinases/crystal structure/PDK1/PDK1-interacting fragment/phosphatidylinositol 3,4,5-trisphosphate

Introduction

3-phosphoinositide dependent protein kinase-1 (PDK1) is a key protein kinase, regulating the activity of a group of related protein kinases through phosphorylation. These kinases include isoforms of protein kinase B (PKB; also known as Akt; Brazil and Hemmings, 2001; Scheid and Woodgett, 2001), p70 ribosomal S6 kinase (S6K; Volarevic and Thomas, 2001), p90 ribosomal S6 kinase (RSK; Frodin and Gammeltoft, 1999) and the serum- and glucocorticoid-induced protein kinase (SGK; Lang and Cohen, 2001). These enzymes are stimulated by hormones and growth factors, and phosphorylate regulatory proteins mediating the various physiological effects of these agonists.

PDK1 possesses an N-terminal kinase catalytic domain and a C-terminal pleckstrin homology (PH) domain (Alessi et al., 1997; Stephens et al., 1998). PDK1 activates its substrates by phosphorylating these kinases at their activation loop (reviewed in Toker and Newton, 2000; Alessi, 2001). The phosphorylation of PKB by PDK1 is dependent upon prior activation of the phosphoinositide 3-kinase (PI-3-kinase) and the production of the second messenger, phosphatidylinositol 3,4,5-trisphosphate [PtdIns(3,4,5)P³], which binds to the PH domains of PDK1 and PKB. PtdIns(3,4,5)P³ does not activate either PKB or PDK1, but instead recruits and co-localizes these enzymes at the plasma membrane, allowing PDK1 to activate PKB. Unlike PKB, the other PDK1 substrates described thus far do not interact with PtdIns(3,4,5)P³ nor is the rate at which they are phosphorylated by PDK1 further enhanced by the binding of PDK1 to PtdIns(3,4,5)P³. Instead the ability of PDK1 to phosphorylate S6K, SGK and RSK is promoted by phosphorylation of these enzymes at a residue located C-terminal to the kinase catalytic domain in a region known as the hydrophobic motif (Alessi et al., 1997; Pullen et al., 1998; Kobayashi and Cohen, 1999). The kinases that phosphorylate the hydrophobic motif of S6K and SGK are unknown, but as the phosphorylation of this residue in vivo is dependent on PI-3-kinase activation, the hydrophobic motif kinases and/or the hydrophobic motif phosphatases may be regulated by PtdIns(3,4,5)P³. In the case of RSK isoforms, phosphorylation by the ERK1/ERK2 MAP kinases induces the phosphorylation of the hydrophobic motif (reviewed in Frodin and Gammeltoft, 1999).

PDK1 belongs to the same subfamily of protein kinases as its substrates, termed the AGC protein kinases, related to the cAMP dependent protein kinase (PKA), cGMP dependent protein kinase and protein kinase C (PKC). PKA is the only AGC kinase whose crystal structure has been solved (Taylor et al., 1992). Like all protein kinases, its catalytic core possesses an N-terminal lobe consisting mainly of β-sheet and predominantly α-helical C-terminal lobe (Taylor et al., 1992; Husen and Kuriyan, 2002). The ATP-binding site is located in between the two lobes (Knighton et al., 1991; Johnson et al., 2001). At the very end of the C-terminus, PKA possesses an extended loop that terminates in the sequence FXXF, which resembles the first part of the hydrophobic motif phosphorylation site of S6K and SGK (FXXFS/TY) in which the Ser/Thr is the phosphorylated residue (Biondi et al., 2000). In the structure of PKA, the FXXF motif is buried in a hydrophobic pocket in the small lobe of the PKA catalytic domain (Knighton et al., 1991) and mutation of either Phe residue drastically reduces PKA activity towards a peptide substrate (Etchebehere et al., 1997;
Unlike other AGC kinases, PDK1 does not possess a hydrophobic motif C-terminal to its catalytic domain. However, there is evidence that PDK1 possesses a hydrophobic pocket in the small lobe of its catalytic domain similar to that in PKA. We have demonstrated biochemically that the interaction of PDK1 with four of its substrates [S6K1, SGK1, PKCζ, and PKC related kinase-2 (PRK2)] is reduced or abolished by mutation of residues predicted to form part of this pocket (Balendran et al., 2000; Biondi et al., 2000). Furthermore, mutation of a central residue in the predicted pocket, Leu155, prevented PDK1 from phosphorylating and activating S6K1 and SGK1 without affecting its ability to phosphorylate either PKB or a short peptide substrate that encompasses the activation loop of PKB (T308tide; Biondi et al., 2000). The hydrophobic pocket on the kinase domain of PDK1 has been termed the ‘PIF-pocket’ after the name of the first AGC-kinase hydrophobic motif-containing peptide (PDK1-interacting fragment) that was found to bind PDK1 (Balendran et al., 1999a). It has been suggested that the PIF-pocket in PDK1 functions as a docking site, enabling PDK1 to interact with some of its physiological substrates. Furthermore, there is evidence that phosphorylation of the hydrophobic motif of S6K1, SGK and RSK2 (Balendran et al., 1999b; Frodin et al., 2000; Biondi et al., 2001) promotes the interaction of these enzymes with PDK1. These findings suggest that the PIF-pocket on PDK1 could contain a phosphate-binding site promoting the binding of PDK1 to a subset of substrates (S6K, SGK and RSK) once these enzymes have been phosphorylated at their hydrophobic motif. This would result in a physiological phosphate-dependent interaction. In addition there is evidence that occupancy of the PIF-pocket activates PDK1, because peptides that encompass the hydrophobic motif of PRK2 (Biondi et al., 2000) and RSK (Frodin et al., 2000) induce a 4- to 6-fold activation of PDK1.

Here, we describe the first crystal structure of the kinase domain of PDK1 at 2 Å resolution. The structure defines the PIF-pocket and reveals an adjacent phosphate-binding site. Furthermore, we performed structure-based mutagenesis and biochemical analysis which support the existence of such a phosphate-binding site, which could mediate the phosphate-dependent docking interaction with substrates such as S6K and SGK. Finally, we have used a novel algorithm to define the conformational state of PDK1 relative to all the reported structures of PKA, which shows that while PDK1 has all the signs of being in an active form in the crystal, its overall conformation is in between an ‘open’ and ‘closed’ state.

Results and discussion

Overall structure

The structure of the catalytic domain of PDK1 was solved by molecular replacement and refined to an R-factor of 0.19 (Rfree = 0.22). PDK1 assumes the classic bilobal kinase fold (Figure 1) and is similar to the only other AGC kinase structure solved, that of PKA [root mean square deviation (r.m.s.d.) of 1.0 Å on Cα atoms with Protein Data Bank (PDB) entry 1STC (Prade et al., 1997)]. The form of PDK1 that was crystallized comprised residues 51–359. The tip of the activation loop (residues 233–236) is disordered, as observed in other kinase structures (Johnson et al., 1996). The N-terminus (residues 51–70), which is pointing into a large void generated by the crystallographic symmetry, is also disordered. In contrast, the N-terminal extension to the kinase domain of PKA assumes an amphipathic α-helix (termed αA-helix), and packs against the kinase core (Knighton et al., 1991). The cluster of hydrophobic residues that mediates this interaction in PKA is not present in PDK1, suggesting that the N-terminus of PDK1 could have a different function from that of PKA. Interestingly, it has recently been shown that the N-terminus of PDK1 (residues 1–50) interacts with Ras guanine nucleotide exchange factors (Tian et al., 2002). Thus, this region may assume a unique conformation in PDK1, which is not defined by the structure described here.

PDK1 was crystallized in the presence of ATP, but in the absence of any divalent cations. In the early stages of the refinement, well-defined density for the entire ATP molecule could be observed. ATP adopts a different conformation from that observed in other kinase–ATP complexes (Figure 1). Perhaps due to the absence of divalent cations, the generally observed kink between the
and phosphate, caused by the interaction with such an ion, is not seen in the PDK1 structure. It is known that PDK1 can phosphorylate itself on residue Ser241 in the activation loop and that this phosphorylation is required for PDK1 activity (Casamayor et al., 1999). Indeed, we observed density for a phosphate attached to this residue (Figure 1), and extensive interactions are observed between this phospho-serine and residues from the C-terminal lobe and \( \alpha_C \)-helix (Figure 1). The interaction between Ser241 and the C-terminal lobe is similar to the equivalent interactions in PKA, but as discussed below the binding to the \( \alpha_C \)-helix differs.

The PIF-pocket

As outlined in the Introduction, PDK1 was postulated to possess a pocket (the ‘PIF-pocket’) in the small lobe of its catalytic domain, required for the binding of PDK1 to the hydrophobic motif of its substrates (Biondi et al., 2000). The PDK1 structure described here indeed reveals such a pocket, and shows that it lies in a location similar to the FXXF-binding pocket in PKA (Figure 2). PDK1 residues Lys115, Ile118, Ile119 on the \( \alpha_B \)-helix (Figure 2), Val124, Val127 on the \( \alpha_C \)-helix and Leu155 on \( \beta \)-sheet 5 form an ~5 Å deep pocket. Previous work has shown that mutation of Leu155 to Glu abolishes the ability of PDK1 to interact with a peptide that encompasses the hydrophobic motif of PRK2 (PIFtide; Biondi et al., 2000) as well as with S6K1, SGK1, PKC\( \varepsilon \) and PRK2 (Balandran et al., 2000; Biondi et al., 2000). In addition, mutation of Lys115, Ile119, Glu150 and Leu155 to Ala, reduced the affinity of PDK1 for PIFtide ~10-fold, but did not significantly diminish the ability of PDK1 to phosphorylate and activate S6K1 and SGK1 (Biondi et al., 2001). These results are in agreement with the crystal structure of the PIF-pocket, since Leu155 is located at the centre and the other residues line the wall of the pocket (Figure 2). Interestingly, in our structure, the PIF-pocket is occupied by helix \( \alpha_G \) of a symmetry-related molecule (Figure 2). Tyr288 and Phe291 make hydrophobic contacts in this pocket with almost all pocket-lining residues, remarkably reminiscent of the interactions of the phenylalanines in the FXXF motif in PKA and their hydrophobic docking site in the equivalent region of the kinase domain (Figure 2). In addition, residues Glu287, Gln292, Ile295 and Lys296 on the symmetry related loop also form contacts with residues lining the PIF-pocket. In total, 244 Å\(^2\) of accessible surface are buried by this contact. The significance of this interaction is not clear as an oligomerization event for PDK1 has not been demonstrated in solution previously. Indeed, both the isolated catalytic domain of PDK1, which was crystallized, and full-length PDK1 migrate in gel filtration chromatography as apparent monomeric species (data not shown).

The phosphate pocket

As outlined in the Introduction, substrates of PDK1, such as S6K1, interact with the PIF-pocket of PDK1 with higher affinity when they are phosphorylated at their hydrophobic motif. This suggested that a regulatory phosphate-binding site may be located close to the PIF-pocket. During refinement of the PDK1 structure, it became clear that next to the PIF-pocket another small pocket was present,
occupied by a tetrahedral oxy-anion (Figure 2). As 2.0 M sulfate was present in the crystallization conditions, this was assigned as a sulfate ion. The ion interacts with four residues lining the pocket, namely Lys76, Arg131, Thr148 and Gln150. Because of its close proximity to the PIF-pocket (~5 Å) it is possible that this sulfate-occupied pocket could represent the binding site for the phosphate on the phosphopeptide. To investigate this further, we mutated Lys76, Arg131, Thr148 and Gln150 to Ala, in order to verify the contribution of each of these residues in enabling PDK1 to interact with a peptide encompassing the hydrophobic motif of S6K1, in which the residue equivalent to Thr412 was phosphorylated (termed S6K-pHM). A quantitative surface plasmon resonance based binding assay (Figure 4A) showed that wild type PDK1 interacted with S6K-pHM, with a $K_d$ of 0.6 μM, but not detectably with the non-phosphorylated form of this peptide (S6K-HM). The PDK1[R131A] and PDK1[Q150A] mutants did not interact detectably with S6K-pHM in this assay (Figure 4B), confirming that the interactions these residues make in the PDK1 structure are of key importance. The PDK1[K76A] mutant interacted with 3-fold lower affinity ($K_d = 1.7$ μM) with S6K-pHM. The PDK1[T148A] mutant, however, possessed ~10-fold higher ($K_d = 0.06$ μM) affinity for S6K-pHM than wild type PDK1. Moreover, the dissociation of PDK1[T148A] from S6K-pHM is markedly slower than that of wild type PDK1 or PDK1[K76A] (Figure 4A). These findings are unexpected as Thr148 is within hydrogen bonding distance of the sulfate ion (Figure 2), but indicate that this residue may play a role in enabling the dissociation of PDK1 from S6K-pHM.

The binding of PDK1 to PIFtide stimulates up to 4-fold the rate at which PDK1 phosphorylates a small peptide that encompasses the activation loop motif of PKB (termed T308tide; Biondi et al., 2000), indicating that occupancy of the PIF-pocket of PDK1 activates the enzyme. Similarly, the binding of a phosphopeptide corresponding to the hydrophobic motif of RSK stimulated PDK1 activity 6-fold (Frodin et al., 2000). We have now also found that the binding of S6K-pHM to wild type PDK1 induces a maximal 5-fold activation, with a half-maximal activation occurring at a concentration of ~50 μM S6K-pHM (Figure 4C). We next assayed the specific activities of PDK1[K76A], PDK1[R131A], PDK1[T148A] and PDK1[Q150A] mutants in the absence or presence of increasing concentrations of S6K-pHM (Figure 4C). The PDK1[K76A] mutant possessed the same specific activity towards T308tide in the absence of S6K-pHM as wild type PDK1, but an ~3-fold higher concentration of S6K-pHM was required to half-maximally activate PDK1[K76A], consistent with the reduced affinity of this form of PDK1 for S6K-pHM (Figure 4C). The PDK1[K76A] mutant possessed similar specific activity towards T308tide as wild type PDK1 in the absence of S6K-pHM. In the presence of S6K-pHM, the PDK1[T148A] mutant was activated similarly compared with

**Fig. 3.** Structure-based sequence alignment. The sequences of PKA and PDK1 are aligned according to a structural superposition performed in WHAT IF (Vriend, 1990). Sequence numbering is according to PDK1. β-strands (arrows) and α-helices (bars) are shown for the PDK1 structure according to a DSSP (Kabsch and Sander, 1983) secondary structure assignment, and labelled consistent with the secondary structure element names proposed for PKA (Taylor and Radzioandzelm, 1994). Residues lining the PIF-pocket are indicated with a black dot. Residues hydrogen bonding the sulfate ion are indicated by arrows. The PDK1 residues equivalent to Ser53 and Gly186 in PKA are labelled with an asterisk.
wild type PDK1. Consistent with the inability of PDK1[K150A] to interact with S6K-pHM, this mutant of PDK1 was not significantly activated by concentrations of S6K-pHM below 0.1 mM, but at a peptide concentration of 0.3 and 1 mM, a 2- to 3-fold activation was observed (Figure 4).

At very high peptide concentrations (0.3–1 mM), the non-phosphorylated S6K-HM peptide induced a small (<2-fold) activation of PDK1 (Figure 4C). Interestingly, despite the PDK1[K76A] and PDK1[R131A] mutants being markedly less able to interact with the phosphorylated S6K-pHM peptide than wild type PDK1, high concentrations of the S6K-HM peptide activated PDK1[K76A] and PDK1[R131A] to a similar extent as wild type PDK1, indicating that the ability of these mutants to interact weakly with the S6K-HM peptide was not affected.

We evaluated the sequence conservation in the phosphate pocket of the insulin/growth factor-activated AGC family kinases (PKBα, S6K1, SGK1 and RSK1). Sequence alignments indicate that this pocket is conserved amongst these kinases (Figure 5B). The most conserved residue is Gln150, which is found in all of these AGC kinases, and the residue equivalent to Lys76 is always a basic residue (Figure 5B). Arg131 is conserved in S6K1, SGK1, RSK1 and PKCβ, but not in PKBα, PKBβ or PKBγ, where it is an Asn or Ser. Thr148 is conserved in PKBα and SGK1, but is an Ala in S6K1/RSK1 and a Cys in PKCβ. Interestingly, we have found that the Thr148Ala mutation in PDK1 did not disrupt the phosphate pocket (Figure 4). As PKBα, S6K1, SGK1 and RSK1 need to be phosphorylated at their hydrophobic motif in order to be maximally activated, it is tempting to speculate that the C-terminal hydrophobic motifs of these enzymes, when phosphorylated, bind to their own PIF/phosphate pockets, thereby generating a network of interactions similar to that of PDK1. Consistent with PKA not possessing a phosphate pocket, Lys76 and Gln150 are not conserved in PKA (Figure 3), and indeed such a pocket is not observed in the PKA structure (Figure 2).

In the case of conventional PKC, phosphorylation of the hydrophobic motif plays a role in stabilizing these enzymes rather than triggering their activation (Bormancin and Parker, 1997; Edwards and Newton, 1997). Recent studies indicate that PDK1 also interacts with isolated C-terminal fragment of PKCβ with an order of magnitude higher affinity after this is phosphorylated at the hydrophobic motif (Gao et al., 2001). Interestingly, however, PDK1 can only interact with full-length PKC when it is not phosphorylated at its hydrophobic motif and
phosphorylation of this motif was shown to somehow mask this region, preventing it from interacting with PDK1 (Gao et al., 2001). It is likely that the hydrophobic motif of conventional PKC isoforms in its dephosphorylated state will be available for interaction with PDK1, but that following its phosphorylation, it will interact more strongly with the equivalent PIF/phosphate-binding site in its own kinase domain, thereby becoming inaccessible for binding to PDK1 (Gao et al., 2001). Our results support the existence of a phosphate-binding site in PKC isoforms that could account for the higher affinity binding of the phosphorylated C-terminal hydrophobic motif to its own pocket. The alignment in Figure 5A predicts that Arg336, Lys391, Cys409 and Gln411 in PKCβ will comprise the hydrophobic motif phosphate-binding site in the PKCβ kinase catalytic domain. Thus, binding of AGC kinases to PDK1 could depend not only on the sequence surrounding the hydrophobic motif and its phosphorylation status, but also on the availability of this sequence for interaction. The structure of PDK1 does not provide a major insight into the explanation as to why PDK1 appears to interact more strongly with the hydrophobic motif of PRK2 (PIF) compared with the equivalent fragments of S6K1 and PKB phosphorylated at their hydrophobic motif (Biondi et al., 2001). Co-crystallization of PDK1 bound to substrates or peptides encompassing hydrophobic motifs could shed light into other contacts between PDK1 and its substrates.

**The αC-helix**

The PDK1 structure shows that, as in other protein kinases (Johnson et al., 2001; Husen and Kuriyan, 2002), the αC-helix (residues 124–136) is a key signal integration motif in the kinase core. One turn of the PDK1 αC-helix (residues 129–131, Figures 3 and 5) links together the N-terminal lobe, the C-terminal lobe and the active site. Arg129 points towards the activation loop and forms two hydrogen bonds with the phosphorylated Ser241, whereas Arg131 forms two hydrogen bonds with the sulfate in the phosphate pocket (Figure 5). Glu130 coordinates Lys111, which forms a hydrogen bond with the α-phosphate of the bound ATP. This interaction is conserved in all protein kinases and was shown to be crucial for activation (Johnson et al., 2001; Husen and Kuriyan, 2002). An additional residue, Tyr126, forms a third hydrogen bond with the phosphorylated Ser241. Val124 and Val127 on the αC-helix are involved in formation of the PIF-pocket (Figure 5). The αC-helix provides a structural link between the putative phosphopeptide-binding pocket and the phosphoserine in the activation loop. The fact that R131A has higher basal activity than wild type PDK1 may indicate that this residue plays a tuning role in the PDK1 structure, not only participating in the activation of PDK1 in the presence of a phosphate ion, but also in keeping the equilibrium of the enzyme towards an inactive conformation in the absence of S6K-phHM. To our knowledge this is the first report of a kinase structure in which the αC-helix is positioned by two regulatory phosphate-binding sites on either side of the helix (Figure 5). This provides a possible sensor mechanism for linking the phosphorylation-state of the activation loop and the phosphopeptide binding event in the PIF-pocket to PDK1 activity. Our findings could provide an explanation for the observation that inactive dephosphorylated PKCβ can interact with PDK1, whilst phosphorylated PKCβ does not (Gao et al., 2001). This would be possible since in the absence of activation-loop phosphorylation, the αC-helix may not be fixed in a position that enables the formation of the hydrophobic and phosphate pocket that permits the binding of the hydrophobic motif to its own kinase domain. Following phosphorylation of the activation loop by PDK1 and autophosphorylation of the hydrophobic motif, the PKCβ hydrophobic and phosphate pocket would be stabilized as in the PDK1 crystal structure.

**Activation state**

All structures of PKA solved to date show a phosphorylated T-loop and are therefore assumed to be in an active state. In addition to the unphosphorylated versus phosphorylated states of PKA, there appear to be two main conformational states possible for the latter (Zheng et al., 1993; Johnson et al., 2001). In the active, closed conformation, all residues are positioned to facilitate phosphoryl transfer. In contrast, the inactive, open conformation is seen in the absence of a nucleotide, and differs from the closed conformation by conformational
changes of the N- and C-terminal domains with respect to each other. In addition, three ‘intermediate’ structures were described from PKA, having either adenosine (PDB entry 1BKX; Narayana et al., 1997) or the inhibitors staurosporine (PDB entry 1STC; Prade et al., 1997) and balanol (PDB entry 1BX6; Narayana et al., 1999) in the ATP-binding site. Taylor et al. (1992) have described a method to distinguish between the active and inactive conformations, based on three distances: His87–pThr197 (αC-helix positioning), Ser53–Gly186 (opening of the glycine-rich loop) and Glu170–Tyr330 (C-terminal tail distance to active site; Johnson et al., 2001). In PDK1, only one of these distances, the opening state of the glycine-rich loop, can be measured due to sequence conservation (Figure 3). This distance is 12.4 Å, similar to a PKA intermediate conformation [this distance in PKA is 14.2 Å for the open, 11.8 Å for intermediate and 10.0 Å for the closed conformation (Johnson et al., 2001)]. To allow a more direct comparison of the PDK1 structure with the available PKA structures, we have analysed the conformational state of PDK1 in detail using a novel approach, which involves a principal component analysis [also called ‘essential dynamics’ (Amadei et al., 1993)] of the crystallographic coordinates. In short, this involves the construction of a covariance matrix containing the correlations between atomic shifts (with respect to an average structure) in the ensemble of all available PKA crystal structures. Diagonalization of this matrix gives eigenvector/eigenvalue sets, which describe concerted shifts of atoms (eigenvectors) together with the corresponding mean square fluctuation of the structures (eigenvalues). This approach allows a condensed description of PKA conformational states using only a few degrees of freedom, as shown previously for a range of other proteins (van Aalten et al., 1997, 2000; de Groot et al., 1998). Diagonalization of a covariance matrix built from the backbone atoms of residues 37–196, 198–283 and 286–305 results in a set of eigenvectors that describe concerted motions of the PKA backbone. In Figure 6A, all PKA structures are projected on a subspace spanned by the first two eigenvectors (i.e. those with the two largest eigenvalues). It appears that the PKA structures cluster in three main areas along the first eigenvector. On the left of the average structure (which by definition has a projection of 0.0 on all eigenvectors) are the structures that are known to be in the ‘open’ conformation (Figure 6A). Around the average structure lie the structures that have been shown to be in an ‘intermediate’ conformation (complexes with the inhibitors staurosporine, balanol and adenosine). More to the right of the average structure are the PKA structures that are known to be in the ‘closed’ conformation. Thus, we have captured the conformational state of PKA in a single variable, the translation along the first eigenvector. This is further clarified by investigation of the atomic shifts described by this eigenvector in Cartesian space (Figure 6B). A hinge-bending motion is observed between the N- and C-terminal lobes, opening and closing the active site. It is now possible to directly compare the PDK1 conformational state by projecting the structure (backbone atoms only) onto the PKA eigenvectors. Figure 6A shows that the conformation of PDK1 is close to the PKA structures that are in an ‘intermediate’ conformation, consistent with the other structural analyses described above.

Conclusions
We have reported the structure of the PDK1 catalytic domain, which, although similar to PKA, has revealed important features that increase our understanding of the mechanism by which PDK1 is regulated. The structure, together with mutational analyses, defines a phosphopeptide-binding pocket, consisting of a separate hydrophobic PIF-pocket and a phosphate-binding site, which mediates the interaction of PDK1 with the phosphorylated hydrophobic motif of S6K. This is consistent with the previous
hypothesis that phosphorylation of S6K and SGK (Biondi et al., 2001) as well as RSK (Frodin et al., 2000) at their FXXFS/T hydrophobic motif is the trigger for their interaction and phosphorylation by PDK1. In this mechanism, the PIF-pocket would physiologically only interact with the Phe residues when the Ser/Thr residue is phosphorylated. Furthermore, as the phosphate pocket is conserved in other AGC kinases, the structural features and network of interaction of the phosphate pocket with the aC-helix on PDK1 could provide insight into the mode of activation of other AGC kinases.

A significant number of human cancers, such as those in which the PTEN tumour suppressor is mutated, possess markedly elevated PtdIns(3,4,5)P_3 levels resulting in constitutive activation of PKB and S6K, which are likely to be major contributors to the proliferation and enhanced survival of these tumour cells (Leslie and Downes, 2002). Inhibitors of PDK1 could prove effective for the treatment of these types of cancer by suppressing PKB and S6K.

Materials and methods

Materials

Mammalian and insect cell culture reagents were from Life Technologies. SensorChips SA were from Biacore AB (Stevenage, UK). Glutathione–Sepharose, as well as pre-packed HiTrap Q HP and Hiloald Superdex 200 prep grade columns were from Amersham Biosciences. Dialysis cassettes were from the Slide-A-Lyzer series (Pierce). Ni-NTA–agarose was from Qiagen. Disposable ultrafiltration devices (polyethersulfone membranes) were from Vivascience. Crystallization research tools (primary antibody screening plates) were from Hampton Research. Peptides were synthesized by Dr G.Blomberg (University of Bristol, UK).

General methods

Molecular biology techniques were performed using standard protocols. Site-directed mutagenesis was performed using a QuikChange kit (Stratagene) following instructions provided by the manufacturer. DNA was used for transfection were purified from bacteria using Qiagen plasmid Mega kit according to the manufacturer’s protocol, and their sequence verified. Used human embryonic 293 cells were cultured on 10 cm diameter dishes in Dulbecco’s modified Eagle’s medium containing 10% fetal bovine serum.

Buffers

Low-salt buffer: 25 mM Tris–HCl pH 7.5, 150 mM NaCl; High-salt buffer: 25 mM Tris–HCl pH 7.5, 500 mM NaCl. Lysis buffer: 25 mM Tris–HCl pH 7.5, 150 mM NaCl 0.07% β-mercaptoethanol, 1 mM benzamidine and 20 μg/ml PMSF. Buffer A: 50 mM Tris–HCl pH 7.5, 1 μM EGTA, 1 mM EDTA, 1% (by mass) Triton X-100, 1 mM sodium orthovanadate, 50 mM sodium fluoride, 5 mM sodium pyrophosphate, 0.27 M sucrose, 1 μM microcystin-LR, 0.1% (by vol.) β-mercaptoethanol and ‘complete’ protease inhibitor cocktail (one tablet per 50 ml; Roche). Buffer B: 50 mM Tris–HCl pH 7.5, 0.1 mM EGTA, 10 mM β-mercaptoethanol and 0.27 M sucrose.

Expression, purification and characterization of the kinase domain of PDK1

A cDNA encoding for human PDK1 amino acid residues 51–359, with a stop codon inserted at position 360, was amplified by PCR using full-length human PDK1 cDNA in the pCMV5 vector (Alessi et al., 1997) as a template. The PCR strategy employed incorporates at the 5′ end of the gene an initiating methionine, a His_6 tag followed by a PreScission protease recognition sequence prior to the residue equivalent to Met51 of PDK1 (GGATCCTATAAATATGGCACATCATCATCATCATCATC-AGCCTCGG). The 3′ primer applied in this reaction was: 5′-GGAGTCTCAGCTAGCTCGAGCTGCTGCTGCGTTCGAGT-3′. The resulting PCR product was ligated into pCR 2.1 TOPO vector (Invitrogen) and then subcloned as a BamHI fragment into pFastbac1 vector (Life Technologies) for baculovirus protein expression. The resulting construct was then used to generate recombinant baculovirus using the Bac-to-Bac system (Life Technologies) following the manufacturer’s protocol. The resulting baculoviruses were used to infect S21 cells at 1.5×10^7/mL. The infected cells were harvested by centrifugation 72 h post-infection. Cell pellets corresponding to 7 l of culture were resuspended in 900 ml of Lysis buffer and cells lysed in a nitrogen cavitation chamber. Cell debris was then pelleted by centrifugation, the supernatant made 0.5 M NaCl by addition of 4 M NaCl and then incubated with Ni–NTA–agarose (10 ml resin) for 1 h. The resin was then washed 10× with 40 ml of Lysis buffer containing 0.5 M NaCl and then placed in a disposable Econo-Pac column (Bio-Rad), where the resin was further washed with 700 ml of High-salt buffer and then with 100 ml of Low-salt buffer, both supplemented with 10 mM imidazole. Elution was performed with 200 mM imidazole in High-salt buffer and 2 ml fractions were collected. Fractions containing protein were pooled, diluted to 200 mM NaCl with 25 mM Tris–HCl pH 7.5, and loaded onto a 5 ml Hi-trap Q Sepharose column. The flow-through from this step, containing PDK1, was concentrated to 4 ml and then chromatographed on a 16/60 Superdex 200 gel filtration column using an AKTA Explorer system (Amersham Biosciences) equilibrated with High-salt buffer with the addition of 1 mM diithiothreitol (DTT). PDK1 eluted in a large symmetric peak at the expected size for a monomer. The PDK1-containing peak was again pooled, concentrated and incubated with 300 μg of glutathione S-transferase (GST)-PreScission protease (expression construct kindly provided by John Heath, University of Birmingham, UK) on ice for 4 h. In order to eliminate the cleaved His-tag sequences as well as any remaining uncleaved His-PDK1 and the GST–PreScission protease, the mixture was incubated with a mixture of 200 μl of glutathione–Sepharose and 200 μl of Ni–NTA–agarose resin for 15 min and the PDK1 protein that did not bind was collected. The resulting protein consists of PDK1 (51–359) preceded by a Gly-Pro at the N-terminus. The protein at this stage of the

| Wavelength (Å) | 0.933 |
| Space group | P3_121 |
| Unit cell (Å) | a = 123.01, b = 123.01, c = 47.62 |
| Resolution (Å) | 25–2.0 (2.07–2.0) |
| Observed reflections | 77 315 |
| Unique reflections | 27 643 |
| Redundancy | 2.8 (2.5) |
| Completeness (%) | 98.0 (93.5) |
| R_{merge} | 0.091 (0.454) |
| I/σI | 7.3 (2.0) |
| R_{free} | 579 |
| R_{free} | 0.195 |
| R_{free} | 0.222 |

| Number of groups | Protein residues 71–359 |
| Water | 200 |
| ATP | 1 |
| SO_4 | 5 |
| Glycerol | 8 |
| Wilson B (Å^2) | 22.4 |
| <B> Protein | 25.6 |
| <B> Water | 35.7 |
| <B> ATP | 38.8 |

R.m.s.d. from ideal geometry

| Bond lengths (Å) | 0.005 |
| Bond angles (°) | 1.34 |
| Main chain B (Å^2) | 1.5 |

Values in parentheses are for the highest resolution shell. All measured data were included in structure refinement.
purification was apparently homogeneous as revealed by a single band after electrophoresis of 20 μg of protein on an SDS-polyacrylamide gel and staining with Coomassie Blue R250 (data not shown). Electrospray mass spectrometry revealed a main peak mass close to the expected size of this fragment of PDK1. The specific activity of PDK1 (51–359) towards the peptide T308ptide and its activation in the presence of PIFptide were identical to wild type full-length PDK1 (Biondi et al., 2000), and tryptic peptide mass finger printing indicated that PDK1 was quantitatively phosphorylated at Ser241 (data not shown). In BioCore experiments, the steady-state binding of PDK1 (51–359) to the peptide PIFptide was similar to that of the His-tag PDK1 (51–556) protein characterized previously (Balendran et al., 1999a).

Crystalization and data collection

The PDK1 (51–359) protein was concentrated to a final concentration of 8.5 mg/ml (as determined by a Bradford assay using bovine serum albumin as a standard). The sitting drop vapour diffusion method was used for producing crystals. Sitting drops were formed by mixing 1 μl of protein solution + 0.2 μl of EDFTA (100 mM) with 1 μl of a mother liquid solution (0.1 M Tris–HCl pH 8.5, 2.0 M ammonium sulfate, 16.6 mM ATP). Hexagonal crystals (Table I) of PDK1 were grown at 20°C from a mother liquid containing 0.1 M Tris–HCl pH 8.5, 2.0 M ammonium sulfate, 16.6 mM ATP. Crystals appeared after 1 day, growing to 0.05 × 0.05 × 0.2 mm over 20 days. Crystals were frozen in a nitrogen gas stream after being soaked in 0.075 M Tris pH 8.5, 1.5 M ammonium sulfate and staining with Coomassie Blue R250 (data not shown). Electrospray mass spectrometry revealed a main peak mass close to the expected size of the S6K-pHM peptide (SESANQVFLGFTYVAPSV). After incubation for 10 min at 1 mM T308ptide in the presence or absence of the indicated concentrations of the inhibitory peptide as a search model (PDB entry 1YDB), against 8–4 Å data. A single, well separated solution was found with an R-factor of 0.479 (correlation coefficient = 0.428). The structure was automatically built using warpNtrace (Perrakis et al., 1999), which found 262 of a possible 309 residues, giving an initial protein model with R = 0.293 (Rmerge = 0.318) after simulated annealing in CNS (Brünger et al., 1998). Iterative protein building in O (Jones et al., 1991) together with refinement in CNS, which included incorporation of a model for ATP, the phosphoserine in the activation loop, solvent molecules and a key sulfate molecule, resulted in a final model with R = 0.195 (Rmerge = 0.222). No electron density was observed for residues 51–70 (the N-terminus of the construct) and 233–236 (the tip of the activation loop).

All figures were made with PyMOL (http://www.pymol.org).

**PDB coordinates**

The coordinates and structure factors have been deposited with the PDB (entry 1h1w).

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