In silico activation of Src tyrosine kinase reveals the molecular basis for intramolecular autophosphorylation

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Abstract
Structural data suggest that important hinge-bending motions of the two lobes that shape the catalytic domain of Src tyrosine kinase, together with reorganization of an alpha helix (helix C), are needed for the activation loop to adopt the catalytically competent conformation. The phosphorylation of a Tyr residue (Tyr-416) in this loop also seems to be essential for enzyme activation. However, no information is available about the dynamics of this activation process. By comparing the inactive and active forms of the catalytic domains of Src and Lck, another member of the Src family, we first identified a short stretch that can act as a hinge for the interlobe motion. The opening of the lobes was then simulated using a targeted molecular dynamics approach. The results obtained suggested that pulling the two lobes apart is not enough to induce the required conformational change in the activation loop. Rather unexpectedly, however, swinging of the lobes situated Tyr-416 in a suitable position for intramolecular autophosphorylation, and further simulation of Tyr-416-phosphorylated Src in the presence of ADP did then result in a conformational change that placed the activation loop in a position similar to that found in the active open conformation of Lck. Taken together, our results establish a physical link between intramolecular autophosphorylation and loop activation.

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1. Introduction
Cellular Src (c-Src) is the normal cellular counterpart of the product of the first oncogene identified from Rous sarcoma virus, v-Src, and also the first protein characterized as being able to phosphorylate tyrosine residues [1]. Src substrates are found in the cytosol or at the inner face of the plasma membrane, or at cell–matrix or cell–cell adhesions. Tyrosine phosphorylation of these proteins can affect their function directly or, alternatively, the phosphorylonyl residues can serve as docking sites for the binding of signaling proteins containing SH2 domains. The resulting complexes then initiate pathways that regulate protein synthesis, gene expression, cytoskeletal assembly and many other aspects of cell function [2]. Because Src family members are involved in many signaling pathways by means of which several surface receptors regulate cell growth and proliferation, their catalytic activity is strictly regulated [3], thus offering unique opportunities for modulation by small molecules in the fight against disease.

Src can also phosphorylate itself at tyrosine residues. Although the major in vivo phosphorylation site on platelet Src was shown to be Tyr-527, which acts as a negative regulator of its kinase activity [4] and is absent in v-Src, the major site of autophosphorylation in vitro is Tyr-416, a residue located at the center of the catalytic domain [5]. Phosphorylation of Tyr-416 in both v- and c-Src results in increased activation of the enzyme [6–8] whereas mutation of this residue to phenylalanine (Y416F) significantly reduces the transforming potential of Src [9]. A tyrosine kinase that specifically phosphorylates Src at Tyr-527, carboxy-terminal Src kinase (Csk), has been identified [10], as have phosphatases that can dephosphorylate this residue [11], highlighting the fact that the phosphorylation status of Tyr-527 is key for in vivo c-Src regulation. However, to the best of our knowledge, no tyrosine kinase specific for Src Tyr-416 has been described yet. Instead, an autophosphorylation mechanism has been proposed based on the finding that the level of phosphorylation of this tyrosine residue correlates with the activity of...
N-terminal lobe and a large α only to facilitate substrate recognition, but also to reorganize
structural mobility in Src regulation. Easily accessible, thus supporting the importance of struc-
tural mobility in Src regulation.

The catalytic domain of Lck (another member of the Src family), and preventing substrate-binding. In an 'open' active con-
formation the two lobes approach each other closely, and suggest considerable interlobe mobility. In the inactive con-
formation the two lobes close together thereby narrowing the cleft
which extends from the base of the catalytic domain. Binding of pTyr-527 to SH2 acts as a safety catch that locks the
catalytic domain under control. As in all known
residues, the activation loop, the phosphorylation of this
residue could induce an important conformational change
responsible for the activation of the enzyme.

The structural domains of Src kinase (and other mem-
bers of the family) are, in order from the N-terminus: the
SH4 (Src homology 4), SH3, SH2 and SH1 domains. SH1 is
the catalytic domain, SH2 and SH3 are both molecular ad-
hesives important for protein–protein interaction, and SH4
plays a role in membrane attachment. SH2 domains bind
phosphotyrosine-containing peptides [15,16] whereas SH3
domains recognize proline-rich peptides [17]. Both SH2 and
SH3 domains were early shown to be involved in the reg-
ulation of the kinase activity of Src [18,19]. The complex
interactions between different domains of c-Src were better
understood when the three-dimensional structures of c-Src
[20], and the Src family tyrosine kinase Hck [21], were solved, both in a catalytically inactive conformation.

Each of these crystal structures revealed a finely adjusted
nanomachine [2] in which all structural elements, i.e., the
SH3 and SH2 domains, the linker between SH2 and SH1
domains, and the carboxy-terminal tail cooperate in order
to keep the catalytic domain under control. As in all known
kinases, the catalytic domain is made up of a small αβ
N-terminal lobe and a large α C-terminal lobe. The linker
joining the SH2 and the catalytic domain adopts a polypro-
line type II helix structure and serves as an adapter to fit to-
gether the N-terminal lobe and the SH3 domain. This confor-
mation places the SH2 domain in a suitable position to bind
the phosphotyrosine residue at the C-terminal tail (pTyr-527)
which extends from the base of the catalytic domain. Bind-
ing of pTyr-527 to SH2 acts as a safety catch that locks the
whole structure in an inactive conformation [2].

The overall structure of the catalytic domain, shaped as
the active site during catalysis. This kind of motion is com-
mon to many other enzymes and protein molecules [23], in
which two or more domains are connected by a few strands
of polypeptide chains that can be considered as hinges [24].
The conformational changes responsible for these motions
are usually limited to the hinge region in so far as the do-
 mains behave as rigid bodies.

Dynamic hinge-bending motions in proteins are not easily
amenable to experimental structural studies but molecular
dynamics (MD) simulations provide a computational alter-
native that can help to gain insight into these processes at
the atomic level. However, these motions take at place at time
scales that are more than one order of magnitude longer than
those currently achieved by state-of-the-art MD simulations.
This is so because the system must surmount the energy bar-
rier that separates the open and the closed forms. Thus, even
though the energy-barrier crossing process itself is normally
quite fast, the time required for random thermal fluctuations
within the system to produce the local atomic momenta re-
quired for overcoming the local energy barrier may be of
the order of milliseconds or longer. In these cases, a tar-
geted MD (tMD) approach [25] can be used to accelerate the
process and simulate the subdomain hinge-bending motion.
This approach has been previously used to study confor-
mational changes in several proteins such as the GroEl chap-
trone [26] and the glutamate receptor ligand-binding core
[27], as well as the coupling between SH2 and SH3 domains
when Src is forced to adopt the open conformation [28].
In the following, we focus our attention on the dynamic
properties of the catalytic domain itself. We first study
its molecular architecture to pinpoint the mobile parts re-
sponsible for the interlobe motion. We then simulate the
hinge-bending motion using tMD to explore how the con-
formational change in the activation loop is triggered, and in
doing so we also probe the feasibility of an intramolecular
mechanism for Tyr-416 autophosphorylation.

2. Results and discussion

2.1. Determination of the hinge regions of Src catalytic
domain

Evidence for a hinge-bending motion associated with the
activation of the Src catalytic domain can be gained from
comparison of the crystal forms of non-active Src [20] and
active Lck catalytic domains [22]. Visual inspection of these
structures shows that, irrespective of changes in interlobe
orientation and activation loop conformation, the overall
structure of each lobe is maintained upon activation (Fig. 1).
Alignment of the catalytic domains of Src and Lck also
shows that not only do they share significant sequence ho-
omology (>64%) but also very high structural homology. In
fact, only 10 out of 263 Cα atoms of the catalytic domain are
in non-equivalent positions, and all of these residues with-
out structural homology precisely belong to the activation
Fig. 1. Comparison of the overall structures of the catalytic domains of the human tyrosine kinase Src in a closed inactive conformation (green ribbon, PDB code 2SRC), and human lymphocyte kinase, Lck, in an open active conformation (red ribbon, PDB code 3LCK).

The root-mean-square deviation (rmsd) between the Cα traces of the Src and Lck catalytic domains, excluding the activation loop, is 2.49 Å, which is higher than that obtained when the structures of Src and Hck catalytic domains (both in an inactive form) are compared (rmsd = 1.0 Å). However, when considered separately, the rmsd between the Cα traces of the N- and C-terminal lobes (excluding the activation loop) of Src and Lck are 1.12 and 0.98 Å, respectively. These results confirm that, despite the overall conformational change, the two lobes do indeed behave mostly as rigid bodies.

More precise information about the mobile parts of the Src catalytic domain can be obtained either by comparing pairs of phi-psi angles for each residue or, more simply, by calculating the differences between equivalent dihedral angles of the Cα traces of ‘active’ Lck and ‘inactive’ Src (Fig. 2):

(i) The first region in which a difference between the dihedral angles of Cα traces of the closed (Src) and open (Lck) forms is observed corresponds to a segment of the Gly-rich loop (residues 273–278; a in Fig. 2) that makes up part of the ATP-binding site (see also Fig. 10).

(ii) The next variation (b1 and b2, in Fig. 2) is detected just before and after the residues corresponding to the only helix present in the N-terminal lobe (residues 304–316), but the helical structure itself (helix C) is completely conserved. This means that helix C can be reoriented without changing its secondary structure. More interesting are the differences in the conformation corresponding to the region that connects the two lobes of the catalytic domain (residues 341–359; c in Fig. 2). This region is adjacent to the loop that is involved in the recognition of the adenine moiety of ATP, and

![Fig. 1. Comparison of the overall structures of the catalytic domains of the human tyrosine kinase Src in a closed inactive conformation (green ribbon, PDB code 2SRC), and human lymphocyte kinase, Lck, in an open active conformation (red ribbon, PDB code 3LCK).](image)

![Fig. 2. Differences in Cα dihedral angles (ΔCα) along the peptide backbone between the open and closed forms shown in Fig. 1. Residue numbering refers to human Src. The horizontal dashed lines represent the mean of the difference for all residues ±2σ.](image)
comprises a short helix flanked by two short strands. This structural element appears suitable for performing the role of a hinge as small changes in the helical angles can give rise to relatively large displacements of the strands (Fig. 11). This peculiar structure, together with the fact that it is located at the interlobe boundary, strongly suggest that the conformational changes in this region could be responsible for the differences found in the relative orientation of the two lobes between the non-active Src structure and the open active conformation of Lck.

(iii) The most remarkable differences, however, are found, as expected, between residues 394 and 425 (regions d1 and d2, Fig. 2). A short segment of this region corresponds to a loop that is in contact with the SH2 domain (d1, residues 294–299), but the segment where the difference is the highest corresponds to the activation loop (d2, residues 405–425).

(iv) The last region for which significant differences are found between the Cα/H9251 trace dihedral angles of closed and open forms (residues 483–496, e in Fig. 2) has no associated function in Src although a homologous region in glycogen synthase kinase, Gsk3, corresponds to the FRATide peptide-binding site [29].

2.2. Stability of the Src holoenzyme

Since both Src and Lck crystal structures have been obtained in the absence of bound ATP or one of its non-hydrolyzable analogues, we placed an ATP molecule in the nucleotide binding site of Src in such a way that it resembled the conformation and orientation found in the crystal structure of cAMP-dependent protein kinase [30]. We also placed two Mg2+ ions in the positions occupied by the two Mn2+ ions in this structure even though it is known that only one is essential for kinase activity [31]. The holoenzyme was then solvated as described in the Section 3, and the behaviour of the whole system was simulated.

No significant conformational changes in the protein were detected during the 2-ns trajectory. The rmsd value of Cu atoms with respect to the initial structure was never higher than 1.25 Å (Fig. 3). At the beginning of the simulation, three water molecules were incorporated to the coordination spheres of the metal ions. As seen for Mn2+ in cAMP-dependent protein kinase, one of the Mg2+ ions is coordinated in a rather regular octahedral coordination sphere by the two oxygens of the β and γ phosphoryl groups of ATP, the two carboxylate oxygens of Asp-404, and two water molecules. The other Mg2+ ion is coordinated in an unusual trigonal bipyramidal geometry by two oxygens from the α and γ phosphoryl groups of ATP, one carboxylate oxygen from Asp-404, the oxygen from the side chain of Asn-391, and a water molecule. These coordination spheres were consistently observed during the whole trajectory, and both distances and angles were similar to those found for Mn2+ in the cAMP-dependent protein kinase crystal structure (data not shown).

The relative position of ATP in the protein did not change with respect to the initial orientation and differs from that found for Ca2+-bound 5′-adenyly-l-mido-triphosphate in the crystal structure of the Src family tyrosine kinase Hck [21] only in the position of the γ phosphate group, as expected from the differences observed in metal ion–phosphate coordination.

2.3. Simulation of Src catalytic domain activation

In order to preserve the characteristics of a typical hinge-bending motion, the overall shapes of the N- and C-terminal lobes were maintained during the tMD simulation by restraining the dihedrals of their Cu traces in the non-mobile regions to the values found in the inactive form of the enzyme, as explained in the Section 3. The hinge motion was ‘activated’ by forcing the Cu trace of the residues making up the hinge region of Src to adopt the conformation found for the equivalent stretch in the open conformation of Lck. Fig. 4a shows the evolution of the rmsd value of Cu atoms belonging to the Src catalytic domain in its complex with ATP/2Mg2+ along the simulation time with respect to the initial modelled structure.
Fig. 4. Monitoring of the hinge-bending motion during interlobe separation along the simulation time. (a) RMSD of Cα atoms belonging to the hinge region (residues 341–359) of Src with respect to equivalent residues in the Lck open active conformation. (b) Interlobe separation measured as the distance between Cα atoms corresponding to residues Pro-333 and Thr-440, each placed in a different lobe. The upper axis in both panels shows the value of the template force constant applied at different times during the simulation.

atoms of the hinge (residues 341–359) with respect to the corresponding atoms in the open ‘active’ conformation of Lck along the simulation time. The adaptive conformational changes in the hinge region are apparent even at very low force constant values and the rmsd becomes progressively and gradually reduced as the force constant applied to the system increases until it reaches a value very close to zero at the end of the simulation.

The hinge-based separation of the N- and C-terminal lobes of the Src catalytic domain was also monitored by measuring the distance between Cα atoms corresponding to residues Pro-333 and Thr-440, each placed in a different lobe (Fig. 4b). This increase in interlobe distance nicely correlates with the extent of the overall conformational change in the hinge region (Fig. 4a). For values above 3.55 kcal mol⁻¹ per hinge atom the distance reaches a plateau, which corresponds to the degree of opening found in ‘active’ Lck. Taken together, these results strongly suggest that interlobe separation in the Src catalytic domain is induced by a conformational change taking place solely in a short polypeptide stretch close to the region that connects the two lobes.

The aim of this work was to understand whether the hinge-bending motions of the lobes are associated with the conformational changes in the activation loop that result in Src catalytic domain activation. In the closed inactive form, the activation loop is packed close to the substrate-binding cleft (Fig. 1) and not far from the active site of the enzyme. Some residues of this loop establish interactions with helix C. For example, Arg-409 forms a salt bridge with Glu-310 whereas in the open active structure the carboxylate of this glutamate residue forms a salt bridge with the ε-amino group of Lys-295, a residue that is involved in the recognition of the phosphate groups of ATP. Current hypotheses based on structural data [20,21] suggest that the interlobe opening motion could give rise to the reorientation of helix C allowing the conformational changes in the activation loop to take place.

Fig. 5. Monitoring of the conformational changes affecting the activation loop. (a) Distance between the Cα atoms of Asn-391, placed at the cleft, and the phosphorylatable Tyr-416 located on the activation loop. (b) Distance between the phenolic oxygen of Tyr-416 and the Mg²⁺ ion that is coordinated by the β and γ phosphoryl oxygens of ATP. The upper axis in both panels shows the value of the template force constant applied at different times during the simulation.
This rearrangement was monitored by the increase in the distance between the Ca atoms of Asn-391, a residue that is involved in the binding of the ATP/2Mg$^{2+}$ complex, and the phosphorylatable Tyr-416 at the activation loop itself (Fig. 5a). However, under our simulation conditions, the activation loop did not swing away from the cleft. On the contrary, this interresidue distance decreased, suggesting that lobe separation favours the packing of the activation loop within the cleft. The interaction between residues of the activation loop and helix C, most notably the salt bridge between Arg-409 and Glu-310, was not disrupted during the simulation either. As a consequence, the motion of the N-terminal lobe dragged the side chains of the activation loop residues deeper into the cleft. To illustrate this, Fig. 5b shows the distance between the phenolic oxygen of Tyr-416 and the Mg$^{2+}$ ion that is coordinated by the $\beta$ and $\gamma$ phosphoryl oxygens of ATP. At the end of the TMD simulation, by the time the two lobes are as separated as in the 'active' conformation (Fig. 6), the phenolic oxygen of Tyr-416 is only 2 Å away from the Mg$^{2+}$ indicating that it has been incorporated into its coordination sphere by displacing a water molecule. Similarly, a second water molecule that also was coordinating the Mg$^{2+}$ ion has been simultaneously replaced by one of the carboxyl oxygen of Asp-386 (Fig. 7). Coincidentally, the Mg$^{2+}$ that is coordinated by the phosphorylatable oxygen of Tyr-416 has been postulated to be the essential ion for catalysis [32], and Asp-386 is a residue that is conserved in all protein kinases and is most likely involved in the catalytic mechanism [33]. The final structure obtained for the activated Src holoenzyme is thus strongly suggestive of an intramolecular mechanism for autophosphorylation at Tyr-416.

2.4. Molecular dynamics simulation of the Tyr-416-phosphorylated kinase domain and ADP

Since the hydroxyl group of Tyr-416 was placed in a position suitable for intramolecular phosphorylation, we used the final structure of the previous complex to model the catalytic domain with a phosphotyrosine at position 416 and ADP in place of ATP, i.e., the immediate products of the reaction. The dynamic behaviour of this new complex was then simulated for 2 ns under the same conditions as before.
The motion of the activation loop was studied by monitoring the separation between the Cα atoms of Asn-391 and pTyr-416 (Fig. 8). A sharp increase of this distance can be seen after about 50 ps, and a steady increase is observed until a plateau is reached at around 0.8 ns. This change comes mostly as a consequence of both the repulsion between the phosphate groups of ADP and the phosphotyrosine and the tendency of these groups to be surrounded by a solvation shell. The final distance is now very similar to that observed in the active open conformation of Lck. In fact, at the end of the simulation, the activation loop does indeed adopt the more extended conformation (Fig. 9) that is observed in the crystal structure of Lck (Fig. 1). Furthermore, in good analogy to what is seen in this experimental structure, the phosphate group of pTyr-416 is involved in two salt bridges with the guanidinium groups of Arg-385 and Arg-404, and another salt bridge is formed between the side chains of Glu-310 and Lys-295 (Fig. 9). It is precisely the interaction...
between residues equivalent to Glu-310 and Lys-295 that has been related to the activation, not only of tyrosine kinases [20,21], but also of Ser/Thr kinases [30].

3. Computational methods

3.1. Atomic data

The atomic coordinates of human tyrosine kinase, c-Src [20], and human lymphocyte kinase, Lck [22], were obtained from the protein data bank (PDB codes: 2SRC and 3LCK, respectively). Both proteins were structurally aligned with the DALI software [35]. Differences in dihedral angles between the Cα traces of open and closed forms were measured, with positive and negative values representing, respectively, clockwise and anticlockwise increments relative to the closed form.

Neither ATP nor ADP nucleotides were present in any of these PDB files. The ATP molecule and two Mg2+ ions were placed in the X-ray Src structure by superimposing those residues involved in ATP binding with equivalent residues from the crystal structure of cAMP-dependent protein kinase (PDB code: 1CDK) which has been co-crystallized with the non-hydrolyzable ATP analogue adenylylimidodiphosphate [30]. Electrostatic potential-derived charges for ATP and ADP (Supplementary Material) were obtained with the RESP methodology [36] using a 6-31G* basis set as implemented in the Gaussian-98 program [37]. Standard Lennard–Jones AMBER parameters [38] and a charge of +2 were used for Mg2+ ions. We used reported bonded and non-bonded parameters for the phosphotyrosine residue [28]. A spherical cap of TIP3P water molecules [39] centered on the Cα of Asp-404, located at the interlobe cleft, was used to simulate the aqueous environment. The 28 Å radius of this water cap was long enough to cover all residues of the activation loop in both active and inactive conformations.

3.2. Energy refinement

To adapt the crystal structure to the AMBER force field [38], and to gradually refine the modelled Src-ATP-2Mg2+ complex, the SANDER module in AMBER [40] was employed for energy minimization using a cutoff of 10.0 Å. First, only the water dipoles were allowed to reorientate in the electric field of the holoenzyme; then the ATP and Mg2+ ions were also allowed to relax; subsequently, the amino acid side chains were also included in the energy refinement, and finally the whole structure was energy-minimized although protein backbone atoms were restrained to their initial positions with a force constant of 2 kcal mol−1 Å−2. Each refinement stage consisted of 2000 steps of steepest descent and 8000 steps of conjugate gradient energy minimization.

3.3. Molecular dynamics simulations

The refined holoenzyme was used as input for the subsequent MD simulations using the same cutoff as above. In
a 6 ps heating phase the temperature was raised from 0 to 298 K, and velocities were reassigned at each new temperature according to a Maxwell-Boltzmann distribution. ATP and Mg$^{2+}$ ions were completely free to move. The Co atoms of the C-terminal lobe were restrained to their initial positions with a force constant of 2 kcal mol$^{-1}$ Å$^{-2}$ (positional restraints) except for residues in the activation loop. The dihedral angles of the Co trace of the N-terminal lobe region considered as a rigid body (see below) were restrained to those found in the initial structure (although a fluctuation of ±20° was allowed) by means of a harmonic potential with a force constant of 300 kcal mol$^{-1}$ rad$^{-2}$ (dihedral restraints).

Identical restraints were applied to the hinge region for the first 500 ps of the simulation but they were removed thereafter, and the MD simulation was continued during 2 ns at the same temperature. The SHAKE algorithm was applied to constrain all bonds to their equilibrium values so that an integration time step of 2 fs could be used throughout. The list of non-bonded pairs was updated every 25 steps, and coordinates were saved every 2 ps.

The tMD approach consisted of forcing the Co atoms of the hinge region of the ‘closed’ structure to adopt the crystallographic positions of equivalent atoms in the ‘open’ structure by means of a harmonic potential with a force constant that was progressively increased (0 25 kcal mol$^{-1}$ Å$^{-2}$ per run) during 10 consecutive runs of 60 ps each. The incremental use of this term, as previously reported for another system [27], tends to avoid biasing the trajectory by overcoming artifactual energy barriers that could deprive the simulation of any physical sense.

For the simulation of pTyr-416-Src in the presence of ADP, the dihedral angles defining the Co trace of hinge-region residues were restrained with a force constant of 3.55 kcal mol$^{-1}$ Å$^{-2}$ per atom in order to keep the interlobe cleft open.

4. Conclusions

The overall molecular architecture of the catalytic domain of all protein kinases, which is shaped as two lobes connected by a short polypeptidic strand, reveals a remarkable plasticity [34] and strongly suggests considerable interlobe mobility. Comparison of Src in a closed inactive form and Lck in an open active conformation provides structural evidence that the motion of the lobes is an important event in the activation of the catalytic domain. The lobes behave essentially as rigid bodies, with most of the conformational changes being restricted to the activation loop and a short interlobe region that is best described as a hinge. Thus, activation of the catalytic domain of Src kinase has all the characteristics of a typical hinge-bending motion similar to that described for a number of other proteins [22].

Following the precise identification of this hinge region in Src, a MD approach was used to pull the two lobes of the catalytic domain apart. Monitoring both the rmsd of the hinge region relative to the open active conformation and the interlobe distance along the simulation showed that the conformational change taking place in this small stretch was enough for the two lobes to swing apart. However, the activation loop did not reach a conformation similar to that found in the active form. On the contrary, this segment was packed even deeper within the cleft. In this location, the phenolic oxygen of the phosphorylatable Tyr-416 was dragged into the coordination sphere of one of the ATP-bound Mg$^{2+}$ ions. This placed the hydroxyl group at short distances of both the carboxylate of the catalytic Asp-386 and the γ-phosphate group of ATP, in a position suitable for attack. When the simulation proceeded with the immediate products of the phosphotransfer reaction, i.e., a phosphorytrosine at position 416 and ADP in the nucleotide binding site, the Src catalytic domain finally achieved a conformation similar to that observed in the crystal structure of Lck.

Taken together, our results support the feasibility of an intramolecular mechanism for the autophosphorylation of Tyr-416 [5,13], and further suggest that the conformational change in the activation loop responsible for kinase activation is triggered by this event rather than by the mere opening of the lobes.

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