Cryo-EM structures of PI3Kα reveal conformational changes during inhibition and activation

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Phosphoinositide 3-kinases (PI3Ks) are lipid kinases essential for growth and metabolism. Their aberrant activation is associated with many types of cancers. Here we used single-particle cryo-electron microscopy (cryo-EM) to determine three distinct conformations of full-length PI3Kα (p110α-p85α): the unliganded heterodimer PI3Kα, PI3Kα bound to the p110α-specific inhibitor BYL-719, and PI3Kα exposed to an activating phosphopeptide. The cryo-EM structures of unbound and of BYL-719-bound PI3Kα are in general accord with published crystal structures. Local deviations are presented and discussed. BYL-719 stabilizes the structure of PI3Kα, but three regions of low-resolution extra density remain and are provisionally assigned to the cSH2, BH, and SH3 domains of p85. One of the extra density regions is in contact with the kinase domain blocking access to the catalytic site. This conformational change indicates that the effects of BYL-719 on PI3Kα activity extend beyond competition with adenosine triphosphate (ATP). In unliganded PI3Kα, the DFG motif occurs in the “in” and “out” positions. In BYL-719-bound PI3Kα, only the DFG-in position, corresponding to the active conformation of the kinase, was observed. The phosphopeptide-bound structure of PI3Kα is composed of a stable core resolved at 3.8 Å. It contains all p110α domains except the adaptor-binding domain (ABD). The p85α domains, linked to the core through the ABD, are no longer resolved, implying that the phosphopeptide activates PI3Kα by fully releasing the nSH2 domain from binding to p110α. The structures presented here show the basal form of the full-length PI3Kα dimer and document conformational changes related to the activated and inhibited states.

Phosphoinositide 3-kinase (PI3K) | activation | inhibition | activity-dependent conformational changes

PI3Ks (phosphoinositide 3-kinases) are lipid kinases that phosphorylate the 3-hydroxyl group of the inositol ring of phosphatidylinositol (1). They are regulators of numerous important cellular functions, including cell replication, movement, and metabolism (2-5). They are also significant factors in several human diseases, notably in cancer (7).

PI3Ks are grouped into three classes (8). Class I encompasses four isoforms that have shared as well as nonredundant functions (9-11). These are dimeric enzymes consisting of a catalytic subunit, p110, bound to a regulatory subunit. The class is further subdivided into IA and IB. Class IA encompasses the p110α, β, and δ isoforms. They are predominantly mediating signaling by RTKs (receptor tyrosine kinases). Class IB consists of a single isoform, p110γ, which is activated exclusively by G protein–coupled receptors.

The p110α isoform and its regulatory subunit p85α have special importance in cancer (12). Human solid tumors often harbor mutations in p110α, and about 80% of these mutations are located in three hotspots (13). These are single-nucleotide substitutions that cause a gain of function in the enzyme (13-17) and are oncogenic in cell culture and in the animal (18-20). p110α is therefore a promising therapeutic target for cancer.

Numerous PI3K inhibitors have been generated and disclosed (21, 22). Early compounds were “pan-specific,” affecting all four isoforms of class I PI3K. More recently, isoform-specific or isoform-selective inhibitors have been identified. All currently US Food and Drug Administration–approved PI3K inhibitors are either isoform-specific or show distinct preference for one or two isoforms (22). One of the most recently approved compounds is BYL-719 (alpelisib), which is p110α-specific and is the only PI3K inhibitor currently in clinical use for solid tumors (23-25).

PI3Ks have long been in the crosshairs of structural biologists. The first to be crystallized and analyzed was p110α and its regulatory subunit p85α (9). But for more than a decade, advances in the field remained limited, and the PI3K structure was largely unknown. The structures presented here show the basal form of the full-length PI3Kα dimer and document conformational changes related to the activated and inhibited states.

Phosphoinositide 3-kinases (PI3Ks) are of critical importance in cell signaling and can function as drivers of disease. Information on the PI3K structure is essential for an understanding of the function of these proteins and for the identification of specific and effective small-molecule inhibitors. Here we present a single-particle cryo-electron microscopy (cryo-EM) analysis of PI3Kα, the dimer consisting of the p110α catalytic subunit bound to the p85α regulatory subunit. We investigated three conformational states of PI3Kα: the unbound dimer, the dimer bound to the isoform-specific inhibitor BYL-719, and the dimer associated with an activating phosphopeptide. Each of these conformational states provides insights into PI3K structure and function.

Significance

Phosphoinositide 3-kinases (PI3Ks) are of critical importance in cell signaling and can function as drivers of disease. Information on the PI3K structure is essential for an understanding of the function of these proteins and for the identification of specific and effective small-molecule inhibitors. Here, we present a single-particle cryo-electron microscopy (cryo-EM) analysis of PI3Kα, the dimer consisting of the p110α catalytic subunit bound to the p85α regulatory subunit. We investigated three conformational states of PI3Kα: the unbound dimer, the dimer bound to the isoform-specific inhibitor BYL-719, and the dimer associated with an activating phosphopeptide. Each of these conformational states provides insights into PI3K structure and function.

Author contributions: S.Y., M.-W.W., and P.K.V. initiated the project; S.Y., J.R.H., M.-W.W., and P.K.V. designed research; S.Y. established the purification protocol for the PI3Kα complexes, shared this skill with X.L. and performed bioassays; X.L. and Y.X. performed research, purified the PI3Kα complexes, screened specimen and prepared the final samples for negative staining and cryo-EM data collection towards the structures; J.R.H., X.Z., H.Z., Q.Z., T.X., and Y.X. analyzed data and performed map calculation; J.R.H. conducted structure analysis and built the models; S.Y. and J.R.H. prepared the figures; S.Y., D.Y., M.-W.W., and P.K.V. supervised the studies; and X.L., S.Y., J.R.H., M.-W.W., and P.K.V. wrote the paper with input from all co-authors.

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regulatory subunit (26). This study also defined the functional domains of class I PI3Ks. The crystallographic analysis of the p110α–p85α heterodimer was achieved in 2007 (27). By now, more than 40 p110α structures have been deposited in the Protein Data Bank (PDB; https://www.rcsb.org) (28).

Small-molecule inhibitors have played a significant role in the investigation of the PI3K structure, because they often stabilize the molecule and allow images with higher resolution (21). We took advantage of this stabilizing effect of small molecular ligands by analyzing PI3Kα (p110α–p85α) bound to the p110α-specific inhibitor BYL-719. These investigations address questions of isomor specificity of inhibitors and present data on the position of the DFG domain located in the activation loop of PI3K. They also provide data suggesting that the inhibitory effect of BYL-719 is not confined to competition with adenosine triphosphate (ATP). Our studies further extend to the interaction of PI3Kα with an RTK-derived phosphopeptide, representing signal-mediated activation of PI3K.

Results

High-Resolution Cryo-EM Structure of PI3Kα. PI3Kα was coexpressed in High Five insect cells from a baculovirus dual-expression vector coding for p110α and p85α. The PI3Kα complex was purified and confirmed by sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE) for purity (Fig. 1A). Native gel electrophoresis showed that the complex is intact (Fig. 1B). The purified PI3Kα exhibits lipid kinase activity as determined by a membrane capture assay (29).

Initial attempts to determine the full-length PI3Kα structure were hampered by the aggregation of particles near the edges of holes. These problems were overcome through trial and error of blotting and freezing conditions to obtain particles suitable for analysis. Similar problems have been seen before with the cryo-EM structure of the Ragulator complex (30).

The cryo-EM structure of PI3Kα was determined by single-particle reconstruction. Vitrified complexes were imaged by cryo-EM. After sorting by constitutive two-dimensional (2D) and three-dimensional (3D) classifications, 3D consensus density maps were reconstructed with global resolutions of 2.65 Å for PI3Kα (Fig. 1C and D and SI Appendix, Table S1) (31). However, local resolution maps showed that different regions of the electron density map have varying resolutions, ranging from 2.0 to 9.3 Å (Fig. 1E). The cryo-EM map allowed us to build an unambiguous model for most regions of the complex, including all domains of p110α and the nSH2 and iSH2 domains of p85α, but not the remaining domains of p85α, which were less well resolved (32). The unmodeled domains are present and observable in the electron density maps (Fig. 1E), but the low resolution of these domains hampers unambiguous assignment (SI Appendix, Fig. S1).

The cryo-EM structure of PI3Kα shares similar features with structures previously determined with other methods (27, 33) (Fig. 1D). The PI3Kα complex exists as an asymmetric heterodimer containing one copy each of p110α and p85α. Comparing the cryo-EM structure of PI3Kα with the crystal structure of p110α bound to the truncated p85α nSH2 (PDB ID code 4OVU) (see SI Appendix, Table S2 for additional details on PDB ID codes) (33), we found an extensive overall concordance of the two (Fig. 2).

Within the catalytic core of p110α (residues 109 to 1050), there is only a 1.201-Å rmsd as compared with the reported crystal structure (Fig. 2A). The ABD domain of p110α, the iSH2 of p85α, and the nSH2 of p85α are significantly displaced, leading to a relatively high, 3.425-Å rmsd for these regions. Overall, the cryo-EM structure as compared with the crystal structure has an rmsd of 2.131 Å.

When the cryo-EM and crystal structure data (PDB ID code 4OVU) are aligned using the catalytic core of p110α, there is a rotation of the iSH2 domain around residue 538 of p85α, leading to a displacement of both termini of the iSH2. Additionally, there is a deflection in the loop region of the iSH2 (p85α residues 498 to 523), resulting in a displacement of 3.30 Å (Fig. 2B). However, when the cryo-EM structure is compared with the crystal structure of the ABD of p110α and iSH2 of p85α (PDB ID code 2IVY) or another structure of PI3Kα (PDB ID code 2RD0), these displacements are not present. We suggest that there is some conformational flexibility of the iSH2 and ABD domains with respect to the catalytic core of PI3K, and motion occurs via a flexible hinge at G106.

The structure of nSH2 of p85α has been investigated many times by both crystallography (34) and NMR (35, 36) and is complex with p110α (15, 33). Within this domain, the most significant difference between the existing crystal structures and the cryo-EM structure is in the conformation of the BC loop (p110α residues 361 to 366) (Fig. 2D). Multiple conformations of this loop have been previously observed, both in isolated structures of this domain and in complex with p110α (34–36). In cryo-EM, we observed the BC loop positioned away from the helical domain and toward the kinase domain.

In addition to the position of this loop, the nSH2 domain is rotated relative to the catalytic core of PI3K. This difference in positioning of the nSH2 has been previously observed (Fig. 2E) (33). The position in the cryo-EM structure more closely aligns with the position seen in the p110α H1047R mutant structures (15).

In the catalytic domain, the β-sheets forming the N-terminal catalytic lobe (N-lobe) are closer to the active site than observed in the crystal structure (Fig. 2C). This is most pronounced in the β3K–β4K turn of p110α residues 774 to 776 with A775 being displaced by 3.1 Å and in the β4K–β5K turn with I788 displaced by 3.8 Å at the Ca positions. These differences might be due to steric interactions with the N-terminal affinity tag present in the crystal structure.

The Cryo-EM Structure of PI3Kα Bound to the p110α-Specific Inhibitor BYL-719. In an attempt to stabilize the PI3Kα complex, we investigated PI3K inhibitors. The binding of BYL-719 to PI3Kα has been previously studied by crystallography (PDB ID code 4IPS) (25). As the resolution of the BYL-719–bound crystal structure was significantly higher than that of the ligand-free complex, we used this compound to stabilize the PI3Kα complex for cryo-EM analysis.

BYL-719 is a potent and isoform-specific p110α inhibitor. As previously reported, BYL-719 almost completely abolishes the lipid kinase activity of PI3Kα (Fig. 3A) (37). Using a similar procedure as with the ligand-free complex, 3D consensus density maps were reconstructed with a global resolution of 2.93 Å for BYL-719–bound PI3Kα (Fig. 3B and C and SI Appendix, Table S1) (38). A molecular model of the BYL-719–bound PI3Kα cryo-EM structure was generated, and two orientations are shown in Fig. 3C (39).

Comparison of the cryo-EM structure with the previously reported crystal structure shows many of the same differences that have been identified with the ligand-free structure resulting in an overall rmsd of 2.197 Å (Fig. 3E). Again, the catalytic core of PI3K remains relatively unchanged with an rmsd of 1.690 Å. The nSH2 and ABD domains have a similar rotation as in the ligand-free structure and lead to an rmsd of 3.631 Å. However, there is no rotation of the nSH2, and the BC loop is unobserved in the crystallography data. Similar to the unbound PI3Kα, BYL-719–bound PI3Kα also shows a displacement of the N-lobe of the kinase domain.

BYL-719 in the cryo-EM structure is in a conformation that is different from the one seen in the crystal structure. However, key interactions designed to confer p110α specificity are retained (Fig. 3F). Specifically, hydrogen bonds between the
amide group and Q859 and a hydrogen bond with the carbonyl oxygen of V851 are preserved. However, the position of the CF$_3$ group is different in the two structures. While the crystal structure has the CF$_3$ group positioned in a hydrophobic pocket created by P778, I800, K802, and I848, we observed the density of this group rotated by 120°. Instead of a CF–HN hydrogen bond with K802, we observed a CF–HO hydrogen bond with S774. Additionally, the ordered water molecules seen with Y705 and the pyridine nitrogen are not observed by cryo-EM.

**Extra Density.** In the confidence maps for both the unbound PI3Kα and BYL-719–bound structures, there are areas of low-resolution extra density (Figs. 3D and 4A). We have divided the extra density into three regions: ED1, ED2, and ED3, shown in Fig. 4B and SI Appendix, Fig. S1. Upon binding of BYL-719, the ED2 region becomes stabilized with local resolution improving from 8 to 5 Å, and the ED3 region can be observed.

There are three domains of p85 left unmodeled in the cryo-EM data: the SH3, BH, and cSH2 domains of p85α. Using the 1% false discovery rate (FDR) confidence map, we have provisionally assigned ED1, ED2, and ED3 to the cSH2, BH, and SH3 domains, respectively (Fig. 4B). The cSH2 domain of p85β in complex with p110β has been previously solved (40). While the cSH2 domain cannot interact with p110α in the same
manner as with p110β, the volume of ED1 is consistent with the cSH2 domain and the position observed in p110β-p85β. ED2 was assigned to the BH domain based upon its larger size of 180 amino acids and ED3 to the SH3 domain, which is composed of 80 amino acids. However, these domains cannot be unambiguously assigned. Linker regions between the nSH2 and BH domains as well as the iSH2 and cSH2 domains are long enough that either could be ED1 or ED2. Additionally, the linker between the BH and SH3 domains is long enough that it could span the distance from ED1 or ED2 to ED3.

The extra density in the BYL-719–bound structure is in contact with the kinase domain and the iSH2 domain as well as with BYL-719 (Fig. 4C). Specifically, ED2 is in contact with both the N- and C-lobes of the kinase domain, effectively blocking access to the catalytic site (Fig. 4C). Contact with BYL-719 is through the CF3 group and may be responsible for the positional difference between the cryo-EM and the crystal structures. The contact with the loops of the N-lobe in the kinase domain are displaced at both A775 and I778. (A) The nSH2 domain is rotated relative to the helical domain by 14°.

**Three-Dimensional Variability Analysis.** In both PI3Kα structures, there is a large range of local resolutions, particularly around the ED1 to ED3 regions. Three-dimensional variability analysis (3DVA) (41) was performed on both datasets using cryoSPARC.

3DVA demonstrates that ED regions in the unbound PI3Kα structure have low local resolution because of large variations in their position and rotation (SI Appendix, Fig. S2A). Additionally, there are variations in the iSH2, RBD, nSH2, and kinase domains, which are detailed in SI Appendix, Figs. S2 and S3. The 3DVA of unbound PI3Kα includes a component where the DFG motif of the kinase domain rearranges (Fig. 5). In both the unbound PI3Kα and BYL-719–bound structures as well as previous structural studies, the DFG motif is in the catalytically active DFG-in conformation (Fig. S4). However, the 3DVA on the unbound structure (component 3) shows that the DFG motif can also adopt a DFG-out conformation (Fig. 5B). This conformation has not been seen in 3DVA of the three components analyzed for the BYL-719–bound structure. In the final electron density map of unbound PI3Kα, there is indeed a suggestion that the DFG-out conformation could be present in a minority of the particles (Fig. 5A).

**The Cryo-EM Structure of Phosphopeptide-Activated PI3Kα.** Activated PI3K was prepared by combining PI3Kα with a phosphopeptide derived from PDGFRβ (platelet-derived growth factor receptor beta) following previously published methods (14). The phosphopeptide increased PI3Kα lipid kinase activity (Fig. 6A), as previously reported (42). The cryo-EM analysis of particles of activated PI3Kα showed a dramatic departure from the 2D class averages observed for other PI3Kα complexes (SI Appendix, Fig. S4), but the particles identified by cryo-EM were not suitable for 3D reconstruction.

However, one 2D class average was obtained with high resolution, and this class is shown alongside similar orientations seen in the PI3Kα and BYL-719–bound data (Fig. 6C, with models for comparison in Fig. 6D). The observed 2D class average closely matches the electron density projection of the catalytic core of p110α without an ABD (PDB ID code 6OAC). Although this orientation is highest in abundance, multiple orientations of the activated complex were observed (SI Appendix, Fig. S4). The data do not indicate a preferred orientation, as the number of particles is well-distributed between different orientations (43, 44). The other orientations are present within the dataset but, because a large proportion of the complex is conformationally flexible with multiple degrees of freedom, these classes have low resolution. In one orientation, the ABD and p85 domains are in a conformation where they do not overlap with the catalytic core, allowing for this particular class to have high resolution. We suggest that this is through the flexibility of the ABD–RBD linker near G106. In theoretical models of possible orientations of the ABD and iSH2 domains, G106 can adopt multiple conformations without clashing or overlapping with the catalytic core while remaining within Ramachandran-allowed regions for glycine. In activated PI3Kα, electron density for the nSH2 domain is not observed and electron density for the ABD and iSH2 domains is also absent. Yet native gel electrophoresis (Fig. 6B) shows that the PI3Kα complex does not dissociate upon binding to the phosphopeptide. These observations are in accord with the interpretation that when activated, the nSH2, iSH2, and ABD domains no longer interact with the catalytic core of p110α (residues 107 to 1068). The iSH2 domain remains bound to the ABD domain as demonstrated by native gel electrophoresis, and the ABD domain can adopt a range of conformations upon phosphopeptide binding. The data are compatible with a hypothetical model of phosphopeptide-bound PI3Kα in which the catalytic core of p110α, consisting of the RBD, C2, helical, and kinase domains, forms a rigid group, whereas the ABD and all p85e domains are flexible, relative to the rigid catalytic core of p110α.

**Discussion.** Our knowledge of the PI3K structure rests on groundbreaking work conducted by X-ray crystallography and hydrogen/deuterium exchange mass spectrometry (HDEx-MS) (14, 25, 27, 42, 45). Isolated domains of PI3K have also been investigated by NMR (35, 46, 47). Here we take advantage of recent developments in single-particle analysis in cryo-EM to present structures of PI3Kα in different functional conformations. The main technical challenge in the cryo-EM analysis of PI3Kα stems from the inherent structural flexibility of several domains, notably the SH3, BH, iSH2, and cSH2 domains of p85α as well as the ABD domain of p110α. Despite these obstacles, cryo-EM analysis
has succeeded in generating data and insights that have not been available by other technologies.

There is extensive concordance between the cryo-EM and crystallography structures. Local deviations have been identified, characterized, and analyzed in detail. One domain that deserves particular attention is the DFG motif. The position of the aspartate of the DFG motif determines the active or inactive state of the kinase. Additionally, small-molecule inhibitors can bind to either the DFG-in or DFG-out conformation (48, 49). Our data show that in unbound PI3Kα, DFG is predominantly in the active “in” position, but 3DVA identified a component that shows DFG in the “out” position in a minority of the particles. In the BYL-719–bound structure only the DFG-in conformation, corresponding to the active form of PI3K, has been observed.

The cryo-EM analysis of PI3Kα was facilitated by binding to the isoform-specific inhibitor BYL-719. A comparison of the cryo-EM and crystal structures of the bound BYL-719 shows two different conformations, but the interactions that make BYL-719 isoform-specific are preserved and identical in both structures. The use of BYL-719 raises the general issue of the molecular basis of isoform specificity of PI3K inhibitors. This important topic has been reviewed recently (21), emphasizing that isoform specificity is determined by multiple factors that control inhibitor–protein interactions, and that there is no single, universal rule for achieving isoform specificity.

BYL-719 stabilizes areas of extra density, allowing the provisional assignment of the cSH2, BH, and SH3 domains to specific portions of this extra density. In the BYL-719–bound form, SH3, BH, and cSH2 are stably positioned near the...
among the possible positions that could be transiently taken up. We speculate that blocking access to the catalytic site is the iSH2 domain. The extra density also interacts directly with BYL-719.

PI3K

The iSH2 domain is known to be involved in the catalytic core of p110α. Many types of cancer carry the gain-of-function mutation H1047R at this site. The mutation leads to the activation of the enzyme through an increased affinity for lipids (15). In contrast to wild-type PI3Kα (33), the C terminus of p110α is flipped in the crystal structure of the H1047R mutant (15, 27). This places the C terminus in a position where it would clash with the ED2 domain. We speculate that the H1047R mutation repositions the ED2 domain, and this may contribute to the activating effect of H1047R.

For our studies with activated PI3Kα, we offer a hypothetical interpretation that is consonant with the available data. The analysis of these data suggests that all p85α domains and the ABD of p110α adopt flexible conformations relative to the p110α catalytic core. Native gel electrophoresis demonstrated that under these conditions p85α and p110α remain bound. Although a 3D reconstruction of the phosphopeptide-activated PI3Kα complex could not be achieved, a high-resolution 2D class average of the p110α catalytic core can be observed and interpreted. Within this class, the nSH2 domain is no longer observed, which is consistent with the proposed mechanism of activation and HDX-MS data of full-length PI3Kα (14). Importantly, this projection also lacks the ABD domain and the iSH2 domains (Fig. 6C). This observation indicates that in addition to the nSH2 and iSH2 dissociating from p110α to bind to an activated RTK, the iSH2 domain dissociates as well. Thus, the catalytic core of p110α is bound via the ABD–iSH2 interaction to an extended structure containing the nSH2 and cSH2 domains and possibly including the SH3 and BH domains. The connection of the ABD–iSH2 complex to the catalytic core functions like a hinge, and we suggest that G106 can act as this hinge. In the long structure extending from the catalytic core, the ABD–iSH2 part is a relatively rigid fiber with flexible globular domains attached to each end by flexible linkers. This long structure could facilitate the conformational flexibility required to reach the phosphorylated tyrosines in the intracellular domain of diverse RTKs (50–52). These tyrosines are located near the membrane but are not necessarily directly adjacent to the membrane. Mobility of the ABD and iSH2 allows for additional conformational flexibility to simultaneously bind RTKs and access phosphatidylinositol (4, 5) bisphosphate (PIP2) in the membrane.
Previously reported structural data for phosphopeptide-activated PI3Kα come from HDX-MS experiments (14). In addition to the expected exposure of helical domain amino acids normally bound by the nSH2 domain, activation by the phosphopeptide results in increased solvent exchange of the ABD-RBD linker and C2 domains and, upon lipid binding, the iSH2 domain as well. These observations are consistent with our model of the activated form of PI3Kα. The profound structural changes induced in PI3Kα by physiological activation raise the question of whether similar changes are mediated by the gain-of-function mutations that show constitutive enzymatic activity, specifically the G106V mutation (53).

Our investigations of the structure of PI3Kα suggest some immediate goals for the cryo-EM analysis of PI3K. Other isoforms of class I PI3K would be rewarding targets, and while this manuscript was in revision, the cryo-EM structure of the single member of class IB of PI3K, PI3Kγ (p110γ-p101), was published (54). The flexibility of PI3K complexes makes the identification of ligands that can stabilize parts or all of the mobile and flexible domains an urgent task. Also on the horizon is the cryo-EM analysis of cancer-specific mutations which would generate further insights into mutant mechanisms of action and could guide the development of mutant-specific inhibitors of therapeutic value. A more generic problem is availability of instrument access. Numerous technical problems still require a time-consuming trial-and-error approach that relies on teamwork and exceptional institutional resources.

However, rapid advances in technology may provide novel solutions.

**Materials and Methods**

**Protein Expression and Purification.** A pFastBac dual vector expressing both p110α and p85α was kindly provided by Peter Shepherd, University of Auckland, Auckland, New Zealand. The pFastBac dual vector was recombined with the parent bacmid in DH10Bac Escherichia coli competent cells to form an expression bacmid containing both p110α and p85α. The expression bacmid was then transfected into Sf9 insect cells for production of recombinant baculovirus. Baculovirus-expressing p110α with 6x His tag and p85α were amplified once in Sf9 insect cells, and then 10 mL amplified baculoviruses were used to infect 1 L BTI-Tn-5B1-4 (High Five) cells at a density of 1.5 × 10^6 cells per milliliter for protein production. Forty-eight hours after infection, High Five cells were harvested by centrifugation at 1,000 × g for 10 min, washed with ice-cold phosphate-buffered saline, and then pelleted by centrifugation at 2,000 × g for 10 min. The pellets from 1 L of High Five cell culture were lysed by Dounce homogenization in 60 mL buffer A (20 mM Tris, pH 8.0, 400 mM NaCl, 5% [vol/vol] glycerol, 0.5% [vol/vol] Triton X-100, and 2 mM β-mercaptoethanol), with ethylenediaminetetraacetate-free protease inhibitor mixture [TargetMol] on ice, followed by centrifugation for 1 h at 140,000 × g at 4°C. The supernatant was then passed through a 0.45-μm cellULOPLCE1 blames acetate membrane filter (Merck Millipore) and incubated with Ni-NTA resin (Cytiva) for 2 h at 4°C. The resin was loaded onto a gravity column (Sangon Biotech), washed with buffer B (20 mM Tris HCl, pH 8.0, 400 mM NaCl, 50 mM imidazole, 5% [vol/vol] glycerol, and 2 mM β-mercaptoethanol), and eluted with buffer C (20 mM Tris HCl, pH 8.0, 100 mM NaCl, 300 mM imidazole, 5% [vol/vol] glycerol, and 2 mM β-mercaptoethanol). The eluate was then loaded onto a 1-mL heparin HP column (Cytiva), washed with buffer D...
The purified PI3κκα complex was combined with [γ-^32P]ATP in a kinase buffer with phosphoinositides prepared into vesicles as described (29). Phosphorylated lipids were measured by spotting 4 μL of reaction mixture onto nitrocellulose membrane and washing away unincorporated ATP. The phosphorylated phosphoinositol was measured by autoradiography using a phosphor imager.

### Cryo-EM Data Acquisition and Image Processing.

#### Sample preparation.

The unbound PI3κκα was used as is from purification. The purified PI3κκα complex (2.5 μL) at a concentration of 1 mg/mL was spotted onto grids as described below. The BYL-719-bound PI3κκα complex was prepared by incubating the PI3κκα complex (1 mg/mL) with 1 mM BYL-719 stock (MedChemExpress) at a molar ratio of 1:10 at 4 °C for 30 min. The phosphopeptide-bound PI3κκα complex was prepared by incubating the PI3κκα complex (1 mg/mL) and the phosphopeptide stock (1 mg/mL) at a molar ratio of 1:15 at room temperature for 5 min before vitrification of the sample. Each complex was added to glow-discharged holey carbon grids (Quantifoil R1.2/1.3, 300 mesh) and subsequently vitrified by plunging into liquid ethane using a Vitrobot Mark IV (ThermoFisher Scientific).

#### Data acquisition.

Automatic data collection was performed on a Titan Krios equipped with a Gatan K3 Summit direct electron detector. The microscope was operated at 300-kV accelerating voltage, at a nominal magnification of 46,685 × in counting mode, corresponding to a pixel size of 1.04 Å. Acquisition parameters are detailed in SI Appendix, Table S1.

#### Image processing.

Dose-fractionated image stacks were subjected to beam-induced motion correction and dose weighting using MotionCor2.1 (55). A subset was then subjected to 3D refinement and Bayesian polishing with a pixel size of 1.04 Å. Local resolution was determined using the Bsoft package (56).

#### Autopicking.

Autopicking yielded particle projections that were subjected to reference-free 2D classification to discard false-positive particles or particles categorized in poorly defined classes, producing projections for further processing. This subset of particle projections was subjected to consecutive rounds of 3D classification with a pixel size of 2.09 Å followed by a round of maximum likelihood-based 3D classifications with a pixel size of 2.09 Å, resulting in one well-defined subset with projections. Further 3D classifications with a mask on the complex produced one good subset. These were subsequently subjected to another round of 3D classifications with a mask on the complex. A selected subset was then subjected to 3D refinement and Bayesian polishing with a pixel size of 1.04 Å. Local resolution was determined using the Bsoft package (v3.0.1) and RELION-3.0-beta2.

#### Figures.

Figures were produced using Chimera (v1.14 build 42094) (61) and ChimeraX (v1.2 2021-05-24) (71). Protein alignments between models were produced using Chimera MatchMaker (72) using the catalytic core of p110α (residues 108 to 1047) instead of PI3κκα.

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