Ternary structure reveals mechanism of a membrane diacylglycerol kinase

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Diacylglycerol kinase catalyses the ATP-dependent conversion of diacylglycerol to phosphatidic acid in the plasma membrane of *Escherichia coli*. The small size of this integral membrane trimer, which has 121 residues per subunit, means that available protein must be used economically to craft three catalytic and substrate-binding sites centred about the membrane/cytosol interface. How nature has accomplished this extraordinary feat is revealed here in a crystal structure of the kinase captured as a ternary complex with bound lipid substrate and an ATP analogue. Residues, identified as essential for activity by mutagenesis, decorate the active site and are rationalized by the ternary structure. The γ-phosphate of the ATP analogue is positioned for direct transfer to the primary hydroxyl of the lipid whose acyl chain is in the membrane. A catalytic mechanism for this unique enzyme is proposed. The active site architecture shows clear evidence of having arisen by convergent evolution.
Dicylglycerol kinase (DgkA) is the smallest known kinase. In Escherichia coli, it is responsible for the adenosine triphosphate (ATP)-dependent phosphorylation of dicylglycerol (DAG) to phosphatidic acid used in periplasm membrane-derived oligosaccharide synthesis in response to osmotic shock. The enzyme has long served as the model for integral membrane enzymology, folding, and stability. It is a trimer with three active sites of the shared sites type. Each site has three active sites of the shared sites type. Each site has individual subunit models and the electron density on which the protein, is a considerably poorer substrate. Hydrophobic interactions between the non-polar surface of the ribose and the methylene of Lys94C, itself a critical residue, acts like a multiply tipped pincer to lock the adenosine of ACP in a specific orientation on the protein. The purine ring of ACP is locked in position by hydrogen bonds between the N1 and N6 of adenine and the backbone linkage of His87C and Glu85C, respectively, in CLC. Such backbone-adenine interactions form the basis of nucleotide recognition by analogy with base pairing in nucleic acids. Backbone interactions are relatively residue non-specific. This explains mutagenesis work where neither residue was identified as essential (Fig. 2b and Supplementary Table 1). Between Glu85 and His87 is Tyr86C whose tyrosyl ring hovers close to the adenyl amides of Ala29C and Ala30C (Supplementary Movie 1), both of which contribute to the helix dipole of H1C.

The two hydroxyls of ribose in ACP are tightly hydrogen bonded to the side-chain carboxyl of the essential Asp95C (Fig. 3a). These contribute to substrate binding and specificity, and explain why 2′-deoxy-ATP, with one less hydroxyl anchor to the protein, is a considerably poorer substrate. Hydrophobic interactions between the non-polar surface of the ribose and the methylene of Lys94C, itself a critical residue, acts like a multiply tipped pincer to lock the adenosine of ACP in a specific orientation on the protein (Supplementary Fig. 3c,f). We refer to these three sequential residues as the clamping triad. The purine is secured too by its N7 interaction with the ε-amino of Lys94C, which also interacts with the ϕ-phosphate of ATP, and is tethered in place by a salt bridge with Asp80C (Fig. 3a). As expected, this position is, by contrast, sensitive to mutation. Together with Glu85C, His87C and Asp95C, Lys94C acts like a multiply tipped pincer to lock the adenosine of ACP in a specific orientation on the protein (Fig. 3a). Measured $k_{\text{cat}}$ values of DgkA for guanosine triphosphate (GTP) and inosine triphosphate (ITP) are reduced by orders of magnitude compared with ATP (ref. 2). This purine selectivity makes sense in light of the ternary structure. Thus, replacing the N6 in adenine with an O6 in guanine or inosine would lead to unfavourable interactions between the two purines and adjacent backbone carbonyls of Gly83, Ser84, Glu85 and Val79. Clear evidence of instability with GTP was observed by MDS. Likewise, the crystal structure is not compatible with tight pyrimidine nucleotide binding, consistent with the observation that while adenine nucleotides dissolved crystals of DgkA, cytidine triphosphate (CTP), uridine triphosphate (UTP) and thymidine triphosphate (TTP) did not. MDS show that ATP is extremely stably bound with the same orientation and interactions as ACP in the crystal structure (Fig. 4a and Supplementary Movie 1). ACP is therefore a reliable ATP analogue.

The ternary complex asBC contains two lipid substrates (MAG1, MAG2; Fig. 1a). MAG1 has been modelled into the electron density map with its headgroup deep in the protein. The magnesium atom in the reactive 1-OH next to the γ-phosphate of ACP (Fig. 1c,f). MAG2 is in the putative lipid substrate-binding pocket, with its headgroup ~4 Å from MAG1. Their acyl chains extend into the membrane along the hydrophobic surface of the protein. Since DAG, with two acyl chains, is the canonical lipid substrate of DgkA, finding a pair of MAGs as noted, implies they demarc the lipid substrate-binding site.
The electron density for MAGs in the complex is variable and in parts, discontinuous (Supplementary Fig. 4a–d). However, in our studies of DgkA to date, we have crystal structures that include a total of 10 trimers with density in the active site for bound lipid that is variable (refs 12,17 and this work). Mapping these onto a single active site makes a convincing case that lipids reside in the binding pocket (Supplementary Fig. 4e–g). Their variable definition likely reflects acyl chain flexibility and lipid movement during reaction.

The two other active sites in the complex lack nucleotide (Fig. 1b), asAB has two MAGs situated similar to those in asBC. Like asBC, asAB includes an almost complete SH that creates a well-defined active site. By contrast, the SH of asCA is not visible in density until Ser17A. Accordingly, this active site appears wide open (Fig. 1b). It contains just one, distant MAG.

### Reaction mechanism
Mutational studies have identified residues in DgkA that impact catalysis (refs 9,13 and this work). They include Thr8, Arg9, Ser17, Gly20, Glu28, Ala30, Glu34, Glu69, Asn72, Ser73, Glu76, Asp80, Gly83, Ser90, Lys94, Asp95, Gly97 and Ala100 (Fig. 2b). Convincingly, 14 out of the 18 are proximal (≤5 Å) to the nucleotide and lipid substrates in the ternary complex and all are exposed to solvent during the MDS (Fig. 3a–c, Supplementary Fig. 5). They demarcate the substrate binding and catalytic sites. The putative catalytic site resides on the protein at the membrane/cytosol interface where the reactive moieties of the two substrates, with disparate polarities, come together for reaction.

The active site includes glutamates and an aspartate predicted to play roles in substrate binding and catalysis (Fig. 3a,b). Concerns have been raised generally regarding possible X-ray damage to anionic residues18 and to irrelevant conformational...
substates trapped as a result of collecting diffraction data with an intense synchrotron microbeam at 100 K (refs 19,20). To address these issues we have obtained a structure by serial femtosecond crystallography (SFX) with an X-ray free-electron laser (XFEL) at room temperature (RT) and ahead of radiation damage21–23 (Table 1). It suggests that damage and ‘frozen’ conformational substates were not major issues under current conditions and that the ternary complex structure reported here is reliable.

Alternative conformers for side chains of three critical residues, Glu34, Glu69 and Glu76, were observed in the XFEL structure at RT that may be functionally relevant, as discussed (Fig. 5a,b,d,e). Convincing kinetic and biochemical evidence supports a direct, in-line phosphoryl transfer mechanism for DgkA (ref. 2). This is in contrast to one that requires the formation of an enzyme-phosphate intermediate with subsequent phosphotransfer to the lipid substrate. The direct route displays a hallmark, trigonal bipyramidal phosphorus intermediate and a random kinetic pathway where one substrate does not exclude the binding of the other. Further, the γ-phosphate of ATP should be proximal to the 1-OH of the lipid substrate. This is what was observed in the ternary complex where the two entities are poised for reaction ~4 Å apart (Fig. 3c). The separation is reminiscent of what has been interpreted for DgkA and for water-soluble kinases, where a tetraphosphate analogue of the bisubstrate (ATP-DAG in the case of DgkA) is a better inhibitor than the bisubstrate itself2. Presumably, the extra space accommodates both substrates, the reactive moieties of which are brought together for reaction by a small conformational change in the protein.

Glu69 is a conserved, essential residue in DgkA (refs 9,13; Fig. 2b, Supplementary Table 1, Supplementary Fig. 1b). In the liganded asBC, its carboxyl is in hydrogen-bonding distance to the 1-OH of MAG1 (Fig. 3c). By analogy with other kinases24, we propose that Glu69 initiates catalysis by abstracting a proton from this primary hydroxyl. In this configuration, the proximal Glu34C (Fig. 5a, Supplementary Fig. 6, and Supplementary Fig. 7) is predicted to have a \( pK_a \) of 7.53 and therefore will be protonated at neutral pH, acting as a hydrogen bond donor to Glu69C at rest. This is a stable configuration in the MDS. Due to the hydrophobic nature of the pocket, deprotonation of Glu34C will elevate the \( pK_a \) of Glu69C making it a stronger base and a better general-base catalyst. We hypothesize, therefore, that deprotonation of Glu34C is the trigger for catalysis in a substrate-loaded enzyme. Interestingly, in the XFEL structure of the apo-enzyme there are two conformations for Glu34C and Glu69C. One of each conformer has a matching partner in the cryo structure described above that can be isolated from the MDS by switching the protonation states of Glu34C and Glu69C. In the case of Glu69, the alternate conformation extends deeper into the membrane.

**Figure 1 | Overall structure of the DgkA-ACP-lipid ternary complex.** (a,b) Views of the kinase trimer parallel to the membrane and from the cytosol, respectively, with subunits represented as brown, blue and green ribbons. ACP and lipid are shown in stick representation. The two grey spheres represent zinc. Putative membrane boundaries are shown as black lines. Active site asBC contains both ACP and lipid substrate. MAG1 and MAG2 are shown with carbons coloured black and violet, respectively. (c) Expanded view of zinc-ACP in the binding site. (d) Feature enhanced map (Methods section) of Zn-ACP at 1 σ. (e) Anomalous density maps for zinc contoured at 4.0 σ. (f) Feature enhanced map for lipid MAG1 in asBC contoured at 1 σ.
and has a predicted pKₐ value of 8.82. This is reflected in the MDS, where protonation of Glu69_c induces a side-chain switch to this conformation and inversion of the hydrogen-bonding configuration with Glu34_c. This is likely to be the side-chain rotamer after the proton has been removed from the primary hydroxyl of the lipid substrate. Alternatively, this may guide substrate lipid entering from the bilayer to the active site (Fig. 5d,e). With Glu34, the two conformers may also provide a rotamer after the proton has been removed from the primary hydroxyl of lipid substrate MAG1 (red) forming a reactive alkoxide which attacks the γ-phosphorus of zinc-ATP (blue). The transition intermediate includes a pentavalent γ-phosphorus where the two substrates are covalently bonded together. On collapse, the ADP and lyso-PA products form and are released returning the enzyme to its original state. The catalytic Glu69_c abstracts a proton from the primary hydroxyl of lipid substrate MAG1 (red) creating a reactive alkoxide which attacks the γ-phosphorus of zinc-ATP (blue). The transition intermediate includes a pentavalent γ-phosphorus where the two substrates are covalently bonded together. On collapse, the ADP and lyso-PA products form and are released returning the enzyme to its original state.

Since both are negatively charged, product egress might be aided by Glu76_c in the active site, accompanied by metal-binding site rearrangement, and by cationic residues Arg9 and Lys12 in the SH proposed to draw the lipid out as the SH flexes away from the site (Fig. 4f). On release, the active site is reset for another round of catalysis, with Glu34 reprotoonating from Glu69 (Supplementary Figs 6,7).

MDS show clearly that ADP is positionally unstable in the binding site (Fig. 4f, Supplementary Movie 1). DFT simulations on MDS-sampled configurations displaying intimate MAG1 contacts (similar to Fig. 3c) suggest that 1-OH proton abstraction by Glu69_c is kinetically more accessible than phosphate cleavage (Supplementary Fig. 8, Methods section). In accord with the proposed pentavalent stabilization by Asn72_c and/or Arg9_b (Fig. 2a), the cleavage step is likely to be rate-limiting.

Two MAGs have been modelled in the liganded asBC. The 1-OH of MAG1 is proximal to the carboxyl of the putative catalytic Glu69_c (Fig. 3c). Ser98_c interacts with the 2-OH of MAG1 which, in the case of the canonical substrate DAG, would be in ester linkage to an acyl chain. The carbonyl oxygen at the ester linkage in MAG1 is within hydrogen-bonding distance of critically important Ser17_B. Together, these interactions with highly conserved and/or critical residues fix the headgroup of the lipid substrate in position for proton abstraction by Glu69_c and for reaction with the γ-phosphorus of ATP (Fig. 3c). The lipid...
Distances for zinc coordination range from 2.0 to 2.2 Å. The asterisks indicate that little or no electron density was observed for the side chains of Arg9 and nitrogen are coloured red and blue, respectively. (as dashed lines with distances between non-hydrogen atoms in Å. Protein and ACP carbons are coloured green and yellow, respectively. Oxygen and hydrogen-bonding distance to the carboxyl group of Glu69, is plausible in light of a careful consideration of the available maps and the nature of the lipid interactions between MAG1 and the enzyme are shown as dashed lines only and without specifying distances. The orientation shown, with the 1-OH in unambiguously in the active site based on the available electron density maps, even at 2.05 Å resolution (Supplementary Fig. 4a–d). Accordingly, the interactions between MAG1 and the enzyme are shown as dashed lines only and without specifying distances. The orientation shown, with the 1-OH in hydrogen-bonding distance to the carboxyl group of Glu69, is plausible in light of a careful consideration of the available maps and the nature of the lipid substrate and the reaction being catalysed.

Comparison with other kinases. In support of the proposed mechanistic model, we note the striking resemblance between critical residues in the active site of DgkA and their equivalents in cAMP-dependent protein kinase (PKA) for which a similar catalytic mechanism is well established24,25 (Fig. 2a). To begin with, the catalytic residue in both kinases is anionic, Glu69 in DgkA and Asp166 in PKA. While magnesium is a common ATP counter-ion in kinases, zinc can serve this role in DgkA (ref. 16). Both zinc ions in the ternary complex bind electrostatically with the γ-phosphate of ACP and, by electron withdrawing effects, render the phosphorus more susceptible to attack by the lipid nucleophile. The same kind of interaction is seen in PKA. Lys94C coordinates with the α-phosphate of ACP in a manner analogous to its counterpart, Lys72, in PKA. Lys94C is stabilized in position by a critical Asp95C, just like the Lys72-Glu91 pair in PKA. The essential Glu76C chelates the two zincs which, in turn, bind the triphosphate of ACP. The equivalent residue in PKA is Asp184. Parenthetically, a second conformation for the side chain of Glu76C is present in the XPFI apo-structure which is more removed from the metal ions; it may play a role in product release (Fig. 5b). Arg9 is functionally important and mutating it dramatically alters DgkA activity. While its side chain lacks full density in the DgkA complex, one of its possible conformers makes a convincing interaction with the α-phosphate of ACP, reminiscent of the role played by Lys168 in PKA. Asn72 is essential in DgkA (Fig. 2b and Supplementary Table 1). Its side-chain carbonyl is 4.1 Å from Zn2 while its amide nitrogen bridges the essential Glu69 and Glu76 (Fig. 5c). The equivalent residue in PKA is Asn171; it bridges Asp166 and Asp184 and its carbonyl is 2.1 Å from manganese (Mn1).

The similarity between the binding sites of DgkA and PKA extends to the adenosine end of the nucleotide. Thus, the aforementioned clamping triad on adenine in the CL of DgkA that includes Glu85-Tyr86-His87 is matched by an almost identical triad in PKA represented by Glu121-Tyr122-Val123 (Figs 2a and 6). At the same time, the two ribose hydroxyls are anchored by Asp95 in DgkA and by Glu127 in PKA. The resemblance between residues in and chemical architecture of the active sites of DgkA and PKA is remarkable given that the two kinases are evolutionarily unrelated and have essentially no primary through quaternary structure similarities.

Mutagenesis. With the ternary complex structure in hand, hypotheses regarding the role of specific residues in DgkA enzymatic activity were tested by systematic mutagenesis. Kinetic measurements were performed with the mutants reconstituted in the LCP. A total of 23 sites were mutated with the types screened dictated by the putative role for the particular residue. The results are summarized in Fig. 2b and Supplementary Table 1 along with data from separate mutational studies9,13. A few notable mutants will be discussed here, as follows. (i) Glu69 is the putative catalytic residue in DgkA (Figs 2a and 3c), in contrast to Asp which serves that function in serine protein kinases. Strikingly, Glu69 tolerated none of the mutations tested, one of which was Asp (pKₐ 3.8). This could be a spatial effect because the side chain of Asp is too short. It may also have to do with the fact that the 1-OH in gycerides like MAG and DAG have pKₐ values (13.6) higher than those of PKA substrates Ser/Thr (pKₐ 13.0). Glucydrides therefore may require the carboxyl of Glu, with a pKₐ of 4.3, or higher enabled by nearby Glu34, for proton abstraction. (ii) The conserved Asp95 coordinates the hydroxyls of ribose in ACP, preserved throughout the MDS, and the carbonyl of Gly91 (Fig. 3a). Mutating it to Ala essentially abolished activity. However, Glu with its longer side chain did substitute but activity was reduced by 76%. By contrast, Asn had a
minor effect which makes sense given that its side-chain amide can hydrogen bond with carbonyls and hydroxyls. (iii) Ala30 resides on H1 in the active site entrance at the membrane interface (Fig. 5f). Its methyl side chain was speculated to enable lipids with small headgroups to enter the site. Mutating it to the bulky leucine reduced activity by 93% consistent with a putative gatekeeping role. (iv) Gly83 in the CL was proposed to facilitate the helix-coil transition enabling adenine binding (Supplementary Fig. 3). Mutating it to Pro was expected to compromise such activity. Indeed, the Gly83Pro mutant was virtually dead enzymatically; MDS show dynamic instability of the CL (Supplementary Movie 2) and (v) finally, Asn72 plays a key role in catalysis. Its side-chain amide bridges Glu69 and Glu76 (Fig. 5c) both of which are essential. These H-bonds bridge the two essential residues throughout the MDS. The Asn72Ala mutant is catalytically inactive. For all intents and purposes, so too are the Asn72Gln and Asn72Asp mutants. These data show how important both the precise chemistry and the architecture of this part of the binding/active site are for catalysis. The structure-based, hypothesis-testing mutations and others in Fig. 2b and Supplementary Table 1 lend credence to the complex model and mechanism of kinase action.

Discussion
DgkA is the tiniest of kinases. With just 121 residues and a trimeric constitution, three kinase catalytic sites have been fashioned (Fig. 1). Each acts at the membrane/cytosol interface. The rest of the protein creates binding pockets on either side that position the reactive ends of two large substrate molecules with dramatically different polarities in juxtaposition for reaction. The ternary structure explains in atomic detail how the nucleotide and
lipid-binding sites interact with substrate molecules. The many, contact site-docking defines a specific orientation for the nucleotide that presumably is optimized for transphosphorylation. Particularly striking is the similarity between the binding site architecture of this bacterial DgkA and mammalian PKA (Figs 2a and 6), kinases that bear no resemblance at any other level of structure. Thus, these defined points of chemical constitution arrayed precisely in three-dimensional space have evolved independently achieving a similar end in entirely different protein contexts – a convincing case of convergent evolution. DgkA’s small size shows that a functioning kinase can be built with very little protein mass. The ternary structure points to a minimum required for that purpose. Inspecting the complex suggests that a pair of helices, each 3-4 turns long, connected by a

Figure 5 | Rationalizing functional roles of highly conserved residues in DgkA not directly involved in catalysis. (a) The anionic carboxyl of Glu34C is proposed to elevate the pKa of catalytic residue Glu69C, making it a stronger base for proton abstraction from the lipid (green protein carbons). Electron density (mesh) for alternative Glu34 conformers was observed in the XFEL data recorded at RT (purple carbons) that may enable a switch in pKa at Glu69. (b) Alternative conformations for Glu76C in the XFEL structure. The first (Glu76_1) is in position to coordinate with Zn2. Black dashed lines correspond to distances 2.0–2.2 Å. The second (Glu76_2) has the coordinating oxygen at some distance (3.8 Å, green dashed line) from Zn2 where it may facilitate product release. The polyphosphate and zinc are superimposed from the ternary complex on the XFEL apo-structure. (c) Side-chain amide nitrogen of Asn72C coordinates with carboxyls of Glu69C and Glu76C, both essential residues. By weakly interacting with γ-phosphate oxygens, the electrophilicity of the γ-phosphorus is elevated making it more reactive. Additionally, the amide may stabilize a transient bisubstrate by being positioned close to where the pentavalent intermediate is likely to form between the 1-OH of MAG1 and the γ-phosphate of ACP (dotted blue line). Likely interactions between MAG1 and the enzyme are indicated by dashed lines only. (d) Alternative conformations for Glu69 observed in the XFEL structure (magenta carbons) superimposed on the DgkA ternary structure (green carbons). The first conformation (Glu69_1) is like that seen in the ternary complex and interacts with MAG1. The second (Glu69_2) extends into the membrane and may guide lipid substrate to the active site. (e) An expanded view of Glu69 in d with electron density corresponding to alternative conformers seen in the XFEL structure superposed. (f) The methyl side chain of Ala30C (red circle) is proposed to provide room for lipid substrate and product to pass between the start of SHB and the top of H1C that define the gateway into and out of the active site. The Cα–Cα distance from Ala30C (red circle) to Ala13B (blue circle) in asBC is 10.0 Å.
substrate loading and product release. The reduction in SH for lipid and nucleotide binding. This should ensure efficient involvement of small segments such as the CL for nucleotide and the SH for turnover, would appear to be relatively minor to a hypothetical solitary active site. Conformational changes, raises the probability of productive encounters between substrates easy access and egress of substrates and products. This naturally periphery (Fig. 1b) opening to the cytosol and the membrane for half in the cytosol, DgkA has features that make it a good kinase A, PDB ID: 1ATP. Only the similarities are highlighted. In a one surface of the adenosine moiety faces the protein, the other faces the cytoplasm. In b the adenosine is sandwiched between the N- and C-terminal lobes of PKA.

7-residue loop (Supplementary Fig. 3) might be all that is needed to create an ATP-binding module. The ternary structure should serve as a template for rational protein and enzyme design, optimizable for solubility, stability, specificity and catalytic efficiency.

Kinetic studies suggest that DgkA functions with a moderate level of heteroallostery. Thus, the binding of one substrate influences (in this case positively) that of its co-substrate. In a multimer like DgkA, it is possible for heteroallostery to come about through inter-subunit signalling. The fact that any given active site is a composite, with components from at least two subunits, suggests a way for the substrate-bound status of one site to be relayed to another; the SH is an obvious means for inter-subunit information transmission. In this way, the enzyme could present simultaneously three active sites in distinct states. We find some evidence for this in the ternary structure where only one of the three active sites is occupied by ACP. Further, two of the sites in the CL region, which clamps onto the nucleotide base, are profoundly different (Supplementary Fig. 3). Neither CL is involved in crystal contacts. Presumably therefore both are physiologically relevant. We propose that the three sites in DgkA are in different states at any given time during catalysis. However, it is possible to model a symmetrical DgkA trimer that is stable in MDS, indicating that the enzyme may adopt a symmetrical configuration. This does not imply that all three sites are catalytically active at the same time. Additional mechanistic and structure work is required to evaluate allosteric coupling within the enzyme.

As a trimer situated with half its bulk in the membrane and half in the cytosol, DgkA has features that make it a good biocatalyst. Its three active sites are arranged around the enzyme’s periphery (Fig. 1b) opening to the cytosol and the membrane for easy access and egress of substrates and products. This naturally raises the probability of productive encounters between substrates and binding sites, thus elevating the reaction rate compared to a hypothetical solitary active site. Conformational changes, where needed for turnover, would appear to be relatively minor involving small segments such as the CL for nucleotide and the SH for lipid and nucleotide binding. This should ensure efficient substrate loading and product release. The reduction in dimensionality, as a result of lipid substrate and enzyme confinement to a two-dimensional membrane, increases overall efficiency of delivery. The amphiphilic nature of the lipid and of the membrane in which it and the kinase reside, ensures the spontaneous and proper alignment of substrate in the lipid-binding site for reaction. These features of trimeric DgkA undoubtedly contribute to making it an efficient biocatalyst, described previously as an evolutionarily optimized integral membrane enzyme.

**Methods**

**Molecular biology and protein production.** All but two of the DgkA mutants were generated by PCR-based site-directed mutagenesis using the plasmid pTrcHisB-DgkA-WT (ref. 14) as the template. In the case of E34D and E34A, constructs with the desired mutation could not be produced after several rounds of mutagenesis trials. Instead, the corresponding genes were synthesized chemically (Genescipt). All mutations were confirmed by DNA sequencing (MWG Biotech). Detailed purification procedures are described in ref. 14. Briefly, E. coli WH1061 cells carrying DgkA plasmids and grown in Luria-Bertani broth were induced at an optical density of 0.6 for 3 h at 37 °C with 1 mM IPTG. All subsequent steps were carried out at 4 °C unless otherwise specified. For crystallization, cells from 11 of culture were lysed by sonication in buffer containing 0.2 mM TCEP, 0.3 M NaCl, 0.2 mg ml⁻¹ lysozyme, 50 µg ml⁻¹ DNAase, 10 µM BHT, 1 mM PMSE, 0.2 mM EDTA, 5 mM MgCl₂ and 75 mM Tris/HCl pH 7.8. Empigen BB was added to the cell lyaste at 3.3% (w/v) to solubilize DgkA for 1 h. Unsolubalized material was removed by centrifugation at 9,000g for 10 min. The supernatant containing N-terminally His-tagged DgkA was allowed to bind to 4 ml of Ni-NTA resin for 1 h. The resin was packed in a gravity column, which was washed with 3% (w/v) Empigen BB, 10 mM imidazole in Buffer A. The detergent Empigen was then exchanged to 0.25% (w/v) DM, 1 mM TCEP, 10 mM HEPES pH 7.5 and 1.5% (w/v) Empigen BB, 40 mM imidazole in Buffer A. The detergent Empigen was then exchanged to 0.25% (w/v) DM, 1 mM TCEP, 10 mM HEPES pH 7.5. Protein was eluted with 0.25 M imidazole in buffer containing 0.5% (w/v) DM, 1 mM TCEP, 50 mM LiCl, 0.2 mM TCEP, 20 mM HEPES pH 7.5. Protein was eluted with 0.25 M imidazole in buffer containing 0.5% (w/v) DM, 1 mM TCEP, 40 mM HEPES pH 7.5. Pooled fractions were concentrated and run through a size-exclusion chromatography with a Superdex 200 16/600 column in buffer containing 0.25% (w/v) DM, 1 mM TCEP, 0.1 M NaCl and 10 mM Tris/HCl pH 7.5. Fractions corresponding to the central peak were pooled. Protein was concentrated to 12 mg ml⁻¹, flashed frozen in liquid nitrogen as 11 l aliquots and stored at −80 °C until use. For functional assays, protein was purified from 0.1 l of culture, omitting the size-exclusion chromatography step. Size-exclusion chromatography had no effect on specific activity.

**Enzyme assays.** In meso measurements of WT and mutant DgkA kinase activity were carried out as follows. DgkA was reconstituted into LCP by mixing the protein solution with a 1.5-fold volume of monodend. Five microliters of IC-treated enzyme was dispersed into each well of a 96-well plate. The coupled enzyme assay reaction was initiated by adding 200 µl of Assay Mix (20 mM ATP, 0.1 mM EDTA, 0.1 mM...
EGTA, 55 mM magnesium acetate, 1 mM phosphonoformic acid, 0.2 mM dithiothreitol, 50 mM LiCl, 0.4 mM NADH, 20 μM of pyruvate kinase and lactate dehydrogenase, and 25 μM PMES 0.35 % w/v was made for reconstitution into the mesophase was 0.066 mg ml⁻¹. For mutants with ~70% of WT activity, protein concentration was adjusted to 0.1 – 5 mg ml⁻¹ to increase measurement sensitivity.

Crystallization. In meso crystallization of the apo form of D4-DgkA for synchrotron radiation data collection was carried out using an in meso robot.

The 90°C glass sandwich crystallization plates were incubated at 4 °C for crystal growth. The precipitant solution contained 7-9% (v/v) 2-methyl-2,4-pentanediol (MPD), 0.1 M NaCl, 0.1 M KOPO₄, 0.1 M Na₂HPO₄, 0.1 M Na₂CO₃, 0.1 M NaCl, 0.6 mM Mg(CH₃COO)₂ (magnesium acetate), 0.05 M NaCl, 0.01 mM MgSO₄, 7.9 MAG. Crystals were harvested and snap cooled in liquid nitrogen.

To produce crystals of DgkA in complex with an ATP analogue, extensive screening was performed that involved exploring protein constructs, crystal forms, host MAGs, analogues, crystallization precipitants and temperatures. Trials with D4-DgkA in 9.9 MAG with zinc-ACP at 4 °C eventually provided crystals and a structure at 2.70 Å resolution. Crystals were soaked directly in glass sandwich plates for 2 h at 4 °C using precipitant solution (7-10% (v/v) MPD, 0.1 M NaCl, 0.1 M KOPO₄, 0.1 M Na₂HPO₄, 0.1 M Na₂CO₃, 0.1 M NaCl, 0.06 M Mg(CH₃COO)₂, 0.1 M NaCl, and 0.05 mM MgCl₂) supplemented with 60 mM Zn(CH₂COO)₂, and 10 mM ACP. Harvested, was as noted above.

For measurements with the XFEL, crystallization was carried out at 20 °C in coupled micro-syringes using 7.9 MAG as host lipid.

The precipitant solution contained 0.1 M NaCl, 0.01 M NaCl, and 0.05 M NaCl, pH 5.6. Crystals were identified by bright-field and polarized light microscopy, SONICC and trypophan fluorescence at the Linac Coherent Light Source (LCLS). Cysteine mutant samples were transferred to an LCP injector and extruded for data collection into an evacuated chamber at 20 °C, as described below.

Synchrotron data collection and processing. Synchrotron diffraction data were collected at beamline 23-ID-B, Advanced Photon Source (USA), I24, Diamond Light Source (UK) and PX11, Swiss Light Source (SLS, Switzerland) at 100 K. At the Advanced Photon Source, data were collected with a 1° oscillation and a 1 s exposure per frame, a collimated beam size of 10 μm, and a sample-to-detector distance of 500–650 mm, with a Pilatus 6M detector using 0.978 Å wavelength X-rays. At the SLS, data were collected with a 0.1° oscillation and 0.1 s exposure per frame, a collimated beam size of 10 × 15 μm² and a sample-to-detector distance of 450–650 mm, with a Pilatus 6M detector using 1.033 Å X-rays.

Synchrotron-derived diffraction data were reduced with xia (ref. 30) using XDS (ref. 31) and XSCALE (Table 1). Optimum data weights were identified by data quality and isomorphism unit cell parameters. Complete data sets for D4-DgkA in 9.9 MAG (Table 1) and 15.0 MAG (Table 2) were obtained by merging data collected from 4 and 6 crystals, respectively. A single crystal was used for the anomalous data collected at the zinc edge (Zn-edge D4-99-MAG-ACP in Table 1).

XFEL data collection and processing. XFEL data were collected using the CXI instrument at the LCLS at SLAC National Accelerator Laboratory. The LCLS was operated at a wavelength of 1.02 Å (9.5 keV) delivering X-ray pulses of 30 fs pulse duration and with 8 × 10¹¹ photons per pulse focused into a spot size of ~1.5 μm × 1.5 μm² using a pair of Kirkpatrick–Baez mirrors. Needle-shaped protein microcrystals ranging in length from 10 to 40 μm dispersed in the LCP were injected into the focus region at 0.17 μm⁻¹ using a continuously flowing LCP injection with a nozzle capillary diameter of 50 μm (ref. 27). Single-shot diffraction patterns of randomly oriented crystals were recorded at a rate of 7,200 pattern per minute (120 Hz) using the Cornell-SLAC hybrid Pixel Array Detector (CPAD) (ref. 28). The sample-to-detector distance of ~122 mm was used to obtain a maximum obtainable resolution of 2.0 Å at 1.02 A wavelength. Data collection required 4.5 h of beamtime and used 42 μl mesophase corresponding to 220 μg protein. A total of 1,987,632 frames were collected, of which 180,031 (9.1%) were integrated without applying a per-pattern resolution cut-off and merged in Point Group C2 (classical) MDS. Accordingly, using ab initio-based DFT molecular-simulation techniques to probe putative reaction paths and energetics is of relevance to help in enzyme mechanism studies. Gaussian 09 (ref. 42) simulations were performed on MDS configurations like Fig. 3c, featuring intimate MAG contacts; other configurations were less energetically plausible. MM (AMBER), ONIOM-QM/MM (ref. 47) and DFT geometry-optimization led to retaining zinc atoms and residues within ~7 Å (244 atoms) and ~8 Å (365 atoms) of MAG, for MAG’s reliable energetic and geometric interactions.

Examining carefully MDS-sampled, ‘cut-down’ configurations of the ternary complex of the active site similar to the energy barriers of reaction 1-0H OH proton transfer, or rearrangement of bonds, or the formation of a (near-) covalent bond between an 1-O oxoanion (or even 1-OH with proton intact) and the γ-phosphate atom to form a covalent bond (Fig. 6). A key point regarding these possible events is their energetic (and kinetic) ‘accessibility’, especially in relation to background thermal energy ~RT. At room temperature (RT) is around 0.6 kcal mol⁻¹. Accordingly, such reaction events accessible for routine observation during geometry optimization would typically have energy barriers of ~2–3 kcal mol⁻¹, or less. Indeed, the connection between reaction-energy barriers and approximate rates is well-established in classical kinetics.

On this basis, we found that the only likely interaction with protein residues for reaction of 1-0H was with Glu69, featuring a prominent hydrogen bond with the 1-0H pointing directly to Glu69. In some cases, geometry optimization on the protein crystal structure (Fig. 3c, Table 1) was followed by a simple optimization entailed the direct transfer of the proton to Glu69, suggesting an electrostatic interaction, especially given the prominent hydrogen bond’s strong electrostatic interaction, especially given the high charge-to-mass ratio of the proton.

Optimization of the alkoxide form led to the formation of a somewhat strained, imperfect covalent bond for the resultant (1-O)-oxoanion (sans proton) with the phosphophorous atom, not unusual for a pentavalent intermediate, suggesting a reasonable kinetic accessibility with a ~2–3 kcal mol⁻¹ barrier. However, further structure relaxation did not suggest any rupture (or meaningful stretch/strain) of

Structure solution, model building and refinement. Structures were solved by molecular replacement (MR) using published models (PDB ID: 3ZEX and 3ZEX for D4-DgkA and 3X0Z and 3X0Z for Phoenix) for manual building and refinement, respectively. Ligands were built based on feature enhanced maps and compositeomit 2Fo-Fc maps. Lang et al. reported that the noise level in 2Fo-Fc maps is generally over estimated. Accordingly, the 2Fo-Fc map contoured at 0.7 σ, together with the strong electron density in the chain region of other DgkA structures (Supplementary Fig. 4), informed the initial model. The atomic model of DgkA trimer crystallized in the absence of nucleotide and an EM was calculated directly from the MR solution using Phenix without adding nucleotide to the model. For the MAG1, the EM was calculated as for the Zn-ACP but with MAG1 in place. Figures were prepared using PyMol.

Molecular dynamics simulations. All MDS were performed using GROMACS v5.0 (ref. 43). Initial Coarse Grained (CG) MD simulations using the Martini 2.2 model were done for run the assembly of the bilayer around DgkA (Fig. 4). The end-snapshot of the CG simulations were then converted to atomic detail with the crystal structure aligned with the CG protein within the assembled lipid bilayer (Fig. 5). The systems were then equilibrated for 1 ns and the MD restrained before normal atomic MD using the Gromos53a6 force field (ref. 46). Almost all cases, ATP, Zn²⁺ and MAG were included. In one instance, GTP was included instead of ATP. In a second case, ADP and lyso-phosphatidic acid were included in place of ATP and MAG. Parameters for catalytic intermediates were also designed, with the associated coordinates simulated. The systems were equilibrated with a 150-mM concentration of NaCl. In silico mutagenesis was performed using PyMol.

Density functional theory. Modelling chemical reactions is not possible with classical (MMD). Accordingly, using ab initio-based DFT molecular-simulation techniques to probe putative reaction paths and energetics is of relevance to help in enzyme mechanism studies. Gaussian 09 software B3LYP/6-31G(dp) simulations were performed on MDS configurations like Fig. 3c, featuring intimate MAG contacts; other configurations were less energetically plausible. MM (AMBER), ONIOM-QM/MM (ref. 47) and DFT geometry-optimization led to retaining zinc atoms and residues within ~7 Å (244 atoms) and ~8 Å (365 atoms) of MAG, for MAG’s reliable energetic and geometric interactions.

Examining carefully MDS-sampled, ‘cut-down’ configurations of the ternary complex of the active site similar to the energy barriers of reaction 1-0H proton transfer, or rearrangement of bonds, or the formation of a (near-) covalent bond between an 1-O oxoanion (or even 1-OH with proton intact) and the γ-phosphate atom to form a covalent bond (Fig. 6). A key point regarding these possible events is their energetic (and kinetic) ‘accessibility’, especially in relation to background thermal energy ~RT. At room temperature (RT) is around 0.6 kcal mol⁻¹. Accordingly, such reaction events accessible for routine observation during geometry optimization would typically have energy barriers of ~2–3 kcal mol⁻¹, or less. Indeed, the connection between reaction-energy barriers and approximate rates is well-established in classical kinetics.

On this basis, we found that the only likely interaction with protein residues for reaction of 1-0H was with Glu69, featuring a prominent hydrogen bond with the 1-0H pointing directly to Glu69. In some cases, geometry optimization was performed using PyMol.

In silico mutagenesis was performed using PyMol.

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the bridging γ-P–O bond. Extending systematically this covalent bond until breakage, the relative potential-energy profile (Supplementary Fig. 8b) suggests a reaction-energy barrier of ~15 kcal mol⁻¹. Unlike the hydrogen bond in protein transfer, there is no appreciable local electrostatic interaction in this bond. The much larger mass of P and O atoms and inertia of the MAGI and to-be-formed ADP moieties compounds this, altering the character of this potential-energy profile. With a large positive barrier to be crossed, it is much less kinetically favourable. Here, the substantial positive-energy ‘hill’ shows clearly the unfavourable nature of extending the bond from a local minimum in the immediate vicinity of the ‘reactant’ state, hindered by less electrostatic interaction, until rupture beyond 0.5–0.6 Å. This is somewhat consistent with previous computational estimates of 26.7 kcal mol⁻¹ (based on potential-energy differences), albeit for cleavage of ATP (only) in explicit water, and not in the current protein milieu[8]. Such ‘as-the-crow-flies’ overestimates of energy barriers vis-à-vis relatively (but not completely) ‘straight’ paths (akin to those shown in Supplementary Fig. 8a and traversed in b) is around double. This would suggest a γ-P–O cleavage energy barrier of ~8–10 kcal mol⁻¹. This substantially larger value compared with proton abstraction indicates that phosphate cleavage is rate-limiting. This is consistent with the proposed stabilization by Asn72 and/or Arg9 of the pentavent intermediate (Fig. 2a), and MDS evidence pointing to Arg9-mediated stabilization.

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Author contributions
D.L. designed experiments, and produced, purified, crystallized and functionally characterized the protein and its variants, collected and processed synchrotron diffraction data, refined and analysed the structures and wrote the manuscript. N.H. assisted with the production and characterization of mutants. A.K. helped with the catalytic mechanism. L.V. contributed toward data processing and structure solution and analysis. J.A.L, D.A. and K.D. advised on data analysis. P.J.S. and M.S.P.S were responsible for MDS. S.B. and N.J.E. performed DFT. V.C. developed the LCP-SFX approach and supervised the XFEL beamline at LCLS (L672). U.W. conceived, designed and developed the LCP injector. R.F., S.B., M.M., G.J.W. and M.M.S. set-up the XFEL experiment, beamline, controls and data acquisition, operated the CXI beamline and performed the data collection. W.L. helped with XFEL sample preparation and participated in LCP-SFX data collection. P.F., R.F., S.B., N.A.Z., I.G., C.K., K.R., C.G, A.B. and O.Y. participated in LCP-SFX data collection, analysis and processing. H.N.C. supervised software development and data processing. M.C. was responsible for the overall project strategy and management and wrote the manuscript. Edits were provided by V.C., K.D., P.F., N.J.E., and P.J.S.

Additional information
Accession codes: Atomic coordinates and structure factors for Δ4-99MAG-ACP, Δ4-99MAG, Δ7-79MAG-FEL and Δ7-79MAG are deposited in the Protein Data Bank under accession codes 4UXX, 4UXW, 4UYO and 4UXZ, respectively.

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