Autoinhibition of Bruton’s tyrosine kinase (Btk) and activation by soluble inositol hexakisphosphate

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Abstract

Bruton’s tyrosine kinase (Btk), a Tec-family tyrosine kinase, is essential for B-cell function. We present crystallographic and biochemical analyses of Btk, which together reveal molecular details of its autoinhibition and activation. Autoinhibited Btk adopts a compact conformation like that of inactive c-Src and c-Abl. A lipid-binding PH-TH module, unique to Tec kinases, acts in conjunction with the SH2 and SH3 domains to stabilize the inactive conformation. In addition to the expected activation of Btk by membranes containing phosphatidylinositol triphosphate (PIP$_3$), we found that inositol hexakisphosphate (IP$_6$), a soluble signaling molecule found in both animal and plant cells, also activates Btk. This activation is a consequence of a transient PH-TH dimerization induced by IP$_6$, which promotes transphosphorylation of the kinase domains. Sequence comparisons with other Tec-family kinases suggest that activation by IP$_6$ is unique to Btk.
Introduction

Activation of B- and T- lymphocytes relies on a chain of tyrosine phosphorylation events initiated by B-cell and T-cell receptors, respectively (Cambier et al. 1993; Weiss & Littman 1994; Kurosaki 2011). These phosphorylation-dependent signals are generated by three kinds of non-receptor tyrosine kinases, linked to the receptor by membrane co-localization (Kurosaki 2011; Dal Porto et al. 2004). Src-family kinases (principally Lyn and Fyn in B-cells, Lck and Fyn in T-cells), resident at the plasma membrane, respond first to receptor activation. They phosphorylate pairs of tyrosine residues in ITAM motifs in the cytoplasmic tails of activated receptors (Chu et al. 1998; Mócsai et al. 2010; Wang et al. 2010), which in turn recruit the tandem-SH2 domain tyrosine kinases Syk (in B-cells) and ZAP-70 (in T-cells). Membrane localization of Syk or ZAP-70 leads to the recruitment and activation of Tec-family tyrosine kinases, Btk (in B-cells) and Itk (in T-cells) (Andreotti et al. 2010).

Btk is present in all hematopoietic cells, except for T-cells and natural killer cells (Tsukada et al. 1993; Lindvall et al. 2005). It is also part of several other receptor-mediated signaling networks, including those initiated by Toll-like receptors (Jefferies et al. 2003), Fc receptors (Kawakami et al. 1994), and G-protein coupled receptors (Tsukada et al. 1994; Bence et al. 1997). Btk is critical for the generation of calcium flux in response to receptor activation, because it activates phospholipase C-γ2 (PLC-γ2), which produces the second messengers diacylglycerol (DAG) and inositol (1,4,5)-trisphosphate (IP₃). Btk dysfunction is the cause of X-linked agammaglobulinemia, a severe disease of primary immunodeficiency (Tsukada et al. 1993).

The Tec family kinases have a lipid-interaction module attached to the N terminus of a Src-like module (i.e., concatenation of an SH3 domain, an SH2 domain and a tyrosine kinase
domain) (Figure 1A). Like the non-receptor tyrosine kinase c-Abl, the Tec kinases lack a C-terminal tyrosine phosphorylation site that in Src kinases engages the SH2 domain in an autoinhibitory interaction (Sicheri et al. 1997; Xu et al. 1997; Nagar et al. 2003). The lipid-interaction module of the Tec kinases is an extended plekstrin homology (PH) domain fused to a Tec homology (TH) domain (Hyvönen & Saraste 1997). The PH-TH binds PIP₃ in membranes, or the soluble head group of PIP₃, inositol tetrakisphosphate (IP₄), in solution. A flexible linker connects the PH-TH module to the Src-like module; its length varies from 26 residues in Itk, the Tec family kinase in T-cells, to 45 residues in Btk.

Tyrosine phosphorylation controls Btk activity. B-cell activation leads to rapid phosphorylation of Tyr 551 in the kinase domain and Tyr 223 in the SH3 domain of Btk (Rawlings et al. 1996; Park et al. 1996). Tyr 551 corresponds to the canonical phosphorylation site in the activation loop of tyrosine kinases; as expected, its phosphorylation increases Btk catalytic activity (Rawlings et al. 1996). Phosphorylation of Tyr 223 has been reported to have no apparent influence on Btk catalytic activity (Park et al. 1996; Middendorp et al. 2003). It lies within the peptide-binding groove of the SH3 domain, and its phosphorylation alters SH3 domain selectivity among binding partners (Morrogh et al. 1999; Middendorp et al. 2003).

Binding of a Tec-kinase PH domain to the membrane lipid PIP₃, produced as a consequence of receptor activation, provides a simple control mechanism for switching on the kinase. Membrane recruitment triggers trans-autophosphorylation, through increased local concentration, or trans-phosphorylation by membrane-bound Src-family kinases, thereby activating the Tec kinase (Rawlings et al. 1996).

We present here an analysis of two crystal structures of Btk, which jointly provide molecular details of its autoinhibition. One structure is of the Src-like module, in an assembled
and inactive conformation, resembling those of inactive c-Abl and of the Src kinases c-Src and Hck (Xu et al. 1997; Sicheri et al. 1997; Nagar et al. 2003). The other is of a Btk construct that lacks the SH3 and SH2 modules and that links the PH-TH module directly to the kinase domain. We find that the PH-TH module stabilizes the inactive conformation of the kinase by interacting with the N lobe. The inactive conformation of Btk is thus a compact one, in which the PH-TH module cooperates with the SH2 and SH3 domains to maintain an inactive structure. Several structures of isolated domains of Btk have been determined previously, as summarized in Table 4, which also lists structures of fragments of other Tec family kinases.

We also report a set of findings that reveal an unanticipated role for the PH-TH module in activating the kinase domain. We find that inositol hexakisphosphate (IP₆), a soluble signaling molecule ubiquitous in animal and plant cells (Vallejo et al. 1987; Szwergold et al. 1987), can activate Btk strongly. We have determined a crystal structure of the PH-TH module of Btk bound to IP₆ and identified a previously undetected binding site on the PH-TH module, distinct from the canonical PIP₃/IP₄ site. The new site is critical for activation by IP₆.

We show that Btk activation by IP₆ is a consequence of a transient, IP₆-induced PH-TH dimerization, which promotes transphosphorylation of the kinase domains. The possibility that PH dimerization might be important for Btk activation was first proposed by Saraste and colleagues (Hyvönen & Saraste 1997), and we now show that the dimer interface identified previously is essential for IP₆-dependent Btk activation. IP₆-triggered transient dimerization of the PH domain may provide a membrane-independent mechanism for activating Btk in the nucleus or cytoplasm or may have a role in feedback stimulation of Btk that has been activated by PIP₃ at the membrane. The prior model for Btk activation involved only a PIP₃-dependent
membrane recruitment as the key step in switching on the kinase. The ability of IP₆ to activate Btk points to an as yet unappreciated complexity to the activation mechanism.
Results and Discussion

Structural analysis of Btk

We have determined four crystal structures as part of this work. Two of these structures help explain how Btk is autoinhibited. These are the structures of the Btk SH3-SH2-kinase module (determined at 2.6 Å resolution), and of the PHTH-kinase unit (determined at 1.7 Å resolution). The other two structures are of the isolated PH-TH module bound to IP$_6$ (determined at 2.3 Å resolution) and the isolated kinase domain with mutations in the activation loop (determined at 1.6 Å). The details of the crystallographic experiments are given in Tables 1 and 2.

Crystal structure of the Btk SH3-SH2-kinase module

We determined the structure of a construct of mouse Btk spanning residues 214 to 659, which starts just before the SH3 domain and ends at the C terminus of full-length Btk. The structure resembles those described previously for autoinhibited, Src-like, non-receptor tyrosine kinases, with the SH2-kinase linker sandwiched between the SH3 domain and the small domain of the kinase (the N lobe of the kinase) (Figure 1B). The crystal structure has two molecules of Btk in the asymmetric unit, intertwined as a domain-swapped dimer in which each SH2 domain has pieces from two molecules (Figure 1-supplement figure A). The domain-swapped SH2 domain packs against the large domain of the kinase (the C lobe of the kinase), but Btk lacks a Src-like C-terminal tail to stabilize this interaction.
The sequence identity between Btk and c-Abl is 42%, 31%, and 46% in the SH3, SH2 and kinase domains, respectively; for Btk and c-Src, it is 44%, 29%, and 40%. Superposition of the C lobes of the Btk and c-Abl kinase domains gives an approximately 20° relative rotation of their SH3 domains; superposition of the SH3 domains, a relative rotation of about 15° for their SH2 domains. Similar results are obtained for a comparison to c-Src. Coupling between the SH3 and SH2 domains may therefore be somewhat different in these kinases (Figure 1C).

The structure of the SH3 domain of Btk is nearly identical to those determined previously using NMR (the r.m.s. deviation for 53 Ca atoms is 0.9 Å) (Hansson et al. 1998; Tzeng et al. 2000), except for the expected differences seen at the N and C termini. When compared with the other autoinhibited SH3-SH2-kinase structures (c-Src, Hck, c-Abl), the conserved features of the SH3/linker interaction in Btk are the polyproline type-II helix conformation adopted by residues 383-387 in the SH2-kinase linker, including the proline at position 385, and the overall hydrophobic character of the peptide-binding pocket on the SH3 domain (Figure 1D). The tryptophan sidechain in the center of this pocket donates a hydrogen bond to a backbone carbonyl of the linker, as in most other SH3-ligand complexes. A cluster of hydrophobic sidechains extends from the binding pocket and the SH3 "RT loop", across the SH2-kinase linker segment, to the β2-β3 loop and strand β4 of the kinase (Figure 1D). Binding of exogenous protein ligands to the Btk SH3 domain would disassemble the Src-like inactive conformation, as seen for the Src family kinase Hck (Moarefi et al. 1997).

Diffuse electron density for the SH2 domain at every stage of our structure determination shows that this part of the polypeptide chain occupies a range of positions and adopts multiple local conformations. The domain itself has a marginally stable internal structure, as shown by the domain swap, and it is probably imperfectly locked in place. The
SH2 βA/αA loop lies between kinase-domain helices αB and αF. Density features for this loop are relatively sharp, and the rest of the domain may pivot somewhat about its contact with the C lobe.

The domain swap occurs in the same location within the SH2 domain as a domain swap seen in a Grb2 SH2-domain crystal structure (Schiering et al. 2000). The SH2 domain opens up at a position within the subsidiary β-sheet, so that helix B of one polypeptide chain packs against the main β-sheet of the other and vice versa (Figure 1-supplement figure B). This exchange occurs in solution for Btk, as only the dimeric species crystallizes. The similar domain swap seen in two SH2 domains not closely related in their specificity or functions suggest that other SH2 domains may also have the same bipartite character.

There is no evidence that a domain-swapped dimer (or, indeed, any stable dimer) is a functional state of intact Btk, and so it is likely that the assembled state involves a single polypeptide chain, rather than two inter-folded chains. The relatively high concentration of purified protein could have promoted dimer formation in our preparations. In the case of Grb2, the structure of the intact protein, a monomer in solution, shows no domain swap, and loss of contacts between the deleted N- and C-terminal SH3 domains can account for destabilization of the central SH2 when it is prepared as an isolated domain (Maignan et al. 1995). For Btk, the PH-TH module might stabilize the Src-like module.

The unphosphorylated activation loop of Btk folds into the mouth of the catalytic cleft, blocking part of the substrate-binding site, and helix αC lies at the margin of the N lobe, preventing formation of a Glu-Lys salt bridge conserved in all catalytically active protein kinases (Taylor & Kornev 2011) (Figure 1D). The activation loop of Btk is in a conformation seen in the structure of a Btk kinase domain-inhibitor complex (PDB: 3OCS; (Di Paolo et al.
2011), and in autoinhibited c-Src (the r.m.s. deviation from c-Src is 2.3 Å for the 24 α-carbons of the activation loop), but the cleft between the two lobes of the kinase is substantially more open in Btk than in c-Src. Helix αC is also displaced from the catalytic cleft, as in inactive c-Src or cyclin-dependent kinases. The segment of the activation loop that contacts the N lobe shifts with it, with respect to c-Src as a reference, by up to 3 Å, away from the C lobe.

A network of polar interactions present in the autoinhibited forms of c-Src and Hck is also present in the Btk conformation in our crystals (Figure 1-supplement figure C). Glu 445, which would buttress Lys 430 in the active conformation, interacts instead with Arg 544 and Tyr 545. Arg 544 stacks in turn on Arg 520, clamping the latter against the phenyl ring of Phe 574. Each of these arginines also appears to have an amino-aromatic interaction: Arg 544 with Phe 517 and Arg 520 with Phe 574. In the active conformation of Btk, the same two arginines are expected (based on the structure of the active Lck kinase domain, PDB: 3LCK; (Yamaguchi & Hendrickson 1996)) to engage in salt bridges with a phosphorylated Tyr 551. Activation thus entails a shuffling of salt-bridge and polar hydrogen-bond partners, described in the case of Src kinases as an “electrostatic switch” (Ozikirimli & Post 2006; Banavali & Roux 2009) (Figure 1-supplement figure C). In addition, the inactive conformation is stabilized by interactions made by the SH2-kinase linker. Leu 390 of the linker (corresponding to Trp 260 in c-Src) is sandwiched between residues Trp 421 and Tyr 461 in the kinase domain (Figure 1-supplement figure D). The hydrophobic packing between Trp 421 and Leu 390 would not be possible in the active conformation of the kinase domain.

The structure just described demonstrates that the SH3-SH2 domain in a Tec-family member can assemble into a regulatory clamp, just as it does in other Src-like kinases. In the crystal structures of c-Src, Hck, and c-Abl, a latch is present – either in the form of a
phosphorylated C-terminal tail (c-Src, Hck) or a myristoyl group bound in a pocket on the C
lobe of the kinase domain (c-Abl). In both cases, the latch secures the SH2-domain/C lobe
contact, precisely the interface that appears to be poorly formed in our Btk crystals. As we
show below, the latch in Btk is provided by the PH-TH module.

Structure of a PH-TH-kinase construct of Btk

We expressed a "PH-TH-kinase" construct of bovine Btk that contains the PH-TH
module connected to the kinase domain by the 13-residue SH2-kinase linker. We replaced six
residues in the activation loop of Btk with the corresponding ones in Itk, shown previously to
stabilize the activation loop of the Btk kinase domain (Joseph et al. 2013). This improved the
poorly ordered crystals obtained with the unmutated form, leading to a structure of the PH-TH-
kinase with a bound ATP-competitive inhibitor (CGI1746, (Di Paolo et al. 2011)) at 1.7 Å
resolution (Figure 2A).

The overall conformation of the kinase domain, as well as the specific conformation of
the activation loop, is very similar in the two structures we have determined (Figure 2-
supplement figure A). We also determined the structure of the isolated Btk kinase domain with
the six mutations described above and found by comparing it to wild-type Btk that the
mutations do not introduce major perturbations in the structure (Figure 2-supplement figure A).
Details of expression, purification, crystallization, and structure determination are in the
materials and methods.

The PH-TH module contacts the kinase N lobe. The interface differs substantially from
the one found in Akt1, another PH-domain regulated kinase (Wu et al. 2010) (Figure 2-
supplement figure B). In Akt1, the PH domain sits in front of the catalytic cleft of the kinase, blocking peptide substrate binding. In Btk, the PH-kinase interface does not directly occlude the active site. The interface combines a set of hydrophobic contacts with a hydrogen-bond network (Figure 2B). In particular, Tyr 134 at the C terminus of helix α2 of the PH-TH packs into a groove between Trp 395 at the linker-kinase junction and helix αC in the kinase N lobe. We expect these interactions to stabilize the inactive conformation of the kinase, by hindering the shift in helix αC that accompanies kinase activation. Experiments confirming these expectations are described below, in the section on autoinhibition.

A structural model for full-length Btk

We combined the two structures just described to build a model for full-length Btk (Figure 3A). Aligning the kinase domains in the two structures gives some local overlap between helix α2 in the PH-TH module and the β3/β4 loop in the SH3 domain, but otherwise compatible positions for the regulatory domains (Figure 3-supplement figure A). We therefore carried out molecular dynamics simulations of the two component structures and of the composite assembly, to determine whether energetically minor adjustments could resolve the remaining clashes. The trajectories ranged in length from 60 to 150 ns (see materials and methods for computational details).

In simulations of the SH3-SH2-kinase module, the initial and final conformations of the SH3 and kinase domains had r.m.s. deviations in Cα positions of only ~3.0 and ~2.0 Å respectively, after first aligning the C lobe of the kinase domain (Figure 3-supplement figure B). The SH2 domain shifted by ~6 Å, with a roughly 10° rotation that brought it closer to the C lobe
of the kinase domain and closer to its position in the inactive forms of c-Src, Hck, and c-Abl.

Some adjustment of the SH2 domain in these simulations was expected, as the domain swap in the crystals probably perturbed the domain to some extent, and the starting structure for the simulations was therefore itself an approximation.

In simulations of the PH-TH-kinase construct, the interface with the kinase domain remained intact, but the PH-TH module wobbled back and forth through a rotation of about 20° (Figure 3-supplement figure C). The averaged r.m.s. deviation between the crystal structure and structures from the end the trajectories, aligned on the kinase domain, was ~ 4 Å for the PH-TH module. The PH-TH module pivots about an apparent anchor point at the C terminus of helix α2, at the center of the autoinhibitory contact (Figure 3-supplement figure C).

The fluctuations in the PH-TH orientation generate transient structures in which there would be essentially no clashes between the SH3 domain and the PH-TH module in the merged structures. We initiated multiple MD trajectories from such a composite model (Figure 3A). The inhibitory interactions between each of the regulatory modules and the kinase domain were preserved, but various intra-domain rearrangements occurred during the course of these trajectories. The overall conformation of Btk at the end of the trajectories is very similar to the initial structure. The averaged r.m.s. deviations of the PH-TH module and the SH3 domain between the crystal structures and the end of trajectories, aligned on the Src-like module of the kinase, were ~4 Å and ~2.5 Å, respectively. The former is comparable to the value found in simulations of the PH-TH-kinase, but the latter is larger than in simulations of the Src-like module, probably because the SH3 domain reorients with respect to the kinase domain. The SH3 rotation results in a polar interface between SH3 and PH-TH, with good electrostatic complementarity, which preserves the interactions between the SH3 domain, the
linker and the N lobe of the kinase (Figure 4B). In these simulations, the r.m.s. deviations of
the SH2 domain and the kinase N lobe are ~2.6 Å and 2.0 Å, respectively, after 360 ns, similar
to those of the Src-like module simulations. We emphasize, however, that restrictions in the
time scale of the simulations allow only qualitative conclusions and do not give a high-
resolution picture of the intact molecule. Structures from several time points in the simulation of
the composite model are deposited on the eLife website (Supplementary File 1).

**Autoinhibition**

We probed the significance of the interactions seen in the crystal structures by studying
how various mutations affect the rate of autophosphorylation and enzymatic activity. We
avoided the heterogenous phosphorylation that accompanies expression in eukaryotic cells by
using bovine Btk expressed in bacteria, which gives a pure, unphosphorylated product with
good yield. Bovine Btk is 98.8% identical to human Btk in sequence, with only eight amino-
acid differences over the entire protein. The mass of the bacterially-expressed full-length Btk
(76379 Da), as determined by mass spectrometry, is consistent with the calculated molecular
weight (76381.2 Da).

Incubation with ATP-Mg\(^{2+}\) initiates autophosphorylation, leading in turn to increased
catalytic activity. We monitored activation in two ways. First, we monitored the
phosphorylation by Btk of a peptide substrate derived from PLC-\(\gamma\)2, using a continuous kinase-
coupled colorimetric assay (Figure 4A). Second, we followed accumulation of tyrosine-
phosphorylated Btk by immunoblotting with a non-specific, anti-phosphotyrosine antibody
(4G10, EMD Millipore) (Figure 4A). The results of the two assays are in good agreement.
The kinase domain of Btk has low catalytic activity and autophosphorylates very slowly, like the c-Abl kinase domain and unlike those of the Src family (Figure 4B). For example, there is no detectable change over 90 minutes in the level of phosphorylation of the Btk kinase domain at 4 μM concentration. Inclusion of the SH2 domain and the SH2-kinase linker (but not the SH3 domain or the PH-TH module) increases Btk autophosphorylation substantially (Figure 4B). We have not studied how the SH2 domain increases activity, but we note that in c-Abl, the SH2 domain docks onto the N lobe of the kinase domain and stabilizes the active conformation and that activation by the SH2 domain is also seen in Csk (Sondhi & Cole 1999) and c-Fes (Filippakopoulos et al. 2008; Nagar et al. 2006). The apparent linear array of domains detected by small-angle x-ray scattering from a partially phosphorylated form of full-length Btk might represent the activated rather than the inactive form, in a conformation similar to that of activated c-Abl (Márquez et al. 2003; Nagar et al. 2006).

The autoactivation rate of the complete Src-like module of Btk is lower than that of the SH2-kinase module (Figure 4B), as expected from the joint clamping effect of the SH3 and SH2 domains. Based on contacts seen in the crystal structure of the Src-like module of Btk, we introduced (separately) two mutations, Y223A and Y268A, into full-length Btk. These SH3-domain residues pack against Pro 385 in the SH2-kinase linker, and their phosphorylation (or mutation to alanine) would destabilize the autoinhibited conformation (Figure 1C). The autoactivation rates of both mutants are indeed about six-fold greater than that of the unmutated enzyme and comparable to that of the SH2-kinase module (the rate of activation is estimated from the slope of activity versus time, and is only a rough measure because of non-linearity; Figure 4C). An analogous interaction regulates c-Abl activation: phosphorylation of Tyr 89, in the SH3-kinase interface, is necessary for full activation (Brasher & Van Etten 2000).
Mutations expected to disrupt the PH-TH-kinase interface likewise increase the autoactivation rate of full-length Btk, consistent with our inferences from examining the interface between the PH-TH and the kinase N lobe (Figure 4D). Removal of the entire PH-TH module, while retaining the Src-like regulatory components, increases the rate of autoactivation about three-fold, as estimated from the slopes of the activity curves in Figure 4.

**Activation of Btk by membrane recruitment**

We studied the effect on Btk activity of PIP$_3$ in small unilamellar vesicles (SUVs). SUVs containing 75% DOPC, 20% DOPS and 5% PIP$_3$ produce a rapid and substantial activation of Btk (Figure 5A). At a Btk concentration of 2 µM, robust phosphorylation can be detected within the first five minutes of reaction in the presence of these vesicles. The extent of activation depends on the concentration of PIP$_3$ in the vesicles, presumably because increasing the PIP$_3$ fraction increases the surface density of Btk on the lipid vesicles and thereby increases the rate of trans-autophosphorylation (Figure 5A). An estimate of the surface density of PIP$_3$ on these vesicles suggests that the local concentration of Btk on the vesicle surface is in the millimolar range, i.e., 100 fold greater than in bulk solution (see materials and methods for detailed calculations).

We next studied the effect of inositol phosphates, the soluble headgroups of various phosphatidylinositol phosphate lipids, on Btk activity. The purpose of these studies was to see whether interactions with the soluble headgroups might activate Btk independent of membrane localization, as is the case with Akt (Calleja et al. 2009; Wu et al. 2010). The soluble headgroup of PIP$_3$ is IP$_4$, and so this was included in our study, as were several other inositol phosphates (Figure 5B). IP$_4$ itself has no effect on the rate of Btk activation in solution,
demonstrating that the role of PIP₃ is to recruit Btk to the membrane without necessarily triggering an allosteric effect.

**Activation of Btk by IP₆ in solution**

We tested the effect of 10 other inositol phosphates on the activity of Btk (Figure 5B). Quite unexpectedly, we discovered that inositol hexakisphosphate (IP₆) promotes Btk autophosphorylation strongly at 2 μM protein concentration, substantially enhancing Btk catalytic activity within the first 5 minutes after addition (Figure 5C). We also saw a milder activation with certain inositol pentakisphosphates (IP₅), but these required higher protein concentration and longer times for appreciable autophosphorylation than did IP₆ (Figure 5B).

The EC₅₀ value for IP₆ activation is around 20 μM (Figure 5D), in the range of the physiological concentration of IP₆ in cells, which is 10 μM to 80 μM (Shears 2001; Irvine & Schell 2001). In the presence of a saturating concentration of IP₆ (100 μM), full-length Btk is activated within the first 5 minutes of reaction time, in contrast to the 30 minutes and 120 minutes that are required for the activation of the SH2-kinase module and full-length Btk, respectively, in the absence of IP₆ or PIP₃-containing vesicles (Figure 5A and 5C). Once fully activated, the catalytic rates of various Btk constructs are the same with or without IP₆ addition, suggesting that the effect of IP₆ is to change the rate of autophosphorylation, and that it does not affect the catalytic machinery directly (Figure 5C).

By comparing constructs, we determined which domains in Btk are responsible for the IP₆ response. Activation by IP₆ required only the PH-TH module, and not the SH3 and SH2 domains (Figure 5-supplement figure A and B). A PH-TH-kinase construct that contained the PH-TH-SH3 linker and the SH2-kinase linker (with a total linker length of 45 residues) had the
same response to IP₆ as did full-length Btk (Figure 5-supplement figure B). PH-TH-kinase constructs with shorter linkers also responded to IP₆. The shortest linker necessary for activation contained only 13 residues from the SH2-kinase linker; we used this construct to determine the crystal structure described earlier (Figure 5-supplement figure C). The catalytic activity of the fully activated PH-TH-kinase construct was about three-fold lower than that of full-length Btk. This lower activity can be explained by the absence of the SH2 domain.

IP₆ has no effect on activation of the isolated kinase domain of Btk, suggesting that it is not a direct allosteric activator of the enzyme. The protein kinase CK2 is activated by IP₆, which binds to the kinase domain and dislodges an inhibitory interaction between CK2 and the protein Nopp140 (Lee et al. 2013). It seems unlikely that IP₆ works by displacing an inhibitory interaction between the kinase and the PH-TH module, because the Src-like module (lacking the PH-TH module) activates only slightly faster than full-length Btk in the absence of IP₆.

Nevertheless, in order to test this possibility, we introduced two mutations (Y133E and R134E) at the PH-TH-kinase interface that are expected to disrupt the inhibitory docking of the PH-TH module on the kinase domain. Full length Btk bearing these two mutations retains activation by IP₆, showing that IP₆ does not act by dislodging the interactions seen in the crystal structure (Figure 5-supplement figure D).

The canonical lipid-binding pocket of the PH-TH module is not critical for IP₆-mediated Btk activation

The structure of the PH-TH-kinase construct does not provide direct clues about how IP₆ might activate Btk. The IP₄-binding site on the PH-TH module (the “canonical binding site”)
is located ~15 Å from the kinase-interacting surface of the PH-TH module (Figure 2A). The structure of the PH-TH-kinase construct, in which no inositol phosphates are bound, resembles closely that of the liganded PH domain (Baraldi et al. 1999) in the vicinity of the PH-kinase interface, consistent with the failure of IP₄ to activate Btk.

We used isothermal titration calorimetry (ITC) to study the interaction between IP₆ and various constructs of Btk (Figure 6A). The ITC measurements show that among the four domains in Btk, only the PH-TH module interacts with IP₆. At 150 mM NaCl concentration, the titration curve for IP₆ binding to the PH-TH module can be fit by a one-site model for binding, with a dissociation constant, Kᵋ, of 250 nM (Table 3). We introduced two mutations, R28C and N24E, at the canonical binding site in the PH-TH module. Arg 28 and Asn 24 have been shown previously to be critical for IP₄ binding (Hyvönen & Saraste 1997; Baraldi et al. 1999). IP₄ binds to the PH-TH module with a Kᵋ value of 30 nM, and the R28C/N24E mutation in the PH-TH module reduces IP₄ binding to an undetectable level (Table 3). The ITC signal of IP₆ binding to the R28C/N24E mutant is reduced greatly but not completely abolished, and the residual signal corresponds to a dissociation constant between 5 μM and 10 μM.

Although the residual binding of IP₆ to the IP₄-binding deficient mutant is relatively weak, the observed affinity is consistent with the EC₅₀ value of 20 μM for IP₆ activation, as determined in our activation assays (Figure 5C). We observed no significant activation when the IP₆ concentration was lower than 5 μM, a level still high enough to saturate the canonical binding site. We also found that the R28C/N24E mutant that is deficient in IP₄ binding remains highly responsive to IP₆ stimulation, with only a slight reduction in the activation kinetics as compared to the wild-type protein (Figure 6B). These observations imply that the canonical
binding site of the PH-TH module may not play a major role in the IP$_6$-mediated activation of
Btk.

Structure of the IP$_6$-bound PH-TH module

We determined a crystal structure of the Btk PH-TH module bound to IP$_6$ at 2.3 Å
(Figure 7). There are four PH-TH modules in the asymmetric unit of the crystal, which form two
dimers, similar to those seen in other structures of the Btk PH-TH module (Figure 7D)
(Hyvönen & Saraste 1997; Baraldi et al. 1999). The only structural changes in the IP$_6$-bound
PH-TH module, relative to the apo- or IP$_4$-bound PH-TH module, are in the loop regions
(Figure 7-supplement figure B).

The four PH-TH modules in the asymmetric unit have six bound IP$_6$ molecules, each
with clearly resolved electron density. Four of the IP$_6$ molecules are in the canonical binding
sites on each module. At those sites, residues Asn 24, Tyr 39, Arg 28 and Lys 12 coordinate
phosphate groups on the 2, 3 and 4 positions of the myo-inositol ring; these are also the key
residues involved in IP$_4$ binding (Baraldi et al. 1999) (Figure 7B). The loop between strands β1
and β2, which is important for IP$_4$ binding, is largely disordered in our structure (Figure 7A).
There are fewer interactions at the canonical site between the PH-TH module and IP$_6$ than
seen for IP$_4$, consistent with the relative affinities of the two ligands (Figure 6A).

The two other IP$_6$ molecules in the crystallographic asymmetric unit are bound at a
peripheral site. This site is on the surface formed by an elongated β hairpin involving strands
β3 and β4 of each PH-TH module. This site has not been seen in any other PH domain
(Lemmon 2008); it differs from the two well-characterized lipid-binding sites in the PH domains
of Pleckstrin (Ferguson et al. 1995) and β-spectrin (Hyvönen et al. 1995). As in the canonical
binding site, interactions between IP$_6$ and the peripheral site are predominantly electrostatic (Figure 7B). Unlike the canonical site, which has a well-defined groove within which the inositol phosphates are bound, the peripheral site is relatively flat (Figure 7B).

A feature of the crystallographic lattice is that IP$_6$ bound at the peripheral site is sandwiched between two PH-TH modules, interacting with the peripheral sites on both (Figure 7-supplement figure A and C). The majority of interactions are with one subunit. Residues Arg 49, Lys 52, Lys 36, and Tyr 40 at the peripheral binding site of one subunit (Figure 7-supplement figure C) coordinate phosphate groups at the 1, 2, 3 and 4 positions of the myo-inositol ring. A subset of these residues, Lys 52 and Lys 36, on the other PH-TH module interact from the opposite side of the myo-inositol ring with the phosphate group at position 3 (Figure 7-supplement figure C). The bridging interactions provided by IP$_6$ generate an open-ended chain of PH-TH dimers in the crystal lattice (Figure 7-supplement figure A). Based on the limited interactions across this crystallographic interface, we do not believe that this open-ended chain is relevant to the function of Btk.

The peripheral binding site is critical for IP$_6$-mediated Btk activation

We mutated Arg 49 and Lys 52, in the peripheral binding site of the PH-TH module, to serine in full-length Btk. This Btk variant autophosphorylates slowly, as does wild-type Btk, and it has very little response to IP$_6$ (Figure 7C). In ITC experiments, we saw no detectable binding when we added IP$_6$ to a PH-TH module with mutations at both the canonical and the peripheral sites, further confirming that the residual binding of IP$_6$ to the IP$_4$-binding deficient mutant is at the peripheral binding site (Figure 7-supplement figure E).
The disposition of the IP₆ molecule can be determined unambiguously from the electron density because the phosphate group at the 2 position juts out from the plane of the myo-inositol ring (Figure 7E). IP₆ at the peripheral binding site interacts with the primary PH-TH module to which it is bound through the phosphates at positions 1, 2, 3, and 4. None of the inositol phosphates used in our experiments, other than IP₆, can fully satisfy this criterion, which may underlie the specificity of the PH-TH module for IP₆. Two isomers of IP₅, I(2,3,4,5,6)P₅ and I(1,2,3,5,6)P₅, have a mildly activating effect on Btk; they are more effective than I(1,3,4,5,6)P₅. These comparisons indicate an important role in Btk activation for the phosphate at position 2 (Figure 5B).

In the cell, the only known enzyme that can add a phosphate to the hydroxyl group at position 2 of a myo-inositol ring is inositol 1,3,4,5,6-pentakisphosphate 2-kinase (York et al. 1999), which produces IP₆. As a consequence, IP₆, and inositol pyrophosphates, such as IP₇ and IP₈, may be the only cellular phosphates that can interact with the peripheral site in the Btk PH-TH module.

A dimer of the PH-TH module is critical for IP₆-mediated Btk activation

Strong activation of Btk by concentration on PIP₃-containing membranes suggests that IP₆ might promote dimerization or oligomerization of Btk in solution. Neither multi-angle light scattering nor size exclusion chromatography gave convincing evidence for IP₆-induced association, at Btk concentrations up to 30 μM. Nevertheless, we found it striking that in every crystal structure of the Btk PH-TH module, as well as in our structure of the PH-TH-kinase construct, the module forms very similar crystallographic dimers (Figure 7D) (PDB entry: 1BTK (Hyvönen & Saraste 1997); PDB entry:1B55 and 1BWN (Baraldi et al. 1999); see also PDB
entry: 2Z0P). This dimer was first identified and discussed by Marko Hyvönen and the late Matti Saraste; we refer to it as the “Saraste dimer”.

The Saraste dimer is formed by symmetric interactions between helix α1 and strands β3 and β4 of the PH-TH module (Figure 7A). The largely hydrophobic dimer interface has a buried surface area of ~1000 Å² on each molecule. Phe 44 and Tyr 42 on strand β3, Ile 9 on strand β1 and Ile 95 on helix α1 create a hydrophobic patch in one subunit of the dimer that packs tightly against the corresponding patch on the other subunit (Figure 8A).

To test whether the Saraste dimer interface is important for IP₆-mediated Btk activation, we made three sets of mutations in which we replaced hydrophobic residues at the Saraste dimer interface with arginines and compared their autophosphorylation rates and their response to IP₆ with those of wild-type Btk. All three sets of mutant proteins showed little or no response to IP₆ (Figure 8B). These mutations do not affect the autophosphorylation rate of Btk in the absence of IP₆. We conclude that the Saraste dimer interface promotes Btk association in the presence of IP₆.

The activation of Class I Histone Deacetylases (HDACs) requires the formation of a multi-subunit co-repressor complex, in which Ins(1,4,5,6)P₄ serves as an intermolecular glue that cements the complex together (Millard et al. 2013; Watson et al. 2012). The mechanism of IP₆-induced transient Btk dimerization is fundamentally different, in that the peripheral IP₆ binding site is just outside the Saraste dimer interface, and IP₆ does not participate in any interactions between the two Btk subunits in that dimer.

**Fusing the Btk PH-TH module to the catalytic module of c-Abl promotes autophosphorylation of c-Abl in the presence of IP₆**
If the effect of IP$_6$ arises from its ability to promote dimerization of the PH-TH module, fusion of a Btk PH-TH module with the catalytic domain of another kinase that autophosphorylates slowly should impart activation by IP$_6$. A good example of such a kinase is c-Abl. Indeed, in chronic myelogenous leukemia, a chromosomal translocation results in fusion of a coiled-coil oligomerization domain with c-Abl, leading to hyper-activation of the BCR-Abl fusion protein by autophosphorylation. Several other fusion variants involving c-Abl, in which different oligomerization domains are present, are associated with hematological malignancies (De Braekeleer et al. 2011).

We generated a PH-TH-Abl fusion construct in which the PH-TH module of Btk is fused to the SH2-kinase module of c-Abl. We compared autophosphorylation rates of the PH-TH-Abl fusion protein with and without IP$_6$ at a saturating concentration (100 μM). The fusion protein at a concentration of 1 μM autophosphorylated slowly in the absence of IP$_6$, while the IP$_6$-bound protein became heavily phosphorylated within the first 30 s, the dead-time of our assay (Figure 8C). We replaced Phe 44 and Tyr 42, at the Saraste dimer interface, with arginine in the fusion protein. The autophosphorylation rate of the PH-TH-Abl mutant in the presence of IP$_6$ was much lower than that of the IP$_6$-bound wild-type protein, further evidence for the importance of the Saraste dimer interface in IP$_6$-mediated activation (Figure 8C).

The Btk and c-Abl kinase domains are 46% identical in sequence, but most of the surface residues are different. We infer that, except for its enzymatic activity, the interaction properties of the Btk kinase domain are not important for the IP$_6$ response. Because IP$_6$ activates both kinases when they are linked to the Btk PH-TH module, we propose that IP$_6$ promotes transphosphorylation of the attached kinase by inducing transient dimerization.
Our preliminary studies indicate that the IP₆-induced dimerization of the Btk PH-TH module is very weak, with a dissociation constant that is likely to be greater than 100 μM. It might appear counter-intuitive that IP₆ can markedly accelerate activation at 2 μM Btk concentration, well below the dissociation constant. A simple calculation shows that even a modest increase in a small dimer population can lead to a sharp change in the rate of activation because the reaction is autocatalytic. The initial production of phosphorylated Btk drives the further production of this species at an accelerating rate (Figure 9 – supplement figure B).

Speculations about the mechanism of IP₆ activation

Our failure to detect stable dimers of the Btk PH-TH module or of full-length Btk in the presence of IP₆ suggests that the monomeric PH-TH module has a conformation different from the one seen in crystal structures, which all show the same dimer contact. The high protein concentration in the crystal lattice (~30 mM) presumably drives the formation of the Saraste dimer, which is otherwise not favored.

Molecular dynamics simulations suggest that the conformation of the PH-TH module seen in the Saraste dimer is unstable when the protein is monomeric. We initiated molecular dynamics trajectories using a monomeric PH-TH module from a crystal structure which contained neither IP₆ nor IP₄. Within the first 30 ns of simulation, the β3-β4 hairpin bent towards the core of the PH-TH module (Figures 9A and supplement figure A), and Phe 44, at the heart of the Saraste dimer interface, changed its conformation and packed against the
other three interfacial residues, Tyr 42, Ile 9 and Ile 95 (Figure 9A). This conformational change closed the dimer interface, burying the interfacial residues.

The hydrophobicity of the dimer interface drove the observed conformational change. Molecular dynamics simulations initiated from crystal structures but with the four interfacial residues substituted by alanine reduced the extent of conformational change in this area (Figure 9A). The peripheral binding site for IP6 is adjacent to the β3-β4 hairpin that folds in and closes the dimer interface in the simulation. Thus, one effect of IP6 binding may be to stabilize the open form of the hairpin.

It also appears that IP6 might act by altering the electrostatic potential of the PH-TH module. We examined the surface electrostatic potential of the Btk PH-TH module by viewing graphical displays of solutions to the Poisson-Boltzmann equation (Dolinsky et al. 2004; Dolinsky et al. 2007). The PH-TH domain carries 7 net positive charges; it is also electrostatically polarized, with the positively charged residues concentrated on the broad, contiguous surface composed of the β3-β4 hairpin, the IP4 binding loop and the N-terminal region of the loop connecting strand β5 and helix α2. In the Saraste dimer, positive electrostatic field lines calculated based on the monomer run into each other at the dimer interface, suggesting that electrostatic forces oppose dimer formation.

We introduced two charge-reversal mutations at the peripheral IP6-binding site (K49E and R52E), to reduce the positive charge on the protein by four units. The autophosphorylation rate of full-length Btk with the K49E/R52E mutations was substantially higher than that of the wild-type protein, with pronounced phosphorylation observed in the first 2 minutes of the reaction (Figure 9B).
We suggest that there are two barriers to the dimerization of the PH-TH module of Btk. First, the dimer interface of the PH domain is probably closed in the monomeric protein. Second, an electrostatic repulsion between PH domains further impedes dimerization. IP\textsubscript{6} binding might stabilize the open form of the PH-TH module allosterically, while also alleviating the electrostatic repulsion.

Conclusions

Our studies on Btk have revealed the structural basis for its autoinhibition. One key finding is that the Src-like module of Btk adopts a compact and autoinhibited form that resembles the autoinhibited structures of the Src kinases and c-Abl, in which the SH3 domain binds to the SH2-kinase linker and holds the kinase domain in an inactive conformation. Our structure of the PH-TH-kinase construct of Btk shows that the PH-TH module stabilizes the inactive conformation of the kinase domain, as well as the assembled conformation of the Src-like module of Btk. A second, and quite unexpected, finding is that IP\textsubscript{6} can activate Btk strongly by binding to a newly identified site on the PH domain, probably by promoting transient dimerization of the PH-TH module. Because IP\textsubscript{6} synthesis requires the precursor molecule IP\textsubscript{3}, which is itself produced as a result of Btk activation, a transient elevation of IP\textsubscript{6} may create a positive feedback loop for Btk activation, either on the membrane or in the cytoplasm. Formation of the Saraste dimer interface is compatible with the simultaneous engagement of PIP\textsubscript{3} by both PH domains in the dimer, raising the possibility that IP\textsubscript{6} can stimulate dimerization of membrane-tethered Btk (Figure 10). IP\textsubscript{6} binding to the peripheral binding site may also enhance PIP\textsubscript{3}
affinity in the canonical binding site and sensitize Btk to PIP₃ levels. The linker connecting the PH-TH unit to the SH3 domain promotes dimerization of Btk by interacting with the SH3 domain of a second molecule, providing an additional mechanism for Btk dimerization at high local concentration (Laederach et al. 2002). High levels of IP₆ may down-regulate the amount of Btk on membranes by competing for PIP₃ binding in the canonical binding site, as proposed for Akt1 regulation by IP₇ (Chakraborty et al. 2010).

The Src-like modules of Btk and Itk have 55% sequence identity, with all the residues critical for maintaining the assembled and autoinhibited Src-like conformation of Btk conserved in Itk (Figure 10-supplement figure). The Src-like module of Itk is therefore likely also to adopt the assembled Src-like conformation when inactive.

The roles of the PH-TH module in regulating Btk activity are not likely to be conserved in Itk. The four residues at the peripheral binding site that coordinate IP₆ in Btk are not conserved in Itk, unlike those that coordinate PIP₃/IP₄ at the canonical lipid-binding site. This lack of conservation also extends to the key residues at the inhibitory interface between the PH-TH module and the kinase domain in Btk and to residues at the Saraste dimer interface (Figure 10-supplement figure). A different soluble inositol phosphate, IP₄, promotes PIP₃ binding by the Itk PH-TH module; this activity is important for Itk regulation in T-cells (Huang et al. 2007). Thus, regulation by soluble inositol phosphates may be a general feature of Tec family kinases, but the specific effector and the molecular mechanism of the kinase response may be different in each case. The regulation of tyrosine kinases by soluble second messengers has not received much attention in the past. Our discovery that IP₆ activates Btk, along with the known effect of IP₄ on Itk, points to the importance of further studies on these and soluble effectors.
Figure 1: Crystal structure of the Src-like module of Btk.

A. Domain architectures of Btk, c-Abl and c-Src.

B. Model for the Src-like module of Btk, based on the crystal structure of the domain-swapped dimer.

C. Comparison of the Src-like modules of Btk and c-Abl (PDB: 1OPK) (Nagar et al. 2003), superimposed on the C lobes of the kinase domains. There is a relative rotation of about 15° for the SH2 domains and 20° for the SH3 domains for the two structures.

D. Details of the SH3/SH2-kinase linker interface and the kinase catalytic cleft in Btk. Left panel: A cluster of hydrophobic residues on the SH3 domain packs against the polyproline type-II helix formed by the SH2-kinase linker. This type of interaction is seen in most SH3-peptide ligand complexes. Right panel: The activation loop of Btk folds into the mouth of the catalytic cleft, blocking part of the ATP-binding cleft. Glu 445, in helix αC, forms an ion pair with Lys 430 in the active conformation but is prevented from doing so in this inactive conformation by the activation loop and helix αC.
**Figure 1- supplement figure**

A. The Src-like module of Btk forms a domain-swapped dimer in the crystal lattice, with one molecule per asymmetric unit. The SH3 domain and the kinase domain are intact, but the SH2 domain forms a domain-swapped dimer with another molecule.

B. Molecular details of the domain-swapped SH2 dimer in Btk and Grb2 (Schiering et al. 2000). The SH2 domain opens up at a position within the β-sheet, so that helix αB of one polypeptide chain packs against the β-sheet of the other.

C. Interactions between the SH2-kinase linker and the kinase domain. Leu 390 of the linker stabilizes the inactive conformation of the kinase domain by being sandwiched between residues Trp 421 and Tyr 461 in the kinase domain.

D. The “electrostatic switch” in the Btk kinase domain. The active conformation of the Btk kinase domain is modeled from the crystal structure of Lck (PDB: 3LCK) (Yamaguchi & Hendrickson 1996) using Modeller (Eswar et al. 2006). Activation entails a shuffling of salt-bridge and polar hydrogen-bond partners.
**Figure 2: Crystal structure of the PH-TH-kinase construct of Btk**

A. Crystal structure of the PH-TH-kinase construct. The PH-TH module is connected to the kinase domain via a 13-residue linker from the SH2-kinase linker of Btk. The kinase domain is bound to the inhibitor CGI1746 (Di Paolo et al. 2011). The canonical PIP₃/IP₄ binding pocket of the PH domain is located distal to the interface with the kinase, and the peripheral binding site for IP₆ is indicated.

B. The interface between the PH-TH module and the kinase domain of Btk.
Figure 2 - supplement figure

A. Crystal structure of the isolated Btk kinase domain with mutations in the activation loop. These mutations (L542M, S543T, V555T, R562K, S564A, and P565S) are based on a previous study (Joseph et al. 2013), and they improved the quality of the crystals of the PH-TH-kinase construct. The structure of the mutant kinase is very similar to that of the wild-type Btk kinase domain (PDB: 3OCS).

B. Comparison of the Btk PH-TH-kinase structure (left) and the Akt1 PH-kinase structure (right). In Akt1, The PH domain is docked between the N lobe and the C lobe of the kinase domain, and it blocks substrate access to the catalytic cleft. The lipid-binding site on the PH domain is buried at the PH-kinase interface, and binding of PIP$_3$/IP$_4$ breaks the PH-kinase interface and activates Akt1 (Wu et al. 2010; Calleja et al. 2009).
Figure 3: A structural model for full-length Btk

A. A composite model for full-length Btk. The PH-TH domain sits on top of the kinase domain and the SH3 domain, serving as a “latch” that presumably stabilizes the Src-like module in the autoinhibited conformation. The 45-residue linker between the PH-TH module and the SH3 domain is represented by the dotted brown line. Two orthogonal views of an instantaneous structure at 360 ns from a molecular dynamics trajectory for the composite model are shown.

B. Inter-domain interactions in an instantaneous structure at 360 ns from a simulation of the composite model. The SH2/kinase interface and the PH-TH /SH3 interface are formed and remain stable during the molecular dynamics simulation.
Figure 3-supplement figure

A. Overlay of the crystal structures of the Btk Src-like module and the PH-TH-kinase construct, using the kinase domain C lobe as the reference. Sidechain clashes (circled) are observed between helix α2 in the PH-TH module and the β3/β4 loop segment in the SH3 domain.

B. Fluctuations in the Btk Src-like module occurred during a 100 ns molecular dynamics simulation. An instantaneous structure (t=89 ns) from the simulation is overlaid on the crystal structure of the Src-like module.

C. Fluctuations in the Btk PH-TH-kinase construct during a 100 ns molecular dynamics simulation. An instantaneous structure (t=12 ns) from the simulation is overlaid on the crystal structure of the PH-TH-kinase construct using the kinase domain N lobe as the reference. The PH-TH module pivots (black arrow) about an apparent anchor point at the C terminus of helix α2.

D. Steps used to generate a composite model for full-length Btk. An instantaneous structure (t=89 ns) of the Src-like module and an instantaneous structure (t=12 ns) of the PH-TH-kinase construct from the molecular dynamics trajectory are overlaid, using the kinase domain C-lobe as the reference. In contrast to a corresponding overlay of the crystal structures, there are no sidechain clashes between helix α2 in the PH-TH module and the loop in the SH3 domain. A model for full-length Btk is obtained by combining the PH-TH module with the Src-like module in the two instantaneous structures, and is subject to further molecular dynamics simulation.
Figure 4: Autoinhibition of Btk

A. Activation of full-length bovine Btk (residues 1 to 659, 2 µM). Reactions are carried out in the presence of 10 mM Mg$^{2+}$, 150 mM NaCl, 1 mM ATP, 25 mM Tris-HCl pH 8.0. The level of autophosphorylation is assayed by immunoblotting an SDS-PAGE gel with a non-specific, anti-phosphotyrosine antibody (4G10, EMD Millipore) (upper panel). The amount of total protein loaded on the gel is measured by coomassie-blue staining. The kinase activity of Btk is assayed by a continuous kinase-coupled colorimetric assay, in the presence of 1 mM PLC-γ2 peptide substrate. See methods for detailed experimental procedures.

B. Comparison of the activation of the Btk Src-like module (residues 217 to 659), SH2-kinase (residues 270 to 659) and the kinase domain (residues 394 to 659). The SH2-kinase construct activates substantially faster than full-length Btk and the Src-like module of Btk. Activated full-length Btk degrades to a small extent over time, which results in some lower molecule-weight bands being detected on the western blot.

C. Activation of full-length Btk with mutations Y223A and Y268A. Tyr 223 and Tyr 268 are on the SH3/SH2-linker interface, and the two mutants activate faster than wild-type Btk.

D. Activation of full-length Btk with a double mutation (R134E/Y133E). Arg 134 and Tyr 133 are located at the PH-TH/kinase interface.
**Figure 5: Activation of Btk**

A. Activation of full-length Btk on membranes. Left panel: Btk (1 µM) autophosphorylates rapidly on lipid vesicles (total lipid concentration, 1 mM) containing 75% DOPC, 20% DOPS and 5% PIP₃. Right panel: end-point autophosphorylation assay of Btk in the presence of lipid vesicles. The molar concentration of DOPC is kept at 75%, and the lipid fractions of PIP₃ is 0%, 2%, 5% and 10%, with the remaining lipid fraction filled with DOPS. The measurements are performed after two minutes incubation with lipid vesicles in the presence of ATP/Mg²⁺.

B. End-point autophosphorylation assays of Btk in solution in the presence of inositol phosphates. Upper panel: Btk (2 µM) autophosphorylates rapidly in the presence of 100 µM IP₆ in solution, with the results after the first five minutes of incubation shown here. Essentially no phosphorylation is detected in Btk samples incubated with IP₃, IP₄ or I(1,3,4,5,6)P₅, and very low phosphorylation is detected in Btk samples incubated with I(1,2,3,5,6)IP₅ and I(2,3,4,5,6)IP₅ in the same experiment. Lower panel: The assay is repeated with other inositol phosphates at a higher protein concentration (4 µM) and longer incubation time (10 minutes), and the gel was exposed for a longer time. There is a small amount of degraded Btk in the sample, as judged by the coomassie staining, leading to multiple bands in the western blot shown here and in Panel C.

C. Activation of full-length Btk (2 µM) in the presence of IP₆ (100 µM).

D. Activation of full-length Btk (2 µM) with increasing concentrations of IP₆
**Figure 5-supplement figure**

A. End-point autophosphorylation assay for full-length Btk, the Src-like module, the SH2-kinase construct and the kinase domain in the presence/absence of IP₆. The measurements are carried out after incubating proteins (2 µM) with and without IP₆ (100 µM) for two minutes in the presence of ATP/Mg²⁺.

B. End-point autophosphorylation assay for Btk PH-TH-kinase constructs of various linker lengths in the presence/absence of IP₆ (100 µM). The measurements are carried out after incubating proteins (2 µM) with and without IP₆ (100 µM) for two minutes at the presence of ATP/Mg²⁺.

C. Activation of the PH-TH-kinase construct (2 µM) with a 13-residue linker in the presence and absence of IP₆ (100 µM).

D. Activation of full-length Btk (2 µM) with mutations R133E/Y134E in the presence and absence of IP₆ (100 µM). Arg 133 and Tyr 134 are located at the PH-TH/kinase interface.
**Figure 6: Binding of IP$_4$ and IP$_6$ to the Btk PH-TH module.**

A. Representative isothermal titration calorimetry data for the Btk PH-TH module and its R28C/N24D variant binding to IP$_4$ or IP$_6$. The protein concentration is 20 μM and that of the inositol phosphates goes up to 300 μM. Experiments are performed at 20 °C. Titration curves are shown with the baseline-corrected raw data. The parameters from fitting a one-site binding model are listed in Table 3.

B. Activation of full-length Btk N24D/R28C mutant (2 μM) in the presence and absence of IP$_6$ (100 μM). Asn 24 and Arg 28 are critical residues for PIP$_3$/IP$_4$ binding in the canonical lipid-binding pocket of the PH-TH module.
Figure 7: Crystal structure of the PH-TH module bound to IP₆.

A. Structure of the PH-TH module bound to two IP₆ molecules. One IP₆ molecule is in the canonical lipid-binding site and the other IP₆ molecule is in the peripheral binding site formed by strands β3 and β4. The β1/β2 loop (dotted lines) is disordered in this structure.

B. IP₆ coordination in the canonical binding site and in the peripheral binding site.

C. Activation of the Btk R52S/K49S mutant (2 μM) in the presence and absence of IP₆ (100 μM). Arg 52 and Lys 49 are two residues interacting with IP₆ in the peripheral binding site.

D. A dimer assembly of the Btk PH-TH module in the crystal structure of the IP₆-bound PH-TH module. The canonical binding site and the peripheral binding site are shown in magenta circle and blue circle respectively. We refer to this dimer as the “Saraste dimer” (Hyvönen & Saraste 1997).

E. An electron density map, using σₐ-weighted 2m|Fo|-|Fc| coefficients (Read 1986) for IP₆ at the peripheral binding site, contoured at 1.5 σ.
**Figure 7-supplementary figure**

A. An open-ended chain of PH-TH dimers in the crystal lattice, mediated by IP₆ at the peripheral site. IP₆ is bound at the peripheral site in such a way that only one IP₆ molecule occupies the two sites at each interface.

B. Structure superposition of the IP₆-bound PH-TH module with apo- and the IP₄-bound PH-TH module. The β1/β2 loop, which coordinates IP₄ binding, adopts a different conformation in the apo-structure, and is partially disordered in the IP₆-bound structure.

C. IP₆ coordination in the peripheral binding site by a crystal symmetry mate of the PH-TH module in the crystal structure of IP₆-bound PH-TH module. IP₆ and the residues involved in coordination are shown in sticks. The hydrogen bonds are shown in dotted lines. This is a detailed view of the interfacial sites shown in the schematic diagram in Panel A.

D. Isothermal titration calorimetry data for Btk PH-TH mutant R52E/K49E/R28C/N24D titrated with IP₆. Arg 52 and Lys 49 are critical residues coordinating IP₆ binding in the peripheral site. Arg 28 and Asn 24 are critical residues coordinating IP₆ binding in the canonical site. Note that IP₆ does not bind to this mutant form of the PH-TH module.

E. Activation of full-length Btk, Btk R52S/K49S mutant, and Btk R28C/N24D mutant in the presence of IP₆ (100 µM).
Figure 8: The Saraste dimer of the PH-TH module is critical for Btk activation by IP$_6$.

A. Molecular details of the Saraste dimer interface in the crystal structure of the IP$_6$-bound PH-TH module.

B. End-point autophosphorylation assays for wild-type Btk, Btk L29R/I9R mutant, Btk I94R/I95R mutant, and Btk Y42R/F44R mutant in the presence and absence of IP$_6$ (100 µM). The measurements were made after incubating the proteins with and without IP$_6$ in the presence of ATP/Mg$^{2+}$ for five minutes. Residues Leu 29, Ile 9, Ile 94, Ile 95 Tyr 42 and Phe 44 are located on the Saraste dimer interface, as shown in Panel A. Activation of Btk Y42R/F44R mutant is shown in the lower panel.

C. Activation of the Btk-Abl fusion construct (1 µM) and that of a variant of this fusion protein in which the Btk PH-TH module is mutated (Y42R/F44R). Measurements were made with 1 µM protein in the presence and absence of IP$_6$ (100 µM).
Figure 9: Possible allosteric and electrostatic effects of binding IP$_6$

A. Fluctuations of the PH-TH module during a molecular dynamic simulation. An instantaneous structure (t=100 ns) from a 100-ns simulation of the PH-TH module shows a conformational change that closes the Saraste dimer interface (middle panel). Replacing these hydrophobic residues (Ile 9, Tyr 42, Phe 44 and Ile 95) with alanine prevents the dimer interface from closing in the simulations (lower panel). Although the β3/β4 loop becomes more dynamic, it remains open (see Figure 9-supplement figure). Binding of IP$_6$ may shift the equilibrium between open and closed conformations.

B. Activation of the Btk R52E/K49E mutant (2 μM). Arg 52 and Lys 49 are two residues in the peripheral binding site. Mutation of these two residues to glutamate reduces the local charge at the peripheral site by a net 4 units, and results in Btk activation.
Figure 9-supplement figure

A. Fluctuations in the β3/β4 loop of the wild-type PH-TH module and its I9A/Y42A/F44A/I95A variant. Each trajectory was sampled every 1 ns. The instantaneous structures are aligned using the crystal structure as the reference. The r.m.s. deviation was calculated for the backbone of residues 36 through 57 of the PH-TH domain.

B. Simulations of the kinetics of Btk autophosphorylation. We consider a very simple two-step Michaelis-Menten model for autophosphorylation, in which two Btk molecules first form an encounter complex, followed by phosphorylation of one of the two molecules in the complex. We assume that the rate of complex formation increases if IP$_6$ is bound, and that the catalytic rate constant for phosphorylation of the other increases if one of the two Btk molecules is phosphorylated. This model is described by the following reaction scheme:

\[
\begin{align*}
(1) & \quad Btk + Btk \xrightarrow{k_1} Btk\cdot Btk \\
(2) & \quad Btk\cdot Btk \xrightarrow{k_{-1}} pBtk + Btk \\
(3) & \quad pBtk + Btk \xrightarrow{k_2} pBtk \cdot pBtk \\
(4) & \quad pBtk\cdot Btk \xrightarrow{k_3} pBtk + pBtk
\end{align*}
\]

The on-rate for a diffusion-limited encounter between two proteins is in the range of $10^7$ to $10^8$ M$^{-1}$sec$^{-1}$. We assume that in the absence of IP$_6$ the on-rate for Btk dimer formation is much slower than this ($k_1 = k_3 = 10^4$ M$^{-1}$sec$^{-1}$) and that IP$_6$ binding increases the on-rate by 10-fold ($k_1 = k_3 = 10^5$ M$^{-1}$sec$^{-1}$). The dissociation rate constant is chosen so that the dissociation constant is 2 mM and 200 μM in the absence and presence of IP$_6$, respective ($k_{-1} = k_{-3} = 20$ sec$^{-1}$). The catalytic constant ($k_4$) for activated Btk is set to 1.0 sec$^{-1}$, comparable to the values
determined for other tyrosine kinases, such as Src and ZAP70 (Mukherjee et al. 2013). We assume that the catalytic constant is 10-fold smaller for unphosphorylated Btk \( k_2 = 0.1 \text{ sec}^{-1} \).

To compare with our experimental data, the initial concentration of Btk was set at 1 \( \mu \text{M} \). The integration of the differential kinetic equations was done using BerkeleyMadonna (http://www.berkeleymadonna.com/). As shown in the diagram, the autophosphorylation kinetics generated by this simple reaction scheme is consistent with the experimentally observed kinetics.
Figure 10: Models of Btk autoinhibition and activation.

A. The PH-TH module stabilizes the assembled conformation of the Src-like module of Btk in its inactive conformation. The activation of Btk involves formation of a transient dimer through the PH-TH module, which promotes trans-autophosphorylation and activates Btk. The stimulation of dimerization could occur at the membrane or in solution, as in our experiments.

B. IP₆ may stimulate membrane-tethered Btk. Formation of the Saraste dimer is compatible with the simultaneous engagement of PIP₃ by both PH domains in the dimer, and is compatible with the binding of IP₆ at the peripheral binding site. This raises the possibility that IP₆ can stimulate dimerization of the membrane-tethered Btk.
Figure 10-supplement figure

Sequence alignment of Btk and Itk. The alignment was performed using ClustalW2 (Larkin et al. 2007) and visualized by Espript3 (Robert & Gouet 2014). The alignment shows that the adoption of the compact autoinhibited conformation by the Src-like module, seen in our crystal structures, is likely to be conserved in Itk. In contrast, residues at the Saraste dimer interface and at the peripheral IP₆ binding site in Btk are not conserved in Itk.
Table 1. Data collection and refinement statistics

The CC$_{1/2}$ values for the PH-TH-kinase dataset, IP$_6$-bound PH-TH dataset and the kinase domain with mutations in the activation loop dataset are 99.9 (86.5), 99.9 (55.4) and 99.9 (90.7) respectively.
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<th>Src-like module of mouse Btk (217-659)</th>
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**Data collection**

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**Refinement**

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**No. Atoms**

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Table 2. Data statistics for the Src-like module of Btk

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Table 3. Thermodynamic parameters for Btk PH-TH module binding to IP$_6$ and IP$_4$

The integrated heat from representative isothermal titration calorimetry experiments was fit with a one-binding site model. See methods for the details of data analysis.

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<th>$K_d$ (nM)</th>
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Table 4: Structures of fragments of Tec family kinases in the Protein Data Bank

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Supplementary File 1: Structures from several time points in the simulation of the composite model for full-length Btk.

The structures are taken every 20 ns from a 360-ns molecule dynamic simulation of the composite model. See methods for the details of simulations.
Materials and methods

Protein expression and purification

*Insect cell expression, purification, and characterization of mouse Btk*

DNA for mouse Btk (residues 214 to 659; the Src-like module) was used to prepare baculovirus. Infected Sf21 cells were harvested by centrifugation 72 hours post-infection, then frozen in liquid nitrogen for storage. Cells were later thawed, resuspended in 25 mM Tris·Cl, pH 8.0, 50 mM NaCl, 5 mM DTT, 5 mM benzamidine, and complete protease inhibitor tablets (Roche Diagnostics), and lysed mechanically by Dounce homogenization followed by brief sonication. All remaining steps were carried out at 4°C. Cellular debris was cleared by centrifugation at 96,000g for one hour, after which the supernatant was passed through a 0.45 μm filter. Cleared supernatant was loaded on to DEAE sepharose pre-equilibrated with 25 mM Tris·Cl, pH 8.0, 25 mM NaCl and 5 mM DTT. Protein was eluted by a linear gradient to 25 mM Tris·Cl, pH 8.0, 0.35 M NaCl, 5 mM DTT over 3.7 column volumes, with the Src-like module of Btk eluting at approximately 2.6 column volumes.

Fractions containing the Src-like module of Btk were identified by Western blot, pooled, brought to 0.7 M ammonium sulfate, then loaded on to a phenyl sepharose column pre-equilibrated with 50 mM Tris·Cl, pH 8.0, 0.7 M ammonium sulfate and 5 mM DTT. Contaminating protein was removed by an initial step to 0.2 M ammonium sulfate and the column eluted with a linear gradient to 50 mM Tris·Cl, pH 8.0 and 5 mM DTT over four column volumes. The Src-like module of Btk eluted as a broad peak starting at approximately 0.18 M ammonium sulfate.

Fractions containing significant amounts of the Src-like module of Btk, identified by SDS-PAGE, were pooled and concentrated using Centricon Plus-80 concentrators (Millipore).
Concentrated protein was dialyzed against 4 x 1 liter changes of 25 mM Tris·Cl, pH 8.5, 25 mM NaCl, 5 mM DTT. That buffer, excepting changes to the NaCl concentration, was maintained through the rest of the purification. After dialysis, protein was loaded on to a mono-Q HR 10/10 column, and eluted with a linear gradient from 50 mM NaCl to 250 mM NaCl over twenty column volumes, with the protein eluting as two peaks at approximately 100 mM NaCl (peak A) and 150 mM NaCl (peak B), respectively. Fractions from the two peaks were pooled separately. Fractions from peak B were dialyzed against buffer containing 20 mM NaCl. The two pools could then be concentrated using Centricon-10 concentrators (Millipore). They were loaded separately onto a HiPrep S-200 16/60 size exclusion column pre-equilibrated in buffer containing 25 mM NaCl. Peak A eluted at the position corresponding to a globular protein of molecular mass 55 kDa, the value expected for a monomer of the Src-like module. Peak B eluted at a position corresponding of 140 kDa, compatible with an extended dimer. The protein in peak A could be concentrated using Centricon concentrators to approximately 20 mg/ml, while peak B required dialysis against a buffer containing no more than 10 mM NaCl, followed by concentration in dialysis tubing laid on PEG 20,000 at 4°C. In this way, the dimeric protein could be concentrated to approximately 10 mg/ml.

Peaks A and B were definitively identified as the Src-like module of Btk monomer and dimer, respectively, by chemical cross-linking using disuccinimidyl glutarate (Pierce Biotechnology) and PAGE analysis (results not shown). All column media and pre-packed columns were purchased from Amersham Biosciences.

*Bacterial expression and purification of full-length bovine Btk and the PH-TH-kinase construct*
DNA encoding bovine Btk (residues 1 to 659) was cloned into pET-28 expression plasmid yielding N-terminally 6-histidine-tagged SUMO-fusion protein. The hexahistidine-SUMO tag was cleavable by Ulp-1 SUMO specific protease. For protein expression, the plasmid was transformed into BL21 DE3* (Novagen) cells that also contain the expression plasmids for GroES/GroEL and phosphatase YopH (Seeliger et al. 2005), and then grown in Terrific Broth (TB) media supplemented with 50 mg/l kanamycin, 50mg/l streptomycin and 50mg/l chloramphenicol at 37°C. At a cell density corresponding to an absorbance of 1.5 at 600 nm, the temperature was reduced to 18°C, and protein production was induced by mixing the bacterial culture with equal volume of 4°C TB media supplemented with 1 mM IPTG, 100 μM ZnCl₂, 50 mg/L Kanamycin, 50mg/L streptomycin and 50mg/L chloramphenicol. After 16 hours, cells were harvested by centrifugation, resuspended in Ni-NTA buffer A (500 mM NaCl, 20 mM imidazole, 25 mM Tris pH 8.5, 5% glycerol and 1mM DTT), and flash-frozen in liquid nitrogen. Cells were then thawed on ice and supplemented with a mixture of protease inhibitors (aprotinin 2 μg/mL, benzamidine 15 μg/mL, leupeptin 2 μg/mL, PMSF 1 mM, pepstatin A, 1μg/mL).

After cell lysis by French press and removal of cell debris by centrifugation, clear lysates were loaded onto a HiTrap Ni-NTA column (GE Healthcare), washed with 100 mL Ni-NTA buffer A. Proteins were eluted with a single step of Ni-NTA buffer A supplemented with 250 mM imidazole. Proteins were buffer exchanged into storage buffer (25 mM Tris-HCl, pH 8.0, 400 mM NaCl) by FastFlow desalting column (GE Healthcare), and affinity tags were removed by incubation with Ulp-1 protease at 4°C overnight. Cleaved proteins were collected in the flow-through during Ni-NTA affinity chromatography, and were subjected to size-exclusion chromatography on a Superdex 200 column (GE Healthcare) equilibrated in storage buffer.
Proteins were concentrated on an AmiconUltra centrifugal filter device (50 kDa cutoff; Millipore) to a final concentration of ∼400 μM. Protein aliquots were frozen in liquid nitrogen and stored at -80°C.

The Src-like module (residues 217 to 659), SH2-kinase module (residues 270 to 659), and the kinase domain (residues 394 to 659) of bovine Btk, various bovine Btk PH-TH-kinase constructs and the PH-TH-Abl fusion construct (BTK residues 1 to 216 fused to human c-Abl residues 140 to 518) were all expressed and purified using the same protocol. The PH-TH module of bovine Btk (residues 1 to 172) was prepared in the same way as described above expect that the co-expression of GroEL/GroES and YopH is not necessary.

**Crystallization and structure determination**

*The Src-like module of mouse Btk*

The crystals of the dimeric protein (peak B) were grown using hanging-drop vapor diffusion at 4°C by mixing purified protein at 5 mg/ml in 25 mM Tris·Cl, pH 8.5, 10 mM NaCl, 5 mM DTT with an equal volume of the precipitant solution, 100 mM Tris·Cl, pH 8.5, 200 mM Na·acetate, 7.5 % PEG 4,000. Wedge shaped crystals grew to approximately 0.1 mm on a side after a period of two weeks. These crystals could be frozen by first transferring them, in four sequential steps, to a solution of 100 mM Tris·Cl, pH 8.5, 200 mM Na·acetate, 15 % PEG 4,000, followed by transfer in six sequential steps to a solution containing 100 mM Tris·Cl, pH 8.5, 200 mM Na·acetate, 15 % PEG 4,000, 30% glucose. The monomeric protein (peak A) did not yield crystals.
Data were collected at the Cornell High Energy Synchrotron Source (CHESS), beamline A1. Anomalous data, when present, were collected by the inverse beam method. Four sets of data contributed to the structure determination, collected from crystals in 5 mM KAu(CN)$_2$, 1 mM dimercuriacetate (DMA), 3 mM Au(NO$_3$)$_4$, and no adduct, respectively (Table 2). A heavy-atom partial structure was solved by inspection of KAu(CN)$_2$ anomalous difference Patterson maps, which revealed two heavy binding sites. Heavy-atom positions were then refined using SHARP (Bricogne et al. 2003), and difference electron density maps were used to locate a single heavy-atom position in the DMA derivative. Individual mercury atoms within the DMA molecule could not be resolved due to high B-values, so the heavy-atom compound was modeled as a point scatterer. Other derivatives were not useful for MIR phasing, due either to a lack of isomorphism or the presence of overlapping heavy-atom sites. Density modification was performed after heavy-atom refinement in SHARP using the default solvent flipping protocol (Bricogne et al. 2003).

The kinase domain was built into the density modified maps with reference to the already determined Btk kinase domain structure (PDB ID 1K2P), after separately placing the large and small lobes. The SH3 domain was built by comparison with the c-Src SH3 domain (PDB ID 2SRC).

We generated a homology model for the SH2 domain from the SH2 domains present in PDB entries for Src, Hck and Abl containing SH3-SH2-kinase constructs (PDB IDs: 1FMK, 1KSW, 2SRC, 2PTK, 1AD5, 1QCF, 2HCK, 1OPK). This model was placed by hand, aligning helices $\alpha$A and $\alpha$B into the unambiguous density, then refined as a rigid-body using a 4.0 Å resolution cutoff. Multiple crystal averaging across four distinct data sets (KAu(CN)$_2$, DMA,
Au(NO₃)₄ and native, using the program DMMULTI (Cowtan & Main 1993; Winn et al. 2011), revealed the domain swap in the secondary β-sheet portion of the SH2 domain.

The structure was refined using alternating cycles of manual rebuilding in the program O (Jones et al. 1991) and torsional simulated annealing and grouped B-factor refinement against data to 2.8 Å using the program CNS_SOLVE (Brunger et al. 1998; Brunger 2007). A final refinement of the model was performed using PHENIX (Adams et al. 2002) to incorporate TLS as three groups, defined as residues 214-260, 261-391, and 392-657. These groups were identified by the automated TLS group search procedure and agreed well with the character of the experimental maps. In all refinement runs in which the SH2 domain was present, an initial temperature factor was refined for the entire SH2 domain, and then the SH2 domain was fixed. SH2 domain geometry is thus strictly a result of the geometry of the initial homology model. Data collection and refinement statistics are summarized in Table 2.

Illustrations were made in Pymol (DeLano Scientific).

Structure of the PH-TH-kinase construct of bovine Btk

The bovine PH-TH-kinase construct (residues 1 to 170 connected to residues 384 to 659) was thawed on ice and mixed with 2-fold excess IP₃ (Avanti lipid) and inhibitor CGI1746 (Selleck Inc), and was incubated at 4°C for 30 minutes before setting up trays. IP₃ binds to the protein, but does not activate it. Although present in the crystallization mix, it is not visible in the final electron density (see below). Needle clusters of crystals were obtained by sitting drop vapor diffusion by mixing equal volume of protein solutions at 15 mg/ml with reservoir solution containing 200 mM NaCl, 50 mM MgCl₂ and 20% PEG3350. To prepare seeds for microseeding, needle clusters were harvested and combined from 20 drops and transferred into a 1.5 mL appendorf tube.
The bovine PH-TH-kinase construct with activation loop mutations (Joseph et al. 2013) was prepared in the same way as was the wild-type protein. Equal volumes of protein solution and the reservoir solution containing 200 mM NaCl, 50 mM MgCl\textsubscript{2} and 20% PEG3350 were mixed and were equilibrated in a hanging drop tray for 24 hours at 4°C. Microseeding was then performed by striping the wild-type crystal seeds using a cat whisker into the equilibrated drops. Large rock-like crystal clusters appeared ~4 days after micro-seeding, and single crystals (typical dimensions of 0.20 mm x 0.15 mm x 0.25 mm) can be isolated. For cryo-protection, crystals were soaked in reservoir solution supplemented with 25% glycerol, and flash-frozen in liquid nitrogen and stored at 100K.

Data were collected at the Lawrence Berkeley National Laboratory, Advanced Light Source (ALS), beamline 8.2.1. Data reduction was carried out with the software package HKL2000 (Otwinowski & Minor, 1997). The structure was determined by molecular replacement using PHENIX (Adams et al. 2002) with two search models. These were the structure of the Btk kinase domain that has mutations in the activation loop (see below) and the structure of the PH-TH module. Refinement in PHENIX and COOT (Emsley & Cowtan 2004) yielded the final models. No density for IP\textsubscript{3} is seen in difference electron density maps, even at low contour levels (2σ above the mean in σ weighted |F	extsubscript{o}|−|F	extsubscript{c}| difference maps). Data collection and refinement statistics are summarized in Table 1. Illustrations were made in Pymol (DeLano Scientific).

Structure of the Btk PH-TH module bound to IP\textsubscript{6}

The bovine Btk PH-TH module (residues 1 to 172) sample was thawed at room temperature and was incubated with 5-fold excess IP\textsubscript{6} (Sigma) for 10 minutes. Plate-like
crystals were obtained by sitting drop vapor diffusion by mixing equal volume of protein sample solution and the reservoir solution containing 25% PEG1500 and 0.1 M DL-malic acid pH 5.5. Crystals were then transferred to cryo-protection solution, which is the reservoir solution supplemented with 30% glycerol and 3 mM IP₆.

Data were collected at the Lawrence Berkeley National Laboratory, Advanced Light Source (ALS), beamline 8.2.1. Data reduction was carried out with the software package HKL2000 (Otwinowski & Minor, 1997). The structure was determined by molecular replacement using PHENIX (Adams et al. 2002) with the structure of the PH-TH module as the search model (PDB: 1BTK) (Hyvönen & Saraste 1997). Refinement in PHENIX and COOT (Emsley & Cowtan 2004) yielded the final models. The structural restraint file for IP₆ was generated using REEL in PHENIX (Adams et al. 2002) based on the conformation of IP₆ seen in the structure of Auxin receptor TIR1 (PDB: 2P1P) (Tan et al. 2007). Data collection and refinement statistics are summarized in Table 1. Illustrations were made in Pymol (DeLano Scientific).

Structure of the Btk kinase domain activation loop mutant

Btk kinase domain (residues 395 to 659) with the activation loop mutations (L542M, S543T, V555T, R562K, S564A, and P565S) (Joseph et al. 2013) was thawed at room temperature and was incubated with 2-fold excess CGI1746 (Selleck Inc) and 5-fold excess IP₆ (Sigma) for 10 minutes. Crystals were obtained by sitting drop evaporation methods by mixing equal volume of protein sample solution and the reservoir solution containing 0.2 M ammonium citrate dibasic and 20% PEG 3350. Crystals were then transferred to cryo-protection solution which is the reservoir solution supplemented with 30% glycerol and 3 mM
IP$_6$.

Data were collected at the Lawrence Berkeley National Laboratory, Advanced Light Source (ALS), beamline 8.2.1. Data reduction was carried out with the software package HKL2000 (Otwinowski & Minor, 1997). The structure was determined by molecular replacement using PHENIX (Adams et al. 2002) with the structure of wild-type Btk kinase domain (PDB: 3OCS) (Di Paolo et al. 2011). Refinement in PHENIX and COOT (Emsley & Cowtan 2004) yielded the final models. No density can be located for IP$_6$ in crystals. Data collection and refinement statistics are summarized in Table 1.

**Preparation of autophosphorylated Btk**

Btk samples were diluted to 4 µM using the buffer that contains 150 mM NaCl, 25 mM Tris, pH 7.5 and 5% glycerol. For experiments involving lipid vesicles and inositol phosphates, additional 500 µM lipid and 200 µM inositol phosphates are supplemented into the dilution buffer, respectively, followed by a 10-minute incubation at room temperature.

The diluted samples were mixed with equal volume of the reaction buffer that contains 150 mM NaCl, 20 mM MgCl$_2$, 2 mM ATP, 25 mM Tris, pH 7.5, 2 mM sodium vandate and 5% glycerol, and incubated at room temperature. The final concentration of Btk samples is 2 µM. At each time point, the samples were either frozen in liquid nitrogen and stored at -80°C for the coupled kinase assays, or mixed with equal volume of quench buffer that contains 2X SDS-PAGE buffer supplemented with 100 mM EGTA, followed by incubating at 95 °C in a heat block for 15 minutes for western blot assays.

**Coupled Kinase Assay**
A continuous pyruvate-kinase coupled assay was performed to measure the kinase activity of the proteins as described (Barker et al. 1995), with minor modifications. The ATP concentration was kept at 1 mM in all the assays. The buffer used contains 150 mM NaCl, 10 mM MgCl₂, 25 mM Tris, pH 7.5, 2 mM sodium vandate. The protein concentrations were kept at 1 μM for all Btk constructs. The peptide sequence is ERDINSLYDVMYVDPSEIN, which was derived from residues 746 to 766 of PLC-γ2. The $K_m$ value for phosphorylated full-length Btk for this peptide is $925 \pm 117$ μM, and the peptide concentration was kept at 1 mM in all experiments. The reactions were performed in a 96-well plate and were monitored by SpectraMax (Molecular Devices). The ATP turn-over rate ($v_0$, min⁻¹) was calculated by fitting the data acquired from first 150 seconds using a linear regression model.

**Western blot**

The levels of autophosphorylation of purified Btk samples were monitored using non-specific anti-phosphotyrosine antibody 4G10 (EMDMillipore). The total amount of Btk was monitored using coomassie-blue stained SDS-PAGE. For each set of experiments, 10 μL of each Btk sample was loaded on two 12% SDS-PAGE gels. The two gels were run at 250 V, 400mA for 35 minutes at room temperature.

One gel was transferred to 50 ml fixing buffer containing 50% ethanol and 10% acetic acid, and was heated for 30 seconds in a microwave. The gel is then transferred to 50 ml staining buffer that contains 5% ethanol, 7.5% acetic acid and 0.00025% coomassie brilliant blue R-250, and was heated for another 30 seconds in a microwave. The gel was then left to cool at room temperature in the staining buffer on a rocker for 1 hour, and was then imaged using ImageLab (Bio-Rad).
The other gel was transferred into western blot transfer buffer that contains 25 mM Tris at pH 7.4, 192 mM glycine and 20% methanol, and was incubated at room temperature for 15 minutes. Protein samples were then transferred to PVDF membranes from the gel using the semi-dry transfer device TRANS-BLOT (Bio-Rad). The membranes were then blocked for 1 hour in TBST buffer (20 mM Tris pH 7.5, 150 mM NaCl, 0.1% Tween-20) supplemented with 5% dry milk, and incubated with primary antibody 4G10 (1:2000 dilution) at 4°C overnight. The membranes were then washed 3 times in TBST buffer, and incubated with HRP-linked anti-mouse IgG antibody (1:5000 dilution) in TBST buffer at room temperature for 1 hour. The membranes were then washed 3 times in TBST buffer, incubated with 0.5 mL WesternBright Quantum (Advansta) for 1 minute, and were imaged using ImageLab (Bio-Rad). The western blots were reproduced at least three times and the most representative ones were chose to present.

**Small unilamellar vesicle preparation**

Three lipids, 18:1 DOPS, 18:1 (Δ9-Cis)(DOPC), and 18:1 Pl(3,4,5) P3 (Avanti) were dissolved in chloroform and stored at −20 °C. For liposome preparations, the three lipids are mixed in a test tube at the desired molar ratio (10:75:5), and chloroform was evaporated under a nitrogen atmosphere for 15 minutes, and the samples were transferred to a vacuum desiccator, and dried overnight at room temperature. Dry films were hydrated in buffer containing 25 mM Tris-HCl (pH 7.4) and 100mM NaCl to a concentration of 10 mg/mL.

The hydrated liposomes are heterogeneous in size and the diameters of the liposomes varies from ~50 nm to ≥ 1 μm. To prepare small unilamellar vesicles, hydrated liposomes were subject to seven freeze–thaw cycles using liquid nitrogen, followed by extrusion through
100 nm filters (Avestin). The liposomes prepared using this method have diameters around 100 nm, as measured using negative staining electron microscopy (data not shown).

**Isothermal titration calorimetry**

The isothermal titration calorimetry experiments were performed using MicroCal-autoITC 200 (GE Healthcare). Various Btk samples are diluted in the binding buffer containing 150 mM NaCl, 25 mM Tris pH 7.5 and 5% glycerol to a final concentration of 20 μM. IP$_4$ and IP$_6$ were diluted using the same buffer to a final concentration of 300 μM. Each set of ITC experiments included three samples: 420 μL of Btk protein sample in the cell, 150 μL of IP$_4$ or IP$_6$ at a concentration 300 μM in the syringe, and 500 μL of binding buffer. All samples are stored at 4°C before the titration experiments.

The ITC experiments were performed at 20°C. An initial injection of 0.5 μL was excluded from data analysis, followed by 14 injections of 3 μL each, separated by 180 s with a filter period of 5 s. The protein solution was stirred at 500 rpm over the course of titration. Titration curves were fit with a one-site binding model. The three fitting parameters are stoichiometry, N, association constant, K$_a$ and binding enthalpy, ΔH. The binding entropy is then calculated using formula: ΔS=(ΔH+RTlnK$_a$)/T. Thermodynamic parameters for various Btk constructs binding to IP$_4$ and IP$_6$ are listed in Table 1.

**Molecular dynamics simulations**

Molecular dynamics trajectories were generated using the Gromacs 4.6.2 package (Pronk et al. 2013) using the ff99SB-ILDN force field (Lindorff-Larsen et al. 2010). All simulations were in water, using the TIP3P water model; appropriate counterions (Na$^+$ and Cl$^-$)
were added to neutralize the net charges. After initial energy minimization, the systems were subjected to 100 ps of constant number, volume and temperature (NVT) equilibration, during which the system was heated to 300K. This was followed by a short equilibration at constant number, pressure and temperature (NPT, 100 ps). Finally, the production simulations were performed under NPT conditions, with v-rescale thermostats in Gromacs 4.6.2 respectively, in the absence of positional restraints. Periodic boundary conditions were imposed, and particle-mesh Ewald summations were used for long-range electrostatics and the van der Waals cut-off is set at 10 Å. A time step of 2 fs was employed and the structures were stored every 2 ps. His 143, Cys 154, Cys 155 and Cys 165, which coordinate Zn^{2+} in the PH domain were set to be deprotonated. No additional positional restraints are applied to the Zn^{2+} ion, and its sp³ tetrahedral coordination remains intact during all MD simulations.

The Visual Molecular Dynamics (VMD) analysis toolkit (Humphrey et al. 1996) was used to make some of the room mean square deviation (RMSD) measurements, others were done using PyMol

Homology modeling

The active conformation of the Btk kinase domain was modeled based on that of the Lck kinase domain (PDB: 3LCK) (Sicheri et al. 1997), using Modeller (Eswar et al. 2006). The sequence of the kinase domains of Lck and Btk were aligned by ClustalW (Larkin et al. 2007). We chose the model with lowest Discrete Optimized Protein Energy (DOPE: -0.84) as representative of the active conformation of Btk.

Estimate of membrane-bound Btk concentration
We estimated the effective concentration ($C$) of Btk on a lipid vesicle by the following equation:

\[ C = \frac{N}{V \times N_a} \]

$N$ is the number of Btk molecules on a lipid vesicle, $N_a$ is the Avogadro number and $V$ is the effective volume in which Btk moves on a lipid vesicle. An estimate for $V$ is given by

\[ V = A \times L \]

Here, $A$ is the surface area of a lipid vesicle and $L$ is the length of a Btk molecule, $\sim 100$ Å, as measured from the composite model of full-length Btk. Assuming all the PIP$_3$ molecules (molar fraction, $f$) are bound to Btk, then

\[ N = f \times N_0 \]

where $N_0$ is the total number of lipids on a vesicle. This is given by

\[ N_0 = \frac{A}{A_0} \]

in which $A_0$ is the surface area of a lipid molecule. We assume all lipids have the same surface area for their headgroups, which is roughly 0.6 nm$^2$ (Gulik-Krzywicki et al. 1967). Therefore $C$ is given by:

\[ C = \frac{N}{V \times N_0} = \frac{f \times A}{A \times L \times N_a} = \frac{f}{A_0 \times L \times N_a} \]

In our experiments, $f = 0.05$, $A_0 = \sim 60$ Å$^2$, $N_a = 6.03 \times 10^{23}$ mol$^{-1}$ and $L = \sim 100$ Å, giving a value of 12 mM for $C$. 

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Bruton's tyrosine kinase displays altered ligand binding properties when auto-


Figure 2.
A

model for full-length Btk

B

PH-TH/kinase interface

SH3/SH2-linker interface

SH3/PH-TH interface

SH2/kinase interface

SH2/C-terminal tail interface

Figure 3.
Figure 4.
Figure 6.
Figure 7.
Figure 8.
Figure 9.
Figure 10.