

The loops of the N-SH2 binding cleft do not serve as allosteric switch in SHP2 activation

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The Src-homology-2 domain-containing phosphatase SHP2 is a critical regulator of signal transduction, being implicated in cell growth and differentiation. Activating mutations cause developmental disorders and act as oncogenic drivers in hematologic cancers. SHP2 is activated by phosphopeptide binding to the N-SH2 domain, triggering the release of N-SH2 from the catalytic PTP domain. Based on early crystallographic data, it has been widely accepted that opening of the binding cleft of N-SH2 serves as the key "allosteric switch" driving SHP2 activation. To test the putative coupling between binding cleft opening and SHP2 activation as assumed by the allosteric switch model, we critically reviewed structural data of SHP2, and we used extensive molecular dynamics (MD) simulation and free energy calculations of isolated N-SH2 in solution, SHP2 in solution, and SHP2 in a crystal environment. Our results demonstrate that the binding cleft in N-SH2 is constitutively flexible and open in solution and that a closed cleft found in certain structures is a consequence of crystal contacts. The degree of opening of the binding cleft has only a negligible effect on the free energy of SHP2 activation. Instead, SHP2 activation is greatly favored by the opening of the central β -sheet of N-SH2. We conclude that opening of the N-SH2 binding cleft is not the key allosteric switch triggering SHP2

 $molecular\ dynamics\ simulations\ |\ phosphatases\ |\ allosteric\ regulation$

S rc-homology-2–containing protein tyrosine phosphatase 2 (SHP2), encoded by the *PTPN11* gene, is a classical nonreceptor protein tyrosine phosphatase (PTP). It has emerged as a key downstream regulator of several receptor tyrosine kinases (RTKs) and cytokine receptors, functioning as a positive or negative modulator in multiple signaling pathways (1–4). Germline mutations in the human *PTPN11* gene have been associated with Noonan syndrome and with Noonan syndrome with multiple lentigines (formerly known as LEOPARD syndrome), two multisystem developmental diseases (5–16). Somatic *PTPN11* mutations were also linked with several types of human malignancies (17–25), such as myeloid leukemia (7, 26–30).

The structure of SHP2 includes two tandemly arranged Src homology 2 domains (SH2), called N-SH2 and C-SH2, followed by the catalytic PTP domain, and a C-terminal tail with a poorly characterized function (Fig. 1) (31). The SH2 domains are structurally conserved recognition elements (32) that allow SHP2 to bind signaling peptides containing a phosphorylated tyrosine (pY) (33). The N-SH2 domain consists of a central antiparallel β -sheet, composed of three β -strands, denoted βB , βC , and β D, flanked by two α -helices, denoted α A and α B (Fig. 1A). The peptide binds in an extended conformation to the cleft that is perpendicular to the plane of the β -sheet (34). N-SH2 contains a conserved affinity pocket covered by the BC loop (also called phosphate-binding loop or pY loop), whose interaction with pY increases the binding of the peptide by 1,000-fold relative to unphosphorylated counterparts (35). Residues downstream of the pY bind to a more variable, less conserved site, which confers binding specificity and which is flanked by the EF and BG loops (36).

In 1998, the first crystallographic structure of SHP2 at 2 Å resolution (Protein Data Bank [PDB] ID 2SHP) revealed that the N-SH2 domain tightly interacts with the PTP domain (Fig. 1B), so that the DE loop of N-SH2, thereafter indicated as "blocking loop," occludes the active site of PTP, forcing SHP2 into an autoinhibited "closed" state (37). SHP2 structures of the active "open" state, obtained for the basally active, leukemia-associated E76K mutant, showed an alternative relative arrangement of N-SH2 and PTP that exposed the active site of the PTP domain to the solvent (Fig. 1C) (38). Unexpectedly, in both the autoinhibited and the active state of SHP2, the N-SH2 ligand binding site is exposed to solvent and does not directly interact with PTP or other domains (37–39). Hence, the need for an allosteric mechanism was proposed, according to which the binding of a phosphopeptide triggers a series of structural rearrangements in the N-SH2 domain to drive its release from PTP and the consequent activation of SHP2 (37, 39).

The comparison of the autoinhibited structure of SHP2 (PDB ID 2SHP) (37) with the existing structures of the isolated N-SH2 domain, either in the absence of (PDB ID 1AYD) (34) or in complex with a phosphopeptide (PDB ID 1AYA, 1AYB, 4QSY) (34), showed that the EF and BG loops of the N-SH2 domain may undergo large conformational changes (Fig. 1B). In the autoinhibited structure of SHP2, the N-SH2 domain shares surface complementarity with the PTP catalytic site, but, as a result of the displacement of the EF loop toward the BG loop, it also contains a closed binding cleft that renders the N-SH2 domain unable to accommodate the C-terminal part of

Significance

SHP2 is a multidomain protein, playing an important role in upregulating cellular processes such as cell survival, proliferation, and programmed cell death. SHP2 mutations cause developmental disorders and are found in many cancer types, including neuroblastoma, breast cancer, and leukemia. In healthy cells, SHP2 mainly takes an autoinhibited, inactive form, and SHP2 is activated upon binding of phosphopeptides to the N-SH2 domain. For the past two decades, the widening of the binding cleft upon peptide binding has been considered the key event driving SHP2 activation. Here, by analyzing crystallographic data and molecular simulations, we demonstrate that the binding cleft in N-SH2 is, instead, already open and accessible in solution, and its degree of opening does not influence SHP2 activation.

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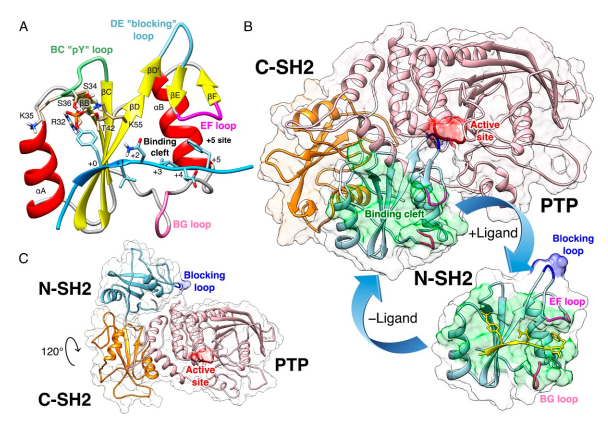


Fig. 1. (*A*) Cartoon representation of the N-SH2 domain in complex with the IRS-1 pY895 peptide (34). The peptide, shown in cyan, comprises the phosphotyrosine and residues from position +1 to +5 relative to the phosphotyrosine (see labels). Functionally important loops are highlighted in color: BC "pY" loop (green), DE "blocking loop" (light blue), EF loop (magenta), and BG loop (deep pink). The phosphotyrosine binds the site delimited by the pY loop and the central β-sheet (βB, βC, βD strands). EF and BG loops delimit the binding cleft (+5 site), where the peptide residue in position +5 is settled. (*B*) Crystal structure of autoinhibited SHP2 (37). The N-SH2 domain (cyan cartoon) blocks the catalytic site (red) of the PTP domain (pink) with the blocking loop (blue). The N-SH2 domain is connected to PTP in tandem with the homologous C-SH2 domain (orange). Closure of the N-SH2 binding cleft (green region), the change in shape of the N-SH2 domain, which accompanies binding of phosphopeptide (yellow), perturbs surface complementarity for the PTP active site, thus promoting N-SH2 dissociation from the PTP domain. (*C*) Crystal structure of SHP2^{E76K} (38). The open conformation reveals a 120° rotation of the C-SH2 domain, relocation of the N-SH2 domain to a PTP surface opposite the active site, and a solvent-exposed catalytic pocket.

the phosphopeptide, in contrast to the isolated N-SH2 domain that exhibits an open binding cleft. Therefore, peptide binding seems only compatible with the conformation of isolated N-SH2, but not with the conformation of N-SH2 in autoinhibited SHP2 (37, 39).

Because 1) the closure of the binding cleft in N-SH2 has been ascribed to its interaction with PTP and 2) the conformation adopted by the EF loop correlates with the activation of SHP2 in available structural data, the EF loop has been suggested as the key allosteric switch that drives the release of N-SH2 from PTP (37, 39). In light of that, conformation selection (9, 21, 40) and induced fit (41) models were put forward for the molecular events leading to functional activation of SHP2; however, both models consider the conformational change involving the EF loop as the key mechanism that drives SHP2 opening (9, 21, 40, 41). In conclusion, it has been widely accepted that the N-SH2 binding cleft, delimited by the EF and the BG loop, plays the role of an allosteric switch for the activation of SHP2 (39, 42). Accordingly, the binding of a ligand at the binding cleft of the N-SH2 domain would induce a transmitted conformational change that prevents PTP domain binding on the other side of N-SH2, and vice versa (37, 39).

However, the allosteric switch model does not explain how the signal, coming from the displacement of the EF loop, is propagated to the rest of the protein (39). In addition, considering that the EF loop might be flexible, the comparison of crystallographic

structures does not provide the energetic penalties involved in the motion of the EF loop and, consequently, the degree of destabilization of the N-SH2/PTP complex upon the binding cleft opening (39). Hence, the role of the EF loop as the key allosteric switch has not been rationalized in energetic terms.

Recently, an allosteric interaction in N-SH2 has been proposed as an alternative mechanism of SHP2 activation (43). Molecular dynamics (MD) simulations have shown that the N-SH2 domain may adopt two distinct conformations, denoted as α - and β -states, which differ primarily in the conformation of the central β -sheet. In the α -state, the central β -sheet is open, adopting a Y-shaped structure; in the β -state, the central β -sheet is closed, adopting a parallel structure. The MD simulations suggested that the β -state of N-SH2 stabilizes the N-SH2/PTP contacts and, hence, the autoinhibited SHP2 conformation. In contrast, the α -state drives the N-SH2 dissociation and SHP2 activation. Notably, the α - β model of activation rationalized modified basal activity and responsiveness to ligand stimulation of certain mutations at codon 42 (15). However, the α - β model seems to contrast with the previously suggested allosteric switch model.

To resolve this discrepancy, we revisited the allosteric switch model. We critically reviewed available crystallographic data of SHP2 in the autoinhibited state. In addition, we used MD simulations, free energy calculations, and enhanced sampling techniques to reveal the conformational dynamics of the binding cleft delimited by the BG and the EF loop in the isolated N-SH2 domain in water, in SHP2 in water, and in SHP2 in a crystal environment. Our results suggest that the binding cleft of N-SH2 is constitutively flexible and the effect of its degree of opening on the activation of SHP2 is negligible. In addition, free energy calculations revealed that, in the crystal environment, the closure of the binding cleft is not due to the allosteric interaction with the PTP domain, but instead a result of the crystal contacts affecting the binding cleft conformation.

Results

N-SH2 Domain Loops Are Characterized by Structural Disorder. In crystal structures, the EF loop and the BG loop have been generally modeled such that the binding cleft of the N-SH2 domain is inaccessible to the ligand (16, 37). To test whether the closed binding cleft is supported by available data, we first evaluated Ramachandran outliers, electron densities, and residue-wise crystallographic R-factors (Z-scores), as well as crystal contacts in the loop regions. The evaluation of the first resolved crystallographic structure of autoinhibited SHP2 (2SHP) (37), performed by MolProbity (44), revealed that the BG loop (residues 84 to 98) adopts a configuration compatible with a closed binding cleft only at the cost of significant energetic penalties. In particular, the Ramachandran angles of the residues Asn⁹² ($\phi = 58.4^{\circ}$; $\psi =$ 118.9°) and Asp⁹⁴ ($\phi = -42.0^{\circ}$; $\psi = 95.5^{\circ}$) represent outliers, being represented in less than 0.01% of the samples in the Protein Data Bank; in the case of His⁸⁴ ($\phi = -145.7^{\circ}$; $\psi =$ 64.1°), Glu⁹⁰ ($\phi = -87.7^{\circ}$; $\psi = -109.2^{\circ}$), and Gly⁹³ ($\phi = 73.9^{\circ}$; $\psi = -46.0^{\circ}$), the conformations are allowed yet disfavored, being represented in only 1.24, 0.07, and 1.4% of the samples, respectively. In the light of the allosteric switch model, these unfavorable conformations may be justified considering that the N-SH2 domain structure is forced by PTP in a tensed conformation, which may be turned into a relaxed conformation only upon ligand binding or upon the release from PTP (37, 39-41). However, inspection of the electron density map showed that the electron densities at the BG loop, the EF loop, and the pY loop were much lower compared to the density of the nearby β -strands (SI Appendix, Fig. S1), indicating increased disorder and therefore a reduced structural definition of these loops.

The 2SHP model was refined to a crystallographic R-factor of 0.2 with an R_{free} of 0.27 (37), characterizing a structure with overall acceptable quality (45). However, the R-factor is a global measure of the model accuracy. In proteins, some regions and strands may be intrinsically disordered such that they cannot be resolved by X-ray diffraction irrespective of the resolution and the overall quality of the experimental data. For this reason, the Z-score has been used as a measure of local model accuracy (46, 47). The Z-score is given by $Z = (R - \langle R \rangle_{res})/\sigma_{res}$ (46–48), where $\langle R \rangle_{\rm res}$ and $\sigma_{\rm res}$ are respectively the expected R-factor and the standard deviation of the R-factor values, calculated with the same amino acid in structures within the same resolution range. For instance, for the case of 2SHP with a resolution of 1.95 Å, the R-factor of each residue is compared with the expected R-factor and standard deviation detected from the same kind of residue in all structures of the database with resolution between 1.8 and 2.0 Å (48). A large, positive spike in a Z-score plot implies that a residue has an R-factor value that is considerably worse than that of the average residue of the same type in structures determined at similar resolution (48). Z-scores are considered as outliers if greater than 2 (46, 48).

The structures of the N-SH2 domain belonging to the two SHP2 chains, A and B, in the 2SHP asymmetric unit are depicted in Fig. 24 (37). The backbone is colored according to the Z-score values reported in Fig. 2B. Evidently, the regions colored in red corresponding to the BG and the pY loop are poorly resolved,

with Z-score values far above the threshold. Similarly, for the EF loop the Z-scores are barely fair in the case of chain B, whereas in chain A the corresponding Z-scores are good.

The allosteric switch model formulation was based on the conformation adopted by the BG and the EF loop in the 2SHP crystal structure (37); thus the validation of the model would require that those loops be univocally defined. However, the analysis of the electron density map and the Z-scores reveal that the loops are poorly resolved compared to the rest of the protein. Instead, the loops have been modeled without adequate electron density and forced into a geometry with significant structural outliers.

The BG Loop Is Generally Poorly Resolved in Crystal Structures of Autoinhibited SHP2. Together with the 2SHP structure (37), we computed Z-scores for other, more recent crystal structures of autoinhibited SHP2, comprising either wild type or functional mutants (16, 49, 50). Table 1 lists the crystal structures considered in this study, spanning resolutions from 1.87 to 2.7 Å. Fig. 3 presents the Z-scores in color scale, for functionally important loop regions, corresponding to the pY loop (Ser³⁴-Asp⁴⁰, dark green), the EF loop (Tyr⁶⁶–Glu⁶⁹, purple), and the BG loop (Gly⁸⁶–Glu⁹⁷, pink). In most of the structures, the residues of the BG loop are either missing (blank spaces) or modeled without adequate electron density (yellow-red squares), confirming that the BG loop is often disordered and therefore poorly resolved by X-ray diffraction. Likewise, the pY loop is sometimes missing or poorly resolved, although it is less affected by structural disorder than the BG loop. In contrast, the short EF loop is marginally affected by structural disorder, being resolved in nearly all crystal structures, although Z-scores for residue Tyr66 are often poor.

Taken together, the Z-scores detected in many crystal structures confirm that loops represent flexible regions of the N-SH2 domain, in particular the BG loop. Notable exceptions are a number of high-quality crystal structures (4DGP, 4OHH, 5I6V) (16, 49, 50), in which the loops have been clearly resolved. These structures with well-resolved loops enabled us to further analyze a putative effect of crystal packing on loop conformations.

N-SH2 Interacts with Other Copies of SHP2 in Crystal Structures. In the crystal environment, proteins do not adopt the same biological assembly according their physiological conditions, but instead every protein is surrounded by other copies of the same protein, establishing close and rather extensive interactions that stabilize the crystal in a particular space group symmetry. If functionally relevant or even flexible regions are involved in such crystal contacts, the conformations of those regions must be interpreted with care.

Fig. 4A shows the surrounding of the N-SH2 domain for the structure 4DGP, which produced good Z-scores even for the BG, EF, and pY loops (16). Evidently, the N-SH2 domain does not bind only the PTP domain belonging to the same chain (colored in white), but it also interacts with a number of other protein replicas (colored in blue, purple, green, and pink). Specifically, the pY, EF, and BG loops, which would be solvent exposed under physiological conditions, are in contact with other chains. Hence, the conformations of these loops are likely influenced by such interactions and not exclusively by the binding of the PTP domain on the other face of N-SH2.

In particular, the Lys³¹ side chain in the BG loop forms salt bridges with charged residues of another replica, whereas the motion of the Tyr⁶⁶ side chain in the EF loop is sterically hindered by the presence of other bulky side chains. Even the pY loop, which in absence of a phosphate group is typically modeled in a partially closed conformation, corresponding to an intermediate conformation between the α - and the β -state (43),

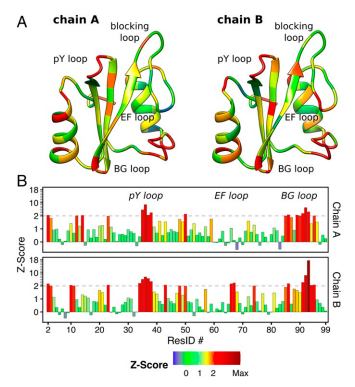


Fig. 2. (*A*) Structures of the N-SH2 domain belonging respectively to the chains A and B of the asymmetric unit reported in the crystal structure of autoinhibited SHP2 (PDB ID 2SHP) (37). The backbone is colored on the base of the R-factor Z-scores reported in *B*. The outlier threshold of 2 is indicated by dashed lines.

interacts with acidic side chains of Glu³¹³ and Glu³⁴⁸ belonging to another protein chain. Therefore, the partially closed loops in the N-SH2 domain might have been stabilized by crystal contacts.

Crystal Packing in Autoinhibited SHP2 Hinders the N-SH2 Motions.

According to the allosteric switch model, opening of the binding cleft would drive the release of N-SH2 from the PTP domain. Hence, a positive correlation in crystal structures might be expected between 1) cleft opening, given by the Gly⁶⁷ $C\alpha$ -Lys⁸⁹ $C\alpha$ distance d_{cleft} , and 2) release of the DE blocking loop from the catalytic PTP loop, given by the Asp⁶¹ $C\alpha$ -Ala⁴⁶¹ $C\alpha$ distance d_{block} . On the other hand, according to the α - β model, a positive correlation might be expected between the opening of the central β -sheet and the distance d_{block} . However, despite a significant spread of d_{block} in available crystal structures of autoinhibited SHP2, no such correlation exists. For instance, d_{cleft} and d_{block} are even (poorly) anticorrelated (Fig. 4B), in disagreement with the expectation from the allosteric switch model. Also, the spread of the N-SH2 central β -sheet lacks the correlation with d_{block} (SI Appendix, Fig. S2).

These data further indicate that crystal packing largely hinders conformational motions of N-SH2 in autoinhibited SHP2 involved in SHP2 activation, and correlations cannot be detected irrespective of the activation mechanism proposed. Hence, 1) to fully resolve the effect of crystal packing on N-SH2 dynamics, and 2) to quantify the influence of N-SH2 dynamics on SHP2 activation, we employed MD simulations in solution and in a crystal.

The N-SH2 Domain Binding Cleft Is Constitutively Flexible. According to the allosteric switch model, the opening and the closure

of the N-SH2 binding cleft is coupled with SHP2 activation or with stabilization of autoinhibited SHP2 (37, 39). Considering that the activation of SHP2 presumably involves a free energy difference of several tens of kilojoules per mole (41, 43), it is mandatory to evaluate the energy required to open the binding cleft in different conditions.

To quantify the intrinsic flexibility of the binding cleft, we used umbrella sampling to compute the potential of mean force (PMF) (also referred to as "free energy profile") of the binding cleft opening for the isolated N-SH2 domain (Fig. 5 A and B). As a reaction coordinate, we used the distance d_{cleft} defined as the $C\alpha\text{--}C\alpha$ distance between Gly⁶⁷ (belonging to the EG loop) and Lys89 (belonging to the BG loop). The PMF covers conformations of a completely inaccessible cleft ($d_{\text{cleft}} = 5 \text{ Å}$; Fig. 5A, cyan cartoon) up to a wide-open cleft ($d_{\text{cleft}} = 14 \text{ Å}$; Fig. 5A, purple cartoon). In Fig. 5B, the PMF shows a single free energy minimum at ~11 Å corresponding to an open cleft, in line with the 1AYD crystal structure of apo N-SH2 (34). This finding is further supported by previous MD simulations of isolated N-SH2 (43), thereby clearly confirming that the cleft is open in the isolated N-SH2 domain.

Notably, the PMF further shows that the binding cleft is highly responsive to external perturbations, as only a small energy of 6 to 10 kJ/mol is required to close the binding cleft into a configuration found in crystal structures of autoinhibited SHP2 with well-resolved loops (e.g., 4DGP.A, 4OHH.A, 5I6V.B; cf. Fig. 3) (16, 49, 50). Therefore, the EF and the BG loops lining the N-SH2 binding cleft are constitutively flexible, and the binding cleft rearrangement in isolated N-SH2 requires only a small amount of free energy.

Binding Cleft Opening Is Favored in Autoinhibited SHP2 in Water. To reveal the effect of the PTP allosteric interaction on the N-SH2 binding cleft, we computed the PMF of the binding cleft opening for SHP2 in water, again along the distance $d_{\rm cleft}$ (Fig. 5 C and D). Critically, for all umbrella windows, the $C\alpha$ - $C\alpha$ distance $d_{\rm block}$ between Asp⁶¹ (belonging to the N-SH2 blocking loop) and Ala⁴⁶¹ (belonging to the PTP catalytic loop) fluctuated around 6 Å (*SI Appendix*, Fig. S3), mainly below the experimental distances observed in autoinhibited structures of SHP2 (Fig. 4B). Therefore, the N-SH2 domain remained closely bound to PTP throughout calculations, and the final PMF in Fig. 5D represents the opening of the binding cleft in autoinhibited SHP2.

Table 1. Experimental data of autoinhibited SHP2 crystal structures

PDB ID	Space group	Resolution (Å)	R-factor	R_{free}	Description
2SHP	P12 ₁ 1	2.00	0.199	0.270	Wild type
4DGP	P2 ₁ 2 ₁ 2	2.30	0.199	0.255	Wild type
4DGX	P2 ₁ 2 ₁ 2	2.30	0.212	0.266	Y279C
4GWF	P12 ₁ 1	2.10	0.211	0.243	Y279C
4H1O	P2 ₁ 2 ₁ 2	2.20	0.211	0.233	D61G
4H34	P2 ₁ 2 ₁ 2	2.70	0.227	0.242	Q506P
4NWF	P2 ₁ 2 ₁ 2 ₁	2.10	0.267	0.309	N308D
4NWG	P2 ₁ 2 ₁ 2 ₁	2.45	0.281	0.326	E139D
40HD	P2 ₁ 2 ₁ 2	2.70	0.206	0.271	A461T
40HE	P2 ₁ 2 ₁ 2	2.51	0.190	0.256	G464A
40HH	P2 ₁ 2 ₁ 2	2.70	0.206	0.266	Q506P
40HI	P2 ₁ 2 ₁ 2	2.20	0.210	0.262	Q510E
40HL	P12 ₁ 1	2.40	0.194	0.259	T468M
516V	P12 ₁ 1	1.87	0.203	0.232	F285S
5IBM	P12 ₁ 1	2.18	0.192	0.240	S502P
5IBS	P12 ₁ 1	2.32	0.189	0.238	E76Q

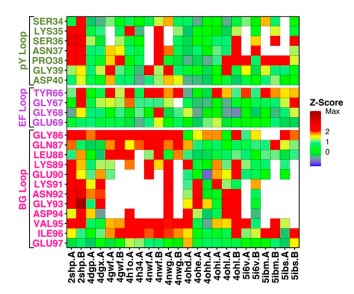


Fig. 3. Tile plot of the R-factor Z-scores, calculated for crystal structures of autoinhibited SHP2. Z-scores are reported in color scale for each residue (blank spaces indicate missing residues), belonging to functionally important loop regions, corresponding to the pY loop (Ser³⁴–Asp⁴⁰, dark green), the EF loop (Tyr⁶⁶–Glu⁶⁹, purple) and the BG loop (Gly⁸⁶–Glu⁹⁷, pink).

Surprisingly, even in tight interaction with PTP, a wide-open binding cleft of N-SH2 corresponds to the free energy minimum (Fig. 5D). This result disagrees with the allosteric switch model (37, 39), which instead expects a closed and inaccessible binding cleft when the N-SH2 is bound to PTP. In addition, the PMF shows that a relatively small amount of free energy of 3 to 9.5 kJ/mol is required to close the binding cleft toward conformations found in crystal structures (Fig. 5D, black dots). This suggests that, on one hand, the binding cleft remains flexible and susceptible to structural rearrangements in the case of ligand binding. On the other hand, the energy involved in

binding cleft rearrangement is presumably many times smaller than the energy required to destabilize the N-SH2/PTP interface, promoting SHP2 activation. In fact, previous molecular dynamics simulations suggested that the complete displacement of N-SH2 away from the PTP active site requires an energy in the order of 70 kJ/mol (41, 43). Hence, it is unlikely that rearrangement of the binding cleft is sufficient to drive SHP2 activation.

Taken together, the PMF shows that 1) even in complex with PTP, the binding cleft of N-SH2 is mostly open and accessible to ligands and 2) the binding cleft is not tensed and, therefore, rearrangement of the binding cleft alone is insufficient to drive SHP2 activation.

Crystal Contacts Hinder the Binding Cleft Opening. The PMF of SHP2 in solution suggests that the binding cleft is more open in solution than expected from available crystal structures (previous paragraph). To shed light on the origin of these discrepancies, we computed the PMF of binding cleft opening for SHP2 in the crystal environment of the 4DGP structure, for which the loops have been well resolved (Fig. 5*E*; cf. Fig. 3). Simulations of a protein in the crystal reveal the influence of the crystal packing on protein structure and dynamics by comparing the results with simulations performed with the same system in solution. Such comparisons are still rare in the literature because researchers are often more interested in the properties of a protein in aqueous solution, corresponding to the most common experimental and physiological conditions (51–54).

The PMF reveals a clear effect of the crystal environment on the binding cleft conformation (Fig. 5 D and F). Namely, the most stable conformation in the crystal corresponds more closely to the crystal structure with a closed binding cleft (i.e., 4DGP) (16), whereas the binding cleft opening is strongly hindered. Hence, the tendency of the binding cleft to adopt a closed and inaccessible conformation in crystal structures is likely a bias due to the crystal contacts between protein replicas and not a consequence of an allosteric mechanism imposed by N-SH2 binding to the PTP active site.

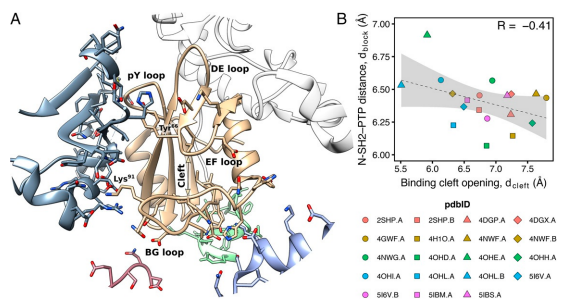


Fig. 4. (A) Cartoon representation of the surrounding of the N-SH2 domain in the crystal structure of autoinhibited SHP2 (PDB ID 4DGP) (16). The PTP domain belonging to the same protein chain is colored in white. Other protein replicas, interacting with the central N-SH2 domain (colored in tan), are depicted respectively in blue, purple, green, and pink. (B) Correlation between the N-SH2 binding cleft opening, d_{cleft} (Gly⁶⁷ Cα–Lys⁸⁹ Cα distance) and the N-SH2 blocking loop distance from the catalytic PTP loop, d_{block} (Asp⁶¹ Cα–Ala⁴⁶¹ Cα distance), as taken from crystal structures of autoinhibited SHP2, comprising either wild type or functional mutants (16, 37, 49, 50).

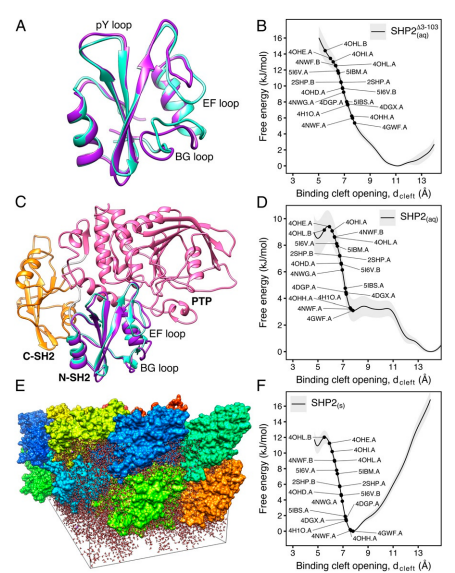


Fig. 5. (A) Simulation snapshot of the N-SH2 domain in water with the binding cleft either closed (cyan cartoon) or open (purple cartoon). Water is not shown for clarity. (B) PMF of the binding cleft opening, d_{cleft} , for the N-SH2 domain in water. Black dots indicate the degree of cleft opening found in various crystal structures. (C) Simulation snapshot of autoinhibited SHP2 in water with the binding cleft either closed (cyan cartoon) or open (purple cartoon). C-SH2 and PTP domains are shown as orange and pink cartoons, respectively. Water is not shown for clarity. (D) PMF of the binding cleft opening, d_{cleft} , for autoinhibited SHP2 in water. (E) Simulation system of SHP2 in the crystallographic quadruple unit cell of the 4DGP structure. The system contains 16 SHP2 copies shown in different colors and water shown as red/white sticks. Some crystal contacts in this structure are shown in Fig. 4. (F) PMF of the binding cleft opening, d_{cleft} , for autoinhibited SHP2 in the crystal environment of E.

Binding Cleft Opening Does Not Promote the Activation of SHP2. Based on the observation that the closed binding cleft is found in crystal structures in correspondence to the autoinhibited SHP2, whereas the crystal structures of the isolated, peptide-bound N-SH2 exhibited a full open, accessible binding cleft, it was hypothesized that the EF loop, together with the BG loop, may serve as an allosteric switch in SHP2 activation (37, 39). According to the allosteric switch model, the interaction with the PTP active site forces the N-SH2 binding cleft into a closed, inaccessible conformation that prevents ligand binding (switch off); binding of a phosphopeptide would push the N-SH2 binding cleft toward an open, fully accessible conformation, with the consequence of weakening the interactions between the blocking loop and the catalytic site, promoting the N-SH2 domain release from PTP (switch on) (37, 39).

To definitely test such a model, we used umbrella sampling to compute the PMF for a critical step of SHP2 activation, that is, the release of the N-SH2 domain from the PTP active site, with the N-SH2 binding cleft restrained either in closed (switch off) or in open (switch on) conformation (Fig. 6). As a reaction coordinate for SHP2 activation, we used the center-of-mass distance between the backbone atoms of the blocking loop and the catalytic PTP loop. Upon pulling the simulation system along this coordinate, the N-SH2 domain moved from its position in the

autoinhibited state (*SI Appendix*, Fig. S4) to different positions on the PTP surface, thereby exposing the active site to the solvent as required for SHP2 activation (Fig. 6 *A* and *B* and Movie S1). Among independent simulation runs, the final position of N-SH2 differed, suggesting that activating SHP2 may involve different pathways in conformational space (*SI Appendix*, Fig. S5).

These final positions do not represent the fully active state of SHP2, which is reached only upon the binding of a bisphosphoryl tyrosine-based activation motif (BTAM) peptide to both the N-SH2 and C-SH2 domains. In fact, the binding of a single phosphopeptide to the N-SH2 domain triggers only a weak activation, since the N-SH2 domain remains in the neighborhood of the catalytic site (36). Only the binding of another phosphopeptide to C-SH2, connected to the first phosphopeptide by a linker of suitable length (55), drives the complete dissociation of the N-SH2 domain from the catalytic site and, thereby, the full activation (36). Hence, the pulling simulations and the PMFs explore early stages of activation but not the complete activation pathway.

In disagreement with the allosteric switch model, the two PMFs for "switch off" (Fig. 6C, yellow line) and "switch on" (Fig. 6C, red line), are almost identical, demonstrating that a switch of the binding cleft does not drive dissociation of the N-SH2 domain from PTP. The only perceptible, small difference at proximal distances is the slight right shift of the switch-on PMF

minimum, likely ascribable to the different tilt of the helix αB , whose effects have been predicted from crystal structure analysis (37). However, such perturbations do not further compromise the interface of the complex at larger distances.

In Fig. 6D we report the $\Delta\Delta \hat{G}$ difference between the two profiles ($\Delta G_{\rm on} - \Delta G_{\rm off}$). The flat profile (Fig. 6D, violet line), whose average is virtually null (Fig. 6D, blue dashed line), confirms that the binding cleft conformation alone has virtually no influence on SHP2 activation. To test whether instead the α - β model rationalizes SHP2 activation, the $\Delta\Delta G$ profile of activation was previously computed with the N-SH2 domain restrained either in the α - or in the β -state (Fig. 6D, green line) (43). Evidently, the $\Delta\Delta G$ values become more and more negative as the distance between N-SH2 and PTP increases, demonstrating that the release of N-SH2 is greatly favored by the α -state.

In conclusion, PMFs indicated that the propensity of the N-SH2 domain to release PTP is essentially unresponsive to the conformation adopted by its binding cleft. Instead, the opening of the central β -sheet, as adopted by the α -state, leads to a larger rearrangement of the N-SH2 shape, therefore promoting SHP2 activation.

Discussion

The conformational transition of disordered regions, occurring upon ligand binding or chemical modifications, is a recurrent molecular mechanism in protein regulation (42, 56, 57), which is why, for a long time, the flexible loops in the binding cleft of N-SH2 have been considered as molecular switches in an allosteric mechanism controlling SHP2 activation (39, 42). However, in this study we have brought several arguments in opposition to

the allosteric switch model: 1) the structural disorder of the N-SH2 loops in crystal structures; 2) the constitutive flexibility of the binding cleft in solution; 3) the unexpected opening of the binding cleft, occurring even in autoinhibited SHP2 in water; 4) the steric hindrance, induced by the crystal contacts, forcing the binding cleft into a closed conformation; and finally 5) the missing destabilization of the N-SH2/PTP interface upon binding cleft opening.

Experimental evidence brought up in favor of the allosteric switch model stated that 1) the phosphopeptide binding is generally stronger to the isolated N-SH2 domain than to the N-SH2 domain as part of SHP2 (21) and 2) the mutations destabilizing the N-SH2/PTP interface not only increase the SHP2 basal activity, but also result in a larger phosphopeptide binding affinity, albeit these mutations do not cluster at the N-SH2 binding site and apparently they do not affect the structure of N-SH2 (40). Thus, the direct correlation between the basal activity of SHP2 and the ligand binding affinity to N-SH2 seems coherent with the allosteric switch model, because an eventual enhancement of the binding cleft opening would consequently promote the ligand binding to N-SH2. However, such experimental evidence would be potentially in agreement with any other allosteric mechanism involving an equilibrium between a stabilizing conformation and any activating conformation of N-SH2, independently of the structural details governing such an equilibrium. As a matter of fact, the assumption of an equilibrium between a stabilizing conformation, blocked by the interaction with PTP, and an activating conformation, induced by the binding of a phosphopeptide, inevitably leads to the conclusion that any weakening of the N-SH2/PTP interaction not only penalizes the stabilizing

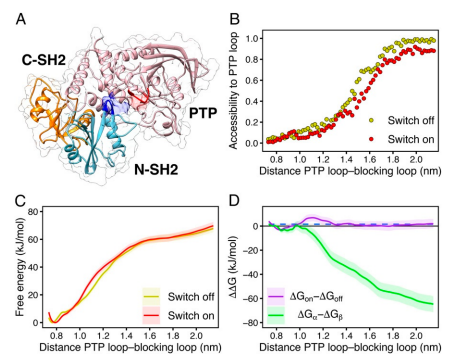


Fig. 6. (A) Exemplary open and active structure of SHP2 as obtained by pulling simulation. The N-SH2 domain (cyan cartoon) moved from its position in the autoinhibited state to a different position relative to the PTP (pink) and C-SH2 domains (orange), leaving the catalytic site (red) solvent exposed. (β) Accessibility of the PTP catalytic site versus the distance between the blocking loop and the catalytic PTP loop used as reaction coordinate in pulling simulation, defined as the backbone centers-of-mass distance between residues 60 to 62 and residues 460 to 462. The N-SH2 binding cleft was restrained either in closed (switch off, yellow dots) or in open (switch on, red dots) conformation. The accessibility of the PTP catalytic site was defined as the ratio of the SAS areas of the PTP loop, obtained including or excluding the N-SH2 domain. The probe radius was 2.38 Å corresponding to a phosphate ion. (C) PMFs of SHP2 opening with the N-SH2 binding cleft restrained either in closed (switch off, yellow line) or in open (switch on, red line) conformation. (D) $\Delta\Delta G$ profiles of SHP2 opening, calculated from the difference of the PMFs for "switch on" relative to the "switch off" conformation in C (violet line), and, for comparison, for the α -state relative to the β -state (green line), calculated from PMFs in ref. (43). Dashed blue line represents the average $\Delta\Delta G$ value of the $\Delta G_{on} - \Delta G_{off}$ profile.

conformation in autoinhibited SHP2, but also indirectly makes N-SH2 more prone to bind a ligand. Such a conclusion is valid in the case of the allosteric switch model, as well as for the α - β model

Certain gain-of-function pathogenic mutations in SHP2 do not cluster at the N-SH2/PTP interface, where they could destabilize the N-SH2/PTP interactions, but seem to have more subtle, allosteric effects, since their impact on SHP2 function cannot be rationalized by mere steric effects. A typical example is the Noonan syndrome (NS)-causing Thr42Ala substitution that replaces a conserved threonine in the central β -sheet (15). Because Thr⁴² forms an H-bond with the phosphotyrosine in wild-type N-SH2, one might expect that Thr42 contributes to the stability of the N-SH2/phosphopeptide complex. However, the substitution with alanine leads to an increase in phosphopeptide binding affinity (15), as documented by the dramatically enhanced catalytic activity of the SHP2A42 mutant upon stimulation with a BTAM peptide (15). Notably, the allosteric switch model has been so far unable to convincingly explain the pathogenicity of the Thr42Ala substitution, in terms of either enhancement of the binding cleft opening or formation of additional favorable contacts with the phosphopeptide (15). Instead, owing to the reduced propensity of alanine for forming β -sheets compared to threonine (58), the α - β model attributes to the Thr42Ala substitution the effect of destabilizing the central β -sheet, thereby rendering N-SH2 more prone to adopting the activating α -state (43).

Analogous to the opening/closure of the binding cleft, the opening and closure of the central β -sheet in N-SH2 are detectable by comparing the crystal structures of autoinhibited SHP2 with crystal structures of the isolated N-SH2 domain bound to a phosphopeptide (34, 37). Thus, it does not come as a surprise that the opening of the central β -sheet emerges for the α - β model as the key determinant of the N-SH2 reshaping, which leads to the loss of complementarity for PTP and thereby to the activation of SHP2. However, according to the α - β model, a coupling exists between the opening of the central β -sheet with both the closure of the pY loop and the tightening of the cleft at the +5 site. Therefore, we can conclude that the role of the binding cleft in N-SH2 is not to directly cause the loss of complementarity for PTP, but to induce, through the recognition and clamping of an activating phosphopeptide, a more extensive, concerted rearrangement in N-SH2 structure, which ends with the central β -sheet opening.

In conclusion, SHP2 is activated by an allosteric mechanism. However, our analysis suggests that the key transition triggering SHP2 activation is not the opening of the N-SH2 binding cleft but instead the opening of the central β -sheet of the N-SH2 domain, as captured by the $\alpha\!-\!\beta$ model. The $\alpha\!-\!\beta$ model includes, apart from central β -sheet opening, also structural elements that were previously considered by the allosteric switch model. As such, the $\alpha\!-\!\beta$ model can be considered as a generalized formulation of an allosteric mechanism, leading to a better comprehension of the structural transitions governing the activation of SHP2.

Materials and Methods

Analyses on Crystallographic Structures. Structures of autoinhibited SHP2, in its wild-type form or mutants, were taken from the Research Collaboratory for Structural Bioinformatics (RCSB) Protein Data Bank: 2SHP (37), 4DGP (16), 4DGX (16), 4GWF, 4H1O, 4H34, 4NWF, 4NWG, 4OHD (50), 4OHE (50), 4OHH (50), 4OHI (50), 4OHL (50), 5I6V (49), 5IBM (49), and 5IBS (49). Dihedral angles were validated using the MolProbity web server (44). Electron densities were inspected with Coot (59). R-values and Z-scores were obtained from the residue-wise electron density using the Uppsala Electron Density Server (48).

MD Simulations of the N-SH2 Domain in Solution. The initial atomic coordinates (residues 3 to 103) were derived from crystallographic structure 1AYD (34). N-SH2 was put at the center of a dodecahedron box, large enough to contain the domain and at least 0.9 nm of solvent on all sides, and then it

was solvated with 5,577 explicit TIP3P (transferable intermolecular potential 3 points) (60) water molecules.

MD simulations were performed with the GROMACS software package (61), using the AMBER99SB*-ILDNP force field (62, 63) if not stated otherwise. Long-range electrostatic interactions were calculated with the particle-mesh Ewald (PME) approach (64). A cutoff of 1.2 nm was applied to the direct-space Coulomb and Lennard-Jones interactions. Bond lengths and angles of water molecules were constrained with the SETTLE algorithm (65), and all other bonds were constrained with LINCS (66). The pressure was set to 1 bar using the Parrinello–Rahman barostat (67). The temperature was controlled at 300 K using velocity rescaling with a stochastic term (68).

The solvent was relaxed by energy minimization followed by 100 ps of MD at 300 K, while restraining protein coordinates with a harmonic potential. The system was then minimized without restraints and its temperature thermalized to 300 K in 10 ns, in a stepwise manner. After an equilibration run of 100 ns while N-SH2 was left completely unrestrained, the protein was equilibrated keeping the binding cleft closed. The $C\alpha$ distance between Gly^{67} (belonging to the EG loop) and Lys^{89} (belonging to the BG loop) was chosen as a reaction coordinate for cleft opening, denoted d_{cleft} . An umbrella potential (k = 10,000 kJ·mol $^{-1}$ ·nm $^{-2}$) was progressively introduced in 10 ns and maintained for other 100 ns, so that the d_{cleft} was finally constrained at 5 Å.

To maintain the N-SH2 domain with the central β -sheet parallel and closed, a flat-bottom potential (k = 10,000 kJ·mol $^{-1}$ ·nm $^{-2}$), was introduced between the carbonyl C atom of Gly 39 and the backbone N atom of Asn 58 , whose equilibrium distance was 4 Å; such a flat-bottom potential, acting only for distances larger than 4.4 Å, remained inactive during most parts of the simulations, and it simply prevented the N-SH2 domain, rarely reaching the transition state, from further relaxing in a conformation with the central β -sheet completely open.

The reference distance of $d_{\rm cleft}=7$ Å was considered for the closed binding cleft [corresponding to the distance in 2SHP (37) crystal structure of autoinhibited SHP2], while a reference distance of 12 Å was chosen for the open binding cleft [corresponding to the distance in 1AYA (34) crystal structure of N-SH2 complexed with a phosphopeptide]. A pulling simulation (69, 70) was carried out for gradually opening the binding cleft over 180 ns. The pull force constant was set to 1,000 kJ·mol $^{-1}\cdot$ nm $^{-2}$, whereas the reference $d_{\rm cleft}$ was gradually increased from 5 to 14 Å with a velocity of 5×10^{-6} nm/os.

To further improve the conformation sampling during these pulling simulations and to obtain more independent starting frames for umbrella sampling (see below), we coupled the pulling simulations with simulated tempering (71). In line with previous studies (71, 72), simulated tempering simulations were carried out at constant volume, and temperature steps of 10 K between 300 and 380 K were applied. Temperature changes were controlled according to the Metropolis algorithm to obtain canonical ensembles at all temperatures. The initial weights were chosen following Park and Pande (72). Temperature transitions were attempted every 1 ps, and the weights were updated throughout the whole simulation according to the Wang–Landau adaptive weighting scheme (73).

The PMFs were obtained by umbrella sampling and the weighted histogram analysis method (WHAM) (74, 75), using 46 windows, equispaced by 0.2 Å. Initial configurations for umbrella sampling were taken from the previous simulated tempering pulling simulations by a cluster analysis, thereby ensuring that the umbrella simulations were triggered from the most representative conformations. Accordingly, the configurations were collected from the subensemble at 300 K, divided into 46 groups on the basis of the umbrella window interval they belong to, and clustered using the GROMOS clustering method (76). The clustering cutoff was chosen on the basis of the root-mean-square deviation distribution in each group, picking the value corresponding to the first relative maximum along the abscissa. The umbrella force constant was set to 1,000 kJ·mol⁻¹·nm⁻², and a sampling of 1.5 $\,\mu s$ was performed for each window. Statistical errors of the PMFs were estimated with the Bayesian bootstrap of complete histograms (74), thereby considering only complete histograms as independent, yielding a maximum error of less than 2 kJ/mol.

MD Simulations of SHP2 in Solution. The initial coordinates of SHP2 were taken from the autoinhibited conformation (2SHP) (37). The protein was positioned at the center of a dodecahedral box, large enough to contain the protein and at least 0.9 nm of solvent on all sides, and solvated with $\sim\!23,000$ explicit water molecules (60) and three Na⁺ ions. The structure was minimized and thermalized to 300 K using the procedure described above for N-SH2. A simulation of 1 μs was performed to ensure a well-equilibrated autoinhibited structure in solution.

Next, the protein was equilibrated maintaining the binding cleft respectively either closed or open. An umbrella potential on d_{cleft} (k = 10,000 kJ·mol⁻¹·nm⁻²) was progressively introduced within 100 ns and maintained for another 100 ns, so that d_{cleft} was finally constrained either at 5 Å or at 14 Å. The procedure was repeated four times providing the same number of independent equilibrated configurations of SHP2 with the N-SH2 binding cleft either in closed or in open conformation. To avoid the opening of the central β -sheet in the N-SH2 domain, a flat-bottom potential was applied as described above.

Four pulling simulations were spawned either for gradually opening or for closing the binding cleft over 180 ns (eight simulations in total). The pull force constant was set to 1,000 kJ·mol⁻¹·nm⁻², whereas the reference d_{cleft} was changed with a velocity of 5×10^{-6} nm/ps. Simulated tempering (71) was used to enhance sampling, using a temperature range from 300 to 380 K $\,$ in steps of 5 K. To avoid the premature activation of SHP2 at higher temperatures and to maintain its autoinhibited structure, a flat-bottom potential (k = 10,000 kJ·mol⁻¹·nm⁻²) was introduced between the C α atom of Asp⁶¹ (belonging to the N-SH2 blocking loop) and the $C\alpha$ atom of Ala⁴⁶¹ (belonging to the PTP catalytic loop); this flat-bottom potential, acting only when the distance d_{block} was larger than 6.4 Å, was removed after the end of the simulated tempering pulling simulations. The PMF along d_{cleft} was computed as described above, except that each umbrella window was simulated for 250 ns.

MD Simulations of SHP2 in Crystal Environment. The initial coordinates of SHP2 were taken from the autoinhibited conformation (4DGP) (16). This SHP2 crystal, which belongs to the $P2_12_12$ space group containing four symmetry-related molecules, has the following orthorhombic unit cell parameters: a = 5.485 nm, b = 22.110 nm, c = 4.036 nm, and $\alpha = \beta = \gamma = 90^{\circ}$. Missing or incomplete residues (strands Leu²³⁶–Gln²⁴⁵, Gly²⁹⁵–Val³⁰¹, Phe³¹⁴– Pro³²³; residues Met¹, Thr², Lys²³⁵, Arg⁵²⁷, Arg⁵²⁸; and the His-tagged tail Leu^{529} -His 536) were modeled by Molecular Operative Environment (MOE) (77) using the AMBER12:EHT force field and enabling the periodic system for the P2₁2₁2 space group.

The starting coordinates of the single-crystal unit cell were obtained by applying the P2₁2₁2 symmetry transformation. Next, the quadruple unit cell was generated by shifting the entire box along the x axis and/or the z axis, resulting in an orthorhombic simulation box, with edges of 10.970, 22.110, and 8.072 nm, containing 16 protein molecules. We used multiple unit cells as a simulation box to fully exclude spurious couplings between the motion of the protein and its periodic images and to exclude unrealistic long-range correlations.

The AMBER99SB* -ILDNP force field (62, 63) was used, while a flat-bottom potential (k = $10,000 \text{ kJ} \cdot \text{mol}^{-1} \cdot \text{nm}^{-2}$) was introduced between the carbonyl C atom of Gly³⁹ and the backbone N atom of Asn⁵⁸. The system was solvated with 26,908 explicit water molecules (60) and 32 Na^+ ions. The solvent box was generated by two successive solvent additions, each followed by a solvent relaxation session. The solvent was relaxed by energy minimization followed by 100 ps MD at 300 K, while restraining protein positions with a harmonic potential. The system was then energy minimized without restraints and its temperature brought in a stepwise manner to 300 K in 10 ns.

To remove the conformational correlation between protein replicas, an equilibration run of 50 ns at 300 K was performed, followed by 50 ns of simulated tempering (71), with temperature ranging from 300 to 360 K in steps of 2 K, and another equilibration run of 50 ns.

Starting from the last system structure, two pulling simulations were spawned either for gradually opening or for gradually closing the

binding cleft in each of the 16 protein replicas (32 simulations in total). The pull force constant was set to 1,000 kJ·mol⁻¹·nm⁻², whereas the reference d_{cleft} was changed with a velocity of 5×10⁻⁶ nm/ps. Simulated tempering (71) was used to enhance sampling, using a temperature range from 300 to 360 K in steps of 2 K. Again, a flat-bottom potential was used to avoid the opening of the central β -sheet and immediately removed after the end of the simulated tempering pulling simulations. The PMF along d_{cleft} was computed as described above, except that each umbrella window was simulated for 20 ns.

MD Simulations of the Activation of SHP2, with the Binding Cleft Restrained into the Closed or Open Conformation. The initial coordinates of SHP2 were taken from the autoinhibited conformation (2SHP) (37). The protein was positioned at the center of a dodecahedral box and solvated with \sim 23,000 explicit water molecules (60) and three Na⁺ ions. Because certain Amber force fields in conjunction with TIP3P have been reported to overstabilize protein-protein contacts (78), we used the CHARMM36m force field (79) for simulations of SHP2 activation. Hydrogen atoms were described as virtual sites, allowing a time step of 4 fs. The structure was minimized and thermalized to 300 K using the procedure described above. A simulation of 1 μs was performed to ensure a well-equilibrated autoinhibited structure in solution.

Next, the binding cleft of the N-SH2 domain was restrained either in closed ($d_{cleft} = 5 \text{ Å}$) or in open conformation ($d_{cleft} = 12 \text{ Å}$), using a harmonic restraint along d_{cleft} (k = 1,000 kJ·mol⁻¹·nm⁻²). Opening of the central β sheet was excluded using a flat-bottom potential, as described above. N-SH2 was pulled away from PTP using 600-ns pulling simulations at constant pull velocity (69). The center-of-mass distance between the backbone atoms of the blocking loop (residues 60 to 62) and the catalytic PTP loop (residues 460 to 462) was chosen as the reaction coordinate. That distance was pulled from 0.73 to 2.15 nm. To obtain statistically independent pathways for SHP2 opening, eight pulling simulations were carried out for each N-SH2 binding cleft state, restrained either in closed or in open conformation (16 simulations in total). To further improve the conformation sampling, we coupled the pulling simulations with simulated tempering (71), using a temperature range from 300 to 400 K in steps of 5 K.

The PMFs for the opening of SHP2 were obtained using umbrella sampling and WHAM (74, 75). Two PMFs were computed with the N-SH2 binding cleft restrained either in closed or in open conformation. For each profile, 72 windows, equispaced by 0.2 Å, were used. Initial configurations for umbrella sampling were taken from the previous simulated tempering pulling simulations by a cluster analysis (76). Because breaking of N-SH2/PTP contacts involves larger forces, we used an increased umbrella force constant set to 4,000 kJ·mol⁻¹·nm⁻². Each window was sampled for 100 ns. Statistical errors of the PMFs were estimated using the Bayesian bootstrap of complete histograms (74).

The normalized solvent-accessible surface (SAS) of the PTP loop (Fig. 6B) was defined as $A/A_{\rm max}$, where A denotes the SAS computed from the MD frame and A_{max} the SAS from the same MD frame after removing the N-SH2 domain.

Data Availability. All study data are included in this article and/or SI Appendix.

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