A conserved allosteric element controls specificity and activity of functionally divergent PP2C phosphatases from Bacillus subtilis

Reversible phosphorylation relies on highly regulated kinases and phosphatases that target specific substrates to control diverse cellular processes. Here, we address how protein phosphatase activity is directed to the correct substrates under the correct conditions. The serine/threonine phosphatase SpoIIE from Bacillus subtilis, a member of the widespread protein phosphatase 2C (PP2C) family of phosphatases, is activated by movement of a conserved α-helical element in the phosphatase domain to create the binding site for the metal cofactor. We hypothesized that this conformational switch could provide a general mechanism for control of diverse members of the PP2C family of phosphatases. The B. subtilis phosphatase RsbU responds to different signals, acts on a different substrates, and produces a more graded response than SpoIIE. Using an unbiased genetic screen, we isolated mutants in the α-helical switch region of RsbU that are constitutively active, indicating conservation of the switch mechanism. Using phosphatase activity assays with phosphoprotein substrates, we found that both phosphatases integrate substrate recognition with activating signals to control metal-cofactor binding and substrate dephosphorylation. This integrated control provides a mechanism for PP2C family of phosphatases to produce specific responses by acting on the correct substrates, under the appropriate conditions.

Protein kinases and phosphatases regulate diverse biological processes by controlling the phosphorylation state of target proteins, requiring that they select specific substrates and act on them at the correct time and place. Protein kinases have conserved allosteric elements that control kinase activity in response to regulatory inputs and have deep active site grooves that facilitate recognition of cognate substrates (1–3). Although all protein kinases belong to a single superfamily of related enzymes, protein phosphatases come from multiple evolutionarily unrelated lineages (4). Determining how each family of phosphatases achieves regulation and specificity will reveal mechanistic differences that distinguish each phosphatase family and may suggest specific strategies for targeting phosphatases with small molecules.

We focus here on the protein phosphatase 2C (PP2C) family of serine/threonine phosphatases, members of which are found across all kingdoms of life (5, 6). Phosphatases from the PP2C family are precisely regulated to control a diverse range of processes including stress response, development, virulence, and cell growth and death in all kingdoms of life, but their mechanisms of regulation and substrate specificity are unknown (4, 7). To address this question, we developed a system to compare substrate selection and regulation of two PP2C phosphatases, SpoIIE and RsbU, which coexist in the same bacterial cell but respond to different signals and act on distinct phosphoprotein substrates.

SpoIIE controls the developmental program of spore development in the bacterium Bacillus subtilis (Fig. 1A) (8). After asymmetric cell division, SpoIIE is stabilized against proteolysis and is activated by multimerization in small cells (9). The multimerization, stabilization, and activation of SpoIIE are mediated by its unique regulatory domain. Upon activation, SpoIIE dephosphorylates SpoIIAA to activate a cell-specific transcription factor (Fig. 1A) (8, 10). We recently found that SpoIIE is activated by rotation of an α-helical element at the base of the phosphatase domain to form the binding site for the catalytically essential metal ions (we refer to the metal ions as “metal cofactor” and use Mn2+ as the metal for all experiments in this study, Fig. 1B and C) (11). The regulatory α-helical element we described is a conserved feature of the PP2C domain, and we hypothesized that it is preserved as a regulatory switch for phosphatases in this family (11).

The phosphatase RsbU has a PP2C domain that is related to SpoIIE (17% sequence identity and 40% similarity) and includes a region predicted to form the α-helical switch (Fig. S1, A and B) but has an unrelated N-terminal regulatory domain (Fig. 1B and Fig. S1A) and responds to different signals to initiate the general stress response in B. subtilis (Fig. 1A) (12). The general stress response describes a broad transcriptional response that is activated by diverse bacterial species in response to changing conditions and coordinates processes including virulence, antibiotic resistance, and biofilm formation (13). A common mechanism for activation of a general stress response relies on a PP2C phosphatase that recognizes the initiating signal. This appears to be the most evolutionarily ancient and broadly
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Figure 1. SpolIE discriminates between cognate and noncognate substrates. A, the pathways controlling the activity of the transcription factors σF and σB are diagrammed. The phosphatases SpolIE (IIE) and RsbU dephosphorylate paralogous substrate proteins SpoIIAA-P (AA-P) and RsbV-P (V-P). The activity of RsbU is dependent upon activation by RsbT. In the unphosphorylated state, SpoIIAA and RsbV bind to anti-sigma factor proteins SpoIIAB (AB) and RsbW (W), displacing and activating the sigma factors. B, domain diagrams of SpolIE and RsbU. The N-terminal regulatory regions of these proteins are unique, while the PP2C domains share 17% identity. The regulatory region of SpoIIE contains an N-terminal degradation tag (black) followed by 10 transmembrane domains (gray) (aa 49–320). Both SpolIE (magenta) (aa 320–589) and RsbU (blue) (aa) contain unrelated regulatory domains. The PP2C domain of SpolIE (aa 590–827) has been crystallized (11) and contains an α-helical regulatory switch (orange), which coordinates the binding of metal cofactor along with metal binding residues D795, D746, and D628 (red) through positioning the backbone carbonyl of G629. The PP2C domain of RsbU (aa 121–335) has not been crystallized but is predicted to contain a similar α-helical regulatory switch. The metal binding residues show conservation between the two proteins (the detailed alignment can be seen in Fig. S1). C, structures of SpolIE in the inactive (top left, PDB ID: 5MQH) and active (top right, PDB ID: 5UCG) states. The α-helical switch is represented in orange and the side chains that coordinate divalent cations in the active site are shown as sticks. Manganese ions are modeled in the active site of SpolIE from the structure of metal-bound RsbX and are represented as spheres. Valine 697 is shown as green spheres. Below each structure is a cartoon representation of the active site as it is controlled by the switch. In the inactive state, residue V697, shown in green, is modeled to be in a hydrophobic pocket, while in the active site, it is exposed to a hydrophilic patch. D, SpolIE590–827 phosphatase activity with SpoIIAA-P (red) or RsbV-P (blue) as the substrate. Reactions with SpoIIAA-P as the substrate were performed as multiple turnover reactions with 0.1 mM SpolIE590–827 and varying concentrations of SpoIIAA-P. Reactions with RsbV-P were performed as single turnover reactions (due to the extremely slow rates of reaction), using trace concentrations of RsbV-P with varying concentrations of SpolIE590–827. All reactions contained 10 mM MnCl2. Data are fit to the linear equation observed rate = (kcat/Km)[SpoIIAA-P or SpolIE]/(Km + [SpoIIAA-P or SpolIE]). F, a reaction coordinate diagram summarizing the data from panels D and E. PP2C, protein phosphatase 2C.

conserved function of the PP2C family of phosphatases across the bacterial kingdom. To activate the general stress response under conditions of environmental stress in B. subtilis, RsbU binds to an activating protein (RsbT) and dephosphorylates RsbV. RsbV then activates the transcription factor σB, which controls the general stress response transcriptional program in B. subtilis (Fig. 1A) (12). It is essential that SpolIE and RsbU act with high fidelity on their cognate substrates (14); aberrant dephosphorylation of RsbV-P and activation of the general stress response during sporulation blocks spore development (15), and dephosphorylation of SpoIIAA-P in nonsporulating cells is toxic.

In addition to sensing different signals and acting on different substrates, SpolIE and RsbU exhibit different
response profiles. Whereas SpoIIE initiates a switch-like developmental transition, RsbU initiates a graded response with amplitudes scaled to the strength of the initiating signal (16, 17). These phosphatase–substrate pairs thus provide an experimentally tractable system within a well-characterized model organism with which to identify principles for how PP2C family of phosphatases are regulated and achieve substrate specificity.

Using a combined forward-genetic and biochemical approach to probe the regulation and specificity of RsbU and SpoIIE, we found that the regulatory switch is a conserved element that ensures each phosphatase is only active at the correct time, under the correct conditions, and when in complex with the correct substrate protein.

**Results**

**SpoIIE principally achieves specificity by stabilizing the transition state**

To assess how SpoIIE achieves specificity for its cognate substrate SpoIIA as opposed to its noncognate substrate RsbV, we used the C-terminal phosphatase domain in isolation (SpoIIE590–827) (Fig. 1C). We selected this construct to test the hypothesis that specificity can be encoded by the PP2C catalytic domain alone and to identify potentially conserved mechanisms of specificity. At least three steps of the phosphatase reaction could in principle contribute to specificity: substrate recognition, chemistry, and metal-cofactor binding. To isolate the contributions of substrate recognition and chemistry, we held the metal-cofactor constant at saturating levels (10 mM MnCl2). Under these conditions, SpoIIE590–827 was specific, dephosphorylating SpoIIA-P approximately 3500-fold more efficiently than the off-pathway substrate, RsbV-P (SpoIIA-P \( k_{\text{cat}}/K_M = 0.68 \pm 0.04 \text{ min}^{-1} \mu\text{M}^{-1} \), RsbV-P \( k_{\text{cat}}/K_M = 0.00020 \pm 0.00002 \text{ min}^{-1} \mu\text{M}^{-1} \)) (Fig. 1D). This specificity was principally the result of an increased \( k_{\text{cat}} \) for the cognate substrate (18.7 ± 0.4 min\(^{-1}\) for SpoIIA, 0.064 ± 0.005 min\(^{-1}\) for RsbV) but was enhanced by a lower \( K_M \) for the cognate substrate (20 ± 1 \( \mu\text{M} \) for SpoIIA, 255 ± 39 \( \mu\text{M} \) for RsbV) (Fig. 1E). Thus, specificity is achieved by directing 3.4 kcal/mol of the specific binding energy for the cognate substrate to stabilize the transition state and 1.5 kcal/mol to stabilization of the enzyme substrate complex (Fig. 1F). Here we discuss \( K_M \) as representing “substrate recognition”, but also note that the lack of a lag phase in presteady state reactions, the \( \mu\text{M} \) \( K_M \) values for substrate, and minute timescale \( k_{\text{cat}} \) are all suggestive of the \( K_M \) being equal to the \( K_D \) of the enzyme–substrate complex.

**The regulatory switch couples substrate recognition to metal-cofactor binding**

The conserved regulatory switch moves to activate SpoIIE by recruiting a metal cofactor to the active site (Fig. 1C). We hypothesized that substrate binding is favored when the switch is in the active conformation, coupling substrate binding to metal-cofactor binding. To investigate this hypothesis, we took advantage of a gain-of-function mutant of SpoIIE (valine 697 to alanine, V697A) in which substitution of a single amino acid in the hydrophobic core of the phosphatase domain biases the switch to the active state (Fig. 1C) (11, 18, 19). The V697A substitution decreased the specificity of SpoIIE, increasing the \( k_{\text{cat}}/K_M \) for RsbV-P 150-fold compared with a 5-fold increase for SpoIIA-P (Fig. 2A). In the experiments described below, we analyzed the effect of the V697A substitution on \( K_M \), \( k_{\text{cat}} \), and metal-cofactor binding with both the cognate and noncognate substrates.

**Substrate recognition (\( K_M \))**

The V697A substitution decreased the \( K_M \) of SpoIIE590–827 for both SpoIIA-P (10-fold) and RsbV-P (5-fold) (Fig. 2A), suggesting that substrate binding is favored when SpoIIE is in the active conformation.

**Chemistry (\( k_{\text{cat}} \))**

The V697A substitution increased the maximal catalytic rate (\( k_{\text{cat}} \)) of SpoIIE590–827 for the noncognate substrate nearly 40-fold while having a modest effect on the cognate substrate (Fig. 2A). Thus, the switch controls the maximum catalytic rate of SpoIIE in addition to controlling metal-cofactor binding. Furthermore, the conformational flexibility of the switch can differentially affect the catalytic rate for cognate and noncognate substrates.

**Metal-cofactor binding**

To monitor metal-cofactor recruitment, we held the substrate concentration constant and measured SpoIIE activity with varied concentrations of MnCl2 (Fig. 2B). Consistent with our previous observations, the V697A substitution decreased the concentration of MnCl2 required for SpoIIE590–827 to dephosphorylate SpoIIA-P (\( K_{1/2} \) 3.1 ± 0.9 mM for WT SpoIIE590–827 and \( K_{1/2} \) 0.16 ± 0.05 mM for SpoIIE590–827 V697A). In contrast, the V697A substitution increases the MnCl2 concentration required for dephosphorylation for the noncognate substrate RsbV-P (\( K_{1/2} \) of 2.0 ± 0.4 mM for WT SpoIIE590–827 and \( K_{1/2} \) of 19 ± 2 mM for SpoIIE590–827 V697A). Cooperativity of metal-cofactor binding (Hill coefficient of two, inset panels Fig. 2B) yields a switch-like response profile, appropriate for initiation of a developmental transition.

Together, these kinetic parameters demonstrate that the switch mediates coupling between metal-cofactor binding, substrate recognition, and chemistry and that the V697A substitution shifts binding energy for the noncognate substrate from metal-cofactor binding to chemistry, reducing specificity at saturating metal-cofactor concentrations (summarized in Fig. 2C).

**The SpoIIE regulatory domain controls metal-cofactor binding, substrate recognition, and chemistry**

PP2C family of phosphatases has regulatory domains that control their activity (4, 11). We therefore repeated the above analysis with the phosphatase domain of SpoIIE together with a portion of the regulatory domain (SpoIIE457–827). We chose this fragment because it is a well-behaved soluble protein whose crystal structure we previously reported (11) (Fig. 3A). Although the regulatory domain is truncated, this fragment allowed us to test how contact with an accessory domain affects the behavior of the switch. Similar to our observations...
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Figure 2. SpoIIIE specificity is controlled by the switch. A, plots showing the dephosphorylation of SpoIIAA-P (red) or RsbV-P (blue). The bar plot (left) summarizes the impact of the V697A substitution on the $k_{cat}/K_M$ of SpoIIIE$^{590–827}$. Data from SpoIIIE$^{697–827}$ are displayed in light colors. B, reactions with SpoIIAA-P as the substrate were performed as multiple-turnover reactions with varying concentrations of SpoIIAA-P and 0.04 μM SpoIIIE$^{592–827}$ V697A. Data for wt SpoIIIE are the same experiment as shown in Figure 1D. All reactions contained 10 mM MnCl$_2$. To calculate $k_{cat}/K_M$, data from concentrations below the KM are fit to the linear equation observed rate = ($k_{cat}/K_M$) * [SpoIIAA-P or SpoIIIE]. Error bars are the error of the fit. Right is a plot of the data but including higher concentrations of SpoIIAA-P and SpoIIIE. Data are fit using nonlinear curve fitting (KaleidaGraph) to the equation observed rate = $k_{cat}$([SpoIIAA-P or SpoIIIE]/(KM + [SpoIIAA-P or SpoIIIE]). B, SpoIIIE specificity as a function of MnCl$_2$ concentration, with SpoIIAA-P (red) or RsbV-P (blue). Multiple turnover reactions with SpoIIAA-P as the substrate included 0.1 μM SpoIIIE and 50 μM SpoIIAA-P. Single turnover reactions with RsbV-P as the substrate included 50 μM SpoIIIE and trace RsbV-P. Data are fit to a cooperative model using the equation $k_{obs} = (k_{cat}/[MnCl_2]^{2}/(K_{1/2} + [MnCl_2])^{2})$. Hill plots from the data are shown to the right and are fit to a linear equation. For SpoIIIE$^{590–827}$ with SpoIIAA-P, h is 2.0 (1.7 for V697A) and 1.7 with RsbV-P (1.8 for V697A). C, a summary of kinetic parameters in the reaction scheme for dephosphorylation by SpoIIIE. $S$ indicates the phosphoprotein substrate, and $P$ indicates the product.

with SpoIIIE$^{590–827}$, SpoIIIE$^{457–827}$ was specific for SpoIIAA-P (SpoIIAA-P $k_{cat}/K_M = 0.039 \pm 0.002$ min$^{-1}$ μM$^{-1}$, RsbV-P $k_{cat}/K_M = 7.2 \pm 2 \times 10^{-5}$ min$^{-1}$ μM$^{-1}$) (Fig. 3A), and its specificity for its cognate substrate was reduced by the V697A substitution ($k_{cat}/K_M$ for RsbV-P increases 100-fold but was unaffected for SpoIIAA-P) (Fig. 3A).

SpoIIIE$^{457–827}$ exhibited several notable differences from the phosphatase domain alone (SpoIIIE$^{590–827}$). First, the regulatory domain increased the $K_M$ of SpoIIIE for both cognate and noncognate substrates, such that the concentration dependence of the phosphatase activity was linear even at concentrations of substrate above 100 μM (Fig. 3A). Second, the regulatory domain altered how the V697A substitution affected metal-cofactor binding; although the substitution decreased the concentration of MnCl$_2$ required for SpoIIIE$^{457–827}$ to dephosphorylate SpoIIAA-P (Fig. 2A), it had no effect on the concentration of MnCl$_2$ required to dephosphorylate RsbV-P (Fig. 3B).

Together, these findings (summarized in Fig. 3C) are consistent with coupling between substrate recognition, chemistry, and metal-cofactor recruitment and indicate that regulatory domains can influence all three steps through the switch.

A role for the switch-element in activating RsbU phosphatase activity

Next, we addressed whether coupled activation, substrate recognition, and metal-cofactor binding through the switch is conserved for the stress-response phosphatase, RsbU (Fig. 1A). First, we tested the hypothesis that the switch mediates RsbT activation of RsbU. We took an unbiased approach to interrogate the mechanism of RsbU activation by performing a genetic screen to isolate variants of RsbU that are active in the absence of RsbT. We reasoned that if the switch is important for RsbU regulation, we would isolate activating mutations in the switch region. We generated a library of randomly mutagenized rsbU variants and induced their expression in a strain harboring a LacZ reporter for σB activity and selected for variants that activated σB. From this screen, we identified two amino acid substitutions in the RsbU phosphatase domain, both at M166 (M166L and M166V), which activated σB in the absence of RsbT as visualized by lacZ-dependent color change on 5-bromo-4-chloro-3-indolyl-β-D-galactopyranoside (X-gal) plates (Fig. 4A shows plates with rsbU$^{M166L/V}$ introduced at the native locus; Fig. S2A shows the plates with strains expressing rsbU$^{M166L/V}$ using the plasmid-based system used in the screen). Activation of σB was dependent on expression of the mutagenized RsbU because both pathways known to dephosphorylate RsbV-P were deleted from the parental strain (and all other strains described below), and activation in plasmid-based systems depended on induction of RsbU expression (Fig. S2A). M166 is in the N-terminal α-helix of the switch and is predicted to pack in the hydrophobic core of the phosphatase domain (Fig. 4B). The position of these
mutations is reminiscent of the activating mutation V697A in SpoIIE (Fig. 4B), and the position of M166 in the RsbU switch provides independent evidence that the switch is required for RsbU activation.

Our analysis of the SpoIIE V697A mutation revealed that it acts by biasing the switch toward the active conformation, allowing for partial activation in the absence of signal, and potentiating the response to low levels of signal (11). Consistent with a similar mechanism for the M166 mutations, we observed that when the plasmids used to isolate the rsbU<sup>M166L</sup>/<sup>V</sup> variants were introduced to strains with rsbT<sup>B</sup>, σ<sup>B</sup> activity was elevated compared with ΔrsbT strains on plates (Fig. S2A). A similar phenomenon was not qualitatively apparent on plates (Fig. S2B, top panel). However, we observed that the σ<sup>B</sup> activity of liquid-grown cells expressing rsbU<sup>M166L</sup> sharply increased between absorbance 0.5 and 1.0 at 600 nm and peaked as the cells entered the stationary phase (absorbance 1.5 at 600 nm) (Fig. 4C, top panel). This cell density–dependent effect was potentiated by RsbT; σ<sup>B</sup> activity was 10-fold lower in strains that lacked rsbT (Fig. 4C, bottom panel). A similar cell density–dependent increase in σ<sup>B</sup> activity was not detected in cells expressing WT rsbU regardless of the presence of rsbT (Fig. 4C, top panel). These data suggest that mutation of M166 sensitizes RsbU to activation by RsbT—even at levels that are normally insufficient to elicit a response.

The rsbU<sup>M166L</sup> mutation also potentiated the RsbT-mediated response to environmental stress. Cells with rsbU<sup>M166L</sup> integrated at the native chromosomal locus had an elevated response to a well-characterized environmental stressor (4% ethanol by volume) that activates RsbU through RsbT (12). Stress was introduced at low cell density (absorbance 0.1–0.2 at 600 nm) to minimize the impact of cell density on σ<sup>B</sup> activity in rsbU<sup>M166L</sup>-expressing cells. The elevated response of cells expressing rsbU<sup>M166L</sup> mirrored the characteristic dynamics of the sharp and transient increase in σ<sup>B</sup> activity of WT cells followed by a gradual ramp-down of activities (Fig. 4D and Fig. S2B, top panel). As this response depends on both RsbT and RsbU, strains lacking either of these genes failed to respond to the stressor (Fig. 4D, bottom panel). Together, the position of M166 in the RsbU regulatory switch and its impact on RsbU activity implicate the switch as a conserved element to control phosphatase activity in response to cellular signals.

**Mechanisms of regulation and specificity are conserved between SpoIIE and RsbU**

If the switch element acts similarly to activate SpoIIE and RsbU, a simple prediction is that activation of RsbU by RsbT would decrease the concentration of the metal cofactor required for the activity. We reconstituted RsbU activation by RsbT and
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Figure 4. Mutation of the switch activates RsbU activity in the absence of RsbT. A, RsbU mutants that are active in the absence of RsbT from a blue-white screen using a strain carrying a lacZ-reporter for α⁺ activity. Colonies of cells expressing rsbUM166L or rsbUM166V at the native rsbU locus on the chromosome exhibit high levels of α⁺ activity both in the presence and absence of rsbT. All strains lack rsbPO and carry a reporter for α⁺ activity (amyE:ctc-lacZ) displayed as green spheres. The α-helical switch is represented in orange and the side chains that coordinate divalent cations in the active site are shown as sticks. C, rsbUM166L leads to increased α⁺ activity as cultures approach the stationary phase. Plots show beta-galactosidase activity from rsbPO deleted strains carrying a lacZ-reporter for α⁺ activity (ΔrsbPO amyE:ctc-lacZ) either encoding WT rsbU (blue), rsbUM166L (red) (top panel), or deleted for rsbU (gray) (bottom panel). Light colors indicate that the strain additionally had rsbT deleted (bottom panel). The plots show cell density–dependent activation of α⁺ from four independent replicates per strain background. Cultures were sampled from the early log phase to the stationary phase, and beta-galactosidase activity is plotted as a function of time of ethanol addition. Error bars reflect SD from the mean. Y-axis scales differ between top and bottom panels to allow better visualization of data. B, Strains were grown on LB plates containing 80 μg/ml X-gal at 37 °C and imaged after 18 h. The presence of a blue pigment indicates α⁺ activity and activation of the stress response pathway. B, a structure of the SpoIIIE PP2C domain (PDB ID: 5UCG), with the co of V697 and S640 (the equivalent of M166 in RsbU) displayed as green spheres. The α-helical switch is represented in orange and the side chains that coordinate divalent cations in the active site are shown as sticks. C, rsbUM166L leads to increased α⁺ activity as cultures approach the stationary phase. Plots show beta-galactosidase activity from rsbPO deleted strains carrying a lacZ-reporter for α⁺ activity (ΔrsbPO amyE:ctc-lacZ) either encoding WT rsbU (blue), rsbUM166L (red) (top panel), or deleted for rsbU (gray) (bottom panel). Light colors indicate that the strain additionally had rsbT deleted (bottom panel). The plots show cell density–dependent activation of α⁺ from four independent replicates per strain background. Cultures were sampled from the early log phase to the stationary phase, and beta-galactosidase activity is plotted as a function of time of ethanol addition. Error bars reflect SD from the mean. Y-axis scales differ between top and bottom panels to allow better visualization of data.

found that RsbT decreased the K_{1/2} of RsbU for Mn^{2+} by nearly 100-fold (K_{1/2} 1.0 ± 0.1 mM in the presence of RsbT and 77 ± 9 mM in the absence of RsbT) (Fig. 5A). In contrast to SpoIIIE, RsbU activity toward its cognate substrate was noncooperative with respect to Mn^{2+} concentration (Fig. 5A, right panels). Although SpoIIIE regulates a developmental transition, for which an all-or-nothing response is important, RsbU regulates a stress response, for which the graded response that noncooperative activation affords may be advantageous.

In our in vitro assays, RsbT additionally enhanced the maximal catalytic activity of RsbU toward RsbV-P 10-fold (k_{cat} 25 ± 2 min⁻¹ in the presence of RsbT and 0.44 ± 0.06 min⁻¹ in the absence of RsbT) (Fig. 5A), consistent with previous observations (12, 20). This suggests that, as we observed for SpoIIIE, control of chemistry and metal-cofactor binding are coupled, providing nearly a 1000-fold increase in the activity of RsbU at concentrations of metal ion below the K_{1/2} (as would be expected in the cytoplasm) (21, 22).

Reconstitution of RsbU activation by RsbT further allowed us to determine how specificity is maintained upon phosphatase activation. Like SpoIIIE, RsbU is specific: RsbU had a higher catalytic activity toward its cognate substrate (k_{cat} 0.44 ± 0.06 min⁻¹ for RsbV-P and 0.0012 ± 0.0004 min⁻¹ for SpoIIAA-P) and bound both cognate and noncognate substrates tightly (K_{M} 0.9 μM for SpoIIAA-P and 1.5 μM for RsbV-P; K_{M} values were difficult to determine accurately because of limitations of detection for substrate proteins) (Fig. 5B).

As with the cognate substrate, RsbT enhanced RsbU catalytic activity toward SpoIIAA-P (a 300-fold increase to k_{cat} 0.32 ± 0.02 min⁻¹) (Fig. 5B). However, RsbT did not substantially change the affinity of metal-cofactor binding with the noncognate substrate (K_{M} 8 ± 3 mM in the absence of RsbT and 20 ± 6 mM in the presence of RsbT), similar to the V697A mutation of SpoIIIE (457–457) (Fig. 5A). We did note partial cooperativity for manganese with SpoIIAA as the substrate.
(Hill coefficient of 1.4 ± 0.1 without RsbT and 1.6 ± 0.3 with RsbT). This mild effect is consistent with our observation that the substrate influences the binding of metal in the active site. RsbT also had little impact on SpoIIAA-P binding (approximately K_{diss} 1 μM in the absence of RsbT and 0.3 μM in the presence of RsbT) (Fig. 5B). Thus, specificity is maintained because RsbT activation fails to stimulate metal-cofactor binding for RsbU in the presence of its noncognate substrate (summarized in Fig. 5C).

Conclusions

By examining two phosphatases, we found that PP2C regulation and specificity is mediated by a conserved allosteric element that is functionally flexible to respond to varied signals and to discriminate between substrate proteins. This regulatory mechanism is orchestrated by an allosteric switch that integrates activating signals and binding to cognate substrate to control metal-cofactor binding in the active site and catalytic activity. This coupling of regulatory inputs, chemistry, and substrate recognition provides a unified mechanism to control diverse signaling pathways.

By what mechanism could the PP2C switch element transduce regulatory inputs to coordinate substrate recognition, metal-cofactor binding, and chemistry? In a parsimonious model, both regulatory domains and substrates might directly contact and influence the conformation of the switch. This is supported by our findings that a fragment of the regulatory domain of SpoIIIE shifts the energetic balance between substrate recognition, chemistry, and metal-cofactor recruitment. Furthermore, the switch is the structural feature of SpoIIIE that mediates these changes in phosphatase activity and substrate recognition. The switch is situated directly below the active site, where it would likely contact substrate proteins and might also control the maximum catalytic rate by precise positioning of the phosphoserine bond and the metal cofactors in the active site. Such an architecture is present in the single available structure of a phosphatase from the PP2C family bound to its substrate protein (the phosphatase Hab1 bound to SnRK2). SnRK2 makes direct contacts with the Hab1 switch and to a variable region (termed “the flap”) that packs against the switch (23).

Dimerization may also be a widespread regulatory mechanism for the PP2C family of phosphatases. SpoIIIE is activated by formation of a dimer in which the switch elements of the two phosphatase domains make direct contacts and lock the switch element in the active state (11). One attractive possibility is that the specific contacts between the switch elements in the phosphatase dimer are conserved. This is supported by the fact that similar contacts have been observed in dimeric structures of *Pseudomonas aeruginosa* phosphatases RssB and SiaA (24, 25). Dimerization also plays an important role in RsbU activation; a fragment of the RsbU regulatory domain crystallized as a dimer and a groove formed at the dimer interface was shown to be critical for RsbT binding to RsbU (20). However, the mechanism of how RsbU dimerization is coupled to phosphatase activation is unknown.

We hypothesize that this conserved allosteric switch may regulate diverse PP2C family of phosphatases across the tree of life. Regulatory domains from diverse PP2C family of phosphatases also pack against the switch, suggesting that coupling between substrate recognition, metal-cofactor binding, and activity could be broadly conserved (11, 26–28). In addition,
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structural analysis and molecular dynamics simulations of human PP2Ca revealed a network of interactions that link the switch element, metal binding in the active site, and the flap subdomain (29). Hydrogen–deuterium exchange experiments revealed increased mobility of the switch α1 and α2 at subsaturating metal-cofactor concentrations and when the binding site for a third metal ion was disrupted by mutation of its coordinating residue, D146 (29). Further highlighting the similarities between the bacterial and human phosphatases, SpoIIE V697 and PP2Ca D146 both sit at the end of β-strand 6, two amino acids away from each other in a sequence alignment. These data suggest that the structural features of the PP2C catalytic domain that we identify as controlling phosphatase activity and specificity are broadly conserved.

The mechanism we describe here for PP2C regulation and specificity, however, is qualitatively different from what is known about evolutionarily unrelated serine/threonine phosphatases from other families. For members of the phosphoprotein phosphatase family including PP1, PP2A, and calcineurin, the catalytic domain itself is thought to be relatively promiscuous for substrates. Substrate specificity is instead imposed by diverse regulatory subunits that tether substrates in proximity of the catalytic center, control subcellular localization of the phosphatase, and/or gate substrate access to the active site (4).

Because phosphatases control signaling pathways that contribute to multiple disease processes, they are attractive drug targets, but their lack of deep and thus classically druggable active site pockets has made development of small-molecule modulators difficult (30). Recent successes, however, have targeted phosphatase regulatory mechanisms: PP1 inhibitors that target the conformation of regulatory subunits (31), PP2A activators that promote the assembly of specific regulatory subunits into holoenzymes (32), and allosteric modulators of active site gating for the protein tyrosine phosphatase, SHP2 (33). The allosteric regulatory mechanism of PP2C family of phosphatases may also similarly provide strategies for the development of modulators. Indeed, allosteric inhibitors of the oncogenic PP2C Wip1 were identified that bind to the flap subregion, although the mechanism of action of these inhibitors is unknown (34, 35). We propose that these and future PP2C modulators may act on the allosteric switch we here describe as coordinating substrate recognition, chemistry, and metal-cofactor binding.

Experimental procedures

Growth conditions

Liquid cultures were grown in Lennox lysogeny broth (LB), whereas colonies were grown on LB containing Bacto agar. The antibiotics used were macrolide-lincosamide-streptogramin (0.5 μg/ml erythromycin, 2.5 μg/ml lincomycin), chloramphenicol (20 μg/ml for Escherichia coli), carbenicillin (100 μg/ml), and kanamycin (10 μg/ml for B. subtilis or 50 μg/ml for E. coli). When indicated, 80 μg/ml of X-gal and 1 mM IPTG were added to the growth medium to visualize σB reporter activity on plates. B. subtilis strains are listed in Supplemental Table S1, E. coli strains are listed in Supplemental Table S2, and primers are listed in Supplemental Table S3.

Protein-expression constructs

SpoIIAA and SpoIIAB were expressed in BL21 (DE3) Rosetta 2 pLysS cells as 6H-sumo fusions in pET23a made by ligation of the coding sequence in pET23a 6H-sumo digested with NotI/Agel. SpoIIAA–P was produced in BL21 (DE3) cells containing pET-YSBLIC 6H-3C::spoIIAA–spoIIAB as reported (36). SpoIIIE V697–827, SpoIIIE 827, RsbT, RsbU, RsbV, and RsbW were produced with 3C cleavable 6H tags in BL21 (DE3) cells (constructs were made by insertion of the coding sequence to pET47b vectors digested with Xmal/Xhol by isothermal assembly). Mutations were introduced to SpoIIIE expression constructs using the QuikChange site-directed mutagenesis kit. RsbV-P was produced by coexpressing RsbV and RsbW using a pET47b vector generated by isolation of the coding sequences for RsbV and RsbW from genomic DNA, yielding a construct with RsbV fused to a 3C cleavable 6H tag followed by untagged RsbW. All strains used are listed in the tables of strains (Supplemental Tables S1 and S2).

Protein expression and purification

Proteins were expressed and purified using slight modifications to previous methods (11). All proteins were purified to greater than 95% purity as assessed by Coomassie Brilliant Blue–stained SDS-PAGE gel. Unless otherwise noted, protein expression was induced with 1 mM IPTG for 14 to 18 h at 14 °C in BL21 (DE3) cells that had been grown in LB to absorbance 0.6 at 60 at room temperature (RT). Details for purification of individual proteins are as follows:

SpoIIE (the same protocol for all variants) was purified from cell pellets lysed (using a microfluidizer) in 50 mM K•Hepes, pH 8.0, 200 mM NaCl, 10% glycerol (v/v), 0.5 mM DTT, 20 mM imidazole, and 0.1 mg/ml PMSF. Lysates were clarified by spinning for 30 min at 16,000 RPM in a Sorvall SS-34 rotor at 4 °C. Protein was bound to a HisTrap HP column on an AKTA FPLC, washed, and then eluted with a gradient to 200 mM imidazole. The 6H tag was cleaved overnight in dialysis with PreScission Protease, and the tag and protease were removed by flowing over Ni-NTA resin. SpoIIE was then purified on a Resource Q column equilibrated in 50 mM Hepes, pH 8.0, 100 mM NaCl, 2 mM EDTA, and 2 mM DTT and eluted with a gradient to 500 mM NaCl. The SpoIIE containing fractions were spin concentrated and gel-filtered on a 120 ml Superdex 75 column equilibrated in 20 mM K•Hepes, pH 8.0, 100 mM NaCl, 10% glycerol (v/v), and 2 mM DTT. Protein was spin-concentrated to approximately 400 μM and flash-frozen and stored at −80 °C.

SpoIIAA was expressed for 4 h at 37 °C. Cell pellets were resuspended in 20 mM K•Hepes, pH 7.5, 200 mM NaCl, 0.5 mM DTT, and 0.1 mg/ml PMSF (5 ml/l of culture). Cells were lysed using a cell disruptor in one-shot mode (Constant Systems), and lysates were clarified by spinning for 30 min at
16,000 RPM in a Sorvall SS-34 rotor at 4 °C. The protein was bound to Ni-NTA resin (1 ml/l of culture) on the column by gravity flow, washed with the buffer containing 20 mM imidazole, and eluted with 200 mM imidazole. The 6H-sumo tag was cleaved using 10 μl of a 100 μM stock of ULP1 in overnight dialysis at 16 °C to 20 mM K•Hepes, pH 7.5, 200 mM NaCl, 10% glycerol (v/v), and 0.5 mM DTT. The 6H-sumo tag was subtracted by elution over Ni-NTA resin equilibrated in the cleavage and dialysis buffer. The cleaved protein was then spin-concentrated and gel-filtered on a 120 ml Superdex 75 column equilibrated in 50 mM K•Hepes, pH 7.5, 175 mM NaCl, 10 mM MgCl2, 10% glycerol (v/v), and 2 mM DTT. The protein was concentrated to approximately 400 μM, supplemented to 50% glycerol (v/v), and flash-frozen and stored at –80 °C.

SpoIIAB was expressed for 4 h at 37 °C. Cell pellets were resuspended in 50 mM K•Hepes, pH 7.5, 200 mM NaCl, 10 mM MgCl2, 0.5% Triton X-100 (v/v), 0.5 mM DTT, and 0.1 mg/ml PMSF. Cells were lysed using a cell disruptor in the one-shot mode (Constant Systems), and lysates were clarified by spinning for 30 min at 16,000 RPM in a Sorvall SS-34 rotor at 4 °C. The protein was bound to Ni-NTA resin (1 ml/l of culture) on the column by gravity flow, washed with the buffer containing 20 mM imidazole, and eluted with 200 mM imidazole. The 6H-sumo tag was left uncleaved to aid in removal from phosphorylation reactions. SpoIIAB was spin-concentrated and gel-filtered on a 120 ml Superdex 75 column equilibrated in 50 mM K•Hepes, pH 7.5, 175 mM NaCl, 10 mM MgCl2, 10% glycerol (v/v), and 2 mM DTT. The protein was concentrated to approximately 400 μM, supplemented to 50% glycerol (v/v), and flash-frozen and stored at –80 °C.

RsbT was purified from cell pellets lysed (using a microfluidizer) in 20 mM K•Hepes, pH 7.5, 200 mM NaCl, 10% glycerol (v/v), 0.5 mM DTT, 20 mM imidazole, and 0.1 mg/ml PMSF. Lysates were clarified by spinning for 30 min at 16,000 RPM in a Sorvall SS-34 rotor at 4 °C. The protein was bound to Ni-NTA resin (1 ml/l of culture) on the column by gravity flow, washed, and eluted with 200 mM imidazole. The 6H tag was cleaved overnight in dialysis with PreScission protease, and the tag and protease were removed by flowing over Ni-NTA resin. RsbT was spin-concentrated and gel-filtered on a 120 ml Superdex 75 column equilibrated in 20 mM K•Hepes, pH 7.5, 150 mM NaCl, 10% glycerol (v/v), and 2 mM DTT. The protein was spin-concentrated to approximately 40 μM (additional concentration led to aggregation) and was flash-frozen and stored at –80 °C.

RsbU was purified from cell pellets lysed (using a microfluidizer) in 50 mM K•Hepes, pH 8.0, 200 mM NaCl, 10% glycerol (v/v), 0.5 mM DTT, 20 mM imidazole, and 0.1 mg/ml PMSF. Lysates were clarified by spinning for 30 min at 16,000 RPM in a Sorvall SS-34 rotor at 4 °C. The protein was bound to a HisTrap HP column on an AKTA FPLC, was washed with the buffer containing 20 mM imidazole, and eluted with 200 mM imidazole. The 6H tag was cleaved overnight in dialysis with PreScission protease, and the tag and protease were removed by flowing over Ni-NTA resin. Cleaved RsbU was spin-concentrated and gel-filtered on a 120 ml Superdex 75 column equilibrated in 20 mM K•Hepes, pH 8.0, 100 mM NaCl, 10% glycerol (v/v), and 2 mM DTT. The RsbU-containing fractions were then purified on a RESOURCE Q column equilibrated in 50 mM Hepes, pH 8.0, 100 mM NaCl, 2 mM EDTA, and 2 mM DTT and eluted with a gradient to 500 mM NaCl. This final step removes a degradation product present in some RsbU preparations. The protein was supplemented with 10% glycerol (v/v), spin-concentrated to approximately 400 μM, and flash-frozen and stored at –80 °C.

RsbV was purified from cell pellets lysed (using a microfluidizer) in 20 mM K•Hepes, pH 7.5, 200 mM NaCl, 10% glycerol (v/v), 0.5 mM DTT, 20 mM imidazole, and 0.1 mg/ml PMSF. Lysates were clarified by spinning for 30 min at 16,000 RPM in a Sorvall SS-34 rotor at 4 °C. The protein was bound to Ni-NTA resin (1 ml/l of culture) on the column by gravity flow, washed, and eluted with 200 mM imidazole. The 6H tag was cleaved overnight in dialysis with PreScission protease, and the tag and protease were removed by flowing over Ni-NTA resin. RsbV was spin-concentrated and gel-filtered on a 120 ml Superdex 75 column equilibrated in 20 mM K•Hepes, pH 7.5, 150 mM NaCl, 10% glycerol (v/v), and 2 mM DTT. The protein was spin-concentrated to approximately 400 μM and flash-frozen and stored at –80 °C.
Control of phosphatase activity and specificity

RsbW was purified from cell pellets lysed (using a microfluidizer) in 20 mM K•Hepes, pH 7.5, 10 mM MgCl₂, 200 mM NaCl, 10% glycerol (v/v), 0.5 mM DTT, 20 mM imidazole, and 0.1 mg/ml PMSF. Lysates were clarified by spinning for 30 min at 16,000 RPM in a Sorvall SS-34 rotor at 4 °C. The protein was bound to a HisTrap HP column on an AKTA FPLC, washed, and eluted with a gradient to 250 mM imidazole. The protein was spin-concentrated to approximately 100 μM and flash-frozen and stored at −80 °C.

RsbV-P was purified from cells coexpressing 6H-3C-RsbV and untagged RsbW that were grown at 37 °C and induced with 1 mM IPTG for 4 h at 37 °C. Cells were lysed (using a microfluidizer) in 50 mM K•Hepes, pH 7.5, 50 mM KCl, 10% glycerol (v/v), 0.5 mM DTT, 20 mM imidazole, and 0.1 mg/ml PMSF. Lysates were clarified by spinning for 30 min at 16,000 RPM in a Sorvall SS-34 rotor at 4 °C. The protein was bound to a HisTrap HP column on an AKTA FPLC, washed, and eluted with a gradient to 250 mM imidazole. RsbVW-containing fractions were then gel-filtered using a 120 ml Superdex 75 column equilibrated in 50 mM K•Hepes, pH 7.5, 150 mM NaCl, 10% glycerol (v/v), and 2 mM DTT. The protein was spin-concentrated to approximately 100 μM and flash-frozen and stored at −80 °C.

Phosphatase assays

Phosphatase assays were performed using methods reported previously (9) with modifications as described. To produce 32P-labeled SpoIIAA-P, 75 μM SpoIIAA, 5 μM SpoIAB, and 50 μCi of γ-32P ATP were incubated overnight at RT in 50 mM K•Hepes, pH 7.5, 50 mM KCl, 750 μM MgCl₂, and 2 mM DTT. Unincorporated nucleotide was removed from the reaction mixture by buffer exchange using a Zeba spin column (Pierce) equilibrated in 50 mM K•Hepes, pH 8.0, 100 mM NaCl, 20 mM imidazole. 6H-tagged RsbW was then washed, and eluted with a gradient to 500 mM NaCl to remove all unincorporated nucleotide and free phosphate. Labeled RsbV-P was aliquoted and frozen at −80 °C for future use.

All phosphatase assays were performed at RT in 25 mM K•Hepes, pH 8, 100 mM NaCl, and 100 μg/ml BSA (to prevent protein sticking to tubes). The concentrations of enzyme, substrate, and MnCl₂ were varied as indicated. 10 μM RsbT was additionally added to reactions as indicated. We note that the magnitude of effects reported for RsbT are lower limits because higher concentrations of RsbT could not be tested because of aggregation of RsbT at a high concentration. Reactions were stopped in 0.5 M EDTA, pH 8.0, and 2% Triton X-100 (v/v) and run on PEI-Cellulose TLC plates developed in 1 M LiCl₂ and 0.8 M acetic acid, and imaged on a Typhoon (GE Life Sciences). Phosphatase assays were performed more than three independent times as separate experiments.

Strain construction

B. subtilis strains were constructed using standard molecular genetic techniques (in the PY79 strain background) and were validated to contain the correct constructs by double-crossover recombination at the correct insertion site. All strains are used in this study are described in the table of strains. A ΔrsbPQ ΔrsbTU rsbV-FLAG amyE::ctc-lacZ reporter strain (KC479 (14)) was used as a parent strain for all B. subtilis strains. A parent strain (ΔrsbPQ ΔrsbTU rsbV-FLAG amyE::ctc-lacZ) was generated for the screen of rsbT-independent rsbU variants. To generate a marked deletion of rsbPQ, the genomic regions 500 bp upstream and downstream of rsbPQ were amplified and ligated to a kanamycin resistance cassette using standard SOE PCR techniques. The linear DNA fragment was inserted into KC479 using standard techniques. Transformants were selected based on kanamycin resistance and confirmed by colony PCR and sequencing. A clean deletion of rsbTU was generated by generating a gBlock (IDT DNA Technologies) of regions corresponding to 500 bp upstream of rsbTU and 500 bp downstream of rsbU ligated together. This fragment was introduced into the HindIII and EcoRI sites of pminiMAD2 (37). Standard genetic techniques were used to excise the chromosomal copy of rsbTU. Mutants were confirmed by colony PCR and sequencing of the genomic region. Similarly, clean deletions of sigB, rsbT, and rsbU were generated, with the modification that we utilized a gBlock (IDT DNA Technologies) of regions corresponding to 500 bp upstream and 500 bp downstream of the appropriate genes. A single FLAG tag was introduced at the C-terminus of rsbV using the pminiMAD2 vector. Point mutations of the rsbU gene were similarly
introduced using the pminiMAD2 vector and confirmed by colony PCR and sequencing.

An inducible expression vector (pKH001-Pspank-rsbU) was generated to screen for rsbT-independent variants of rsbU. A gBlock (IDT DNA Technologies) was generated containing a super-folding gfpl gene flanked by AgeI and NotI sites and introduced into plasmid pDR110 (at the HindIII and SphI sites) using standard genetic techniques. The fragment of the resulting plasmid containing the lacI gene and the inducible sfgfp gene was PCR-amplified and inserted into vector phB201 (38) at the SacI and KpnI sites using standard genetic techniques to generate pKH001-Pspank-sfgfp. To generate expression constructs for rsbU and rsbTU, the appropriate genes were PCR-amplified from genomic DNA from PY79 using a high-fidelity polymerase and introduced at the AgeI and NotI sites using isothermal assembly.

To generate libraries of pKH001-Pspank-rsbU containing mutagenized rsbU, the rsbU gene was PCR-amplified from genomic DNA from PY79 using error-prone PCR, utilizing the natural error rate of Taq polymerase. Mutagenized rsbU was introduced at the AgeI and NotI sites using standard genetic techniques. Plasmids were introduced into the parent strain (ΔrsbPQ ΔrsbTU rsbV-FLAG amyE::ctc-lacZ) using standard B. subtilis transformation techniques.

Screen

Six independent pools of mutagenized rsbU were generated by amplifying rsbU under mutagenic PCR conditions using GoTaq polymerase (Promega) and an additional 2 mM MgCl2. These pools of mutagenized rsbU were subcloned into linearized plasmid using isothermal assembly to generate pKH001-P (spank)-rsbU and transformed separately into E. coli DH5α cells to propagate plasmids. The mutation rate was confirmed to be approximately one SNP per kilobase of DNA by sequencing selected clones. E. coli colonies were pooled to generate mixed populations of mutagenized pKH001-Pspank-rsbU, and this DNA was introduced separately into independent clones of the parent strain. Cultures were grown in selective media and plated on the selective medium containing IPTG and X-gal for 2 days at 37 °C.

Cells were lyzed in the Z-buffer containing a final concentration of 10 mg/ml lysozyme for 30 min at 37 °C. β-D-galactoside was dissolved in Z-buffer (60 mM Na2HPO4 7•H2O, 40 mM NaH2PO4 H2O, 10 mM KCl, 1 mM MgSO4, 50 mM 2-mercaptoethanol) and kept on ice. Cells were arrayed in a 96-well plate, and the absorbance at 600 nm was measured to assess cell culture density. Cells were lyzed in the Z-buffer containing a final concentration of 10 mg/ml lysozyme for 30 min at 37 °C. β-D-galactoside was dissolved in Z-buffer (4 mg/ml) and added to lysed cells to a final concentration of 0.67 mg/ml. Absorbance at 420 nm (to measure β-o-nitrophenyl-β-D-galactoside cleavage) and 550 nm (to control for light scattering caused by cell debris) was measured over 40 min, and the rate of LacZ production was calculated from the slope of the linear phase of A420 − 1.75*A550 (between 300 and 1500 s). Miller units were calculated using the formula that 1 Miller Unit = 1000*(A420 − (1.75*A550))/(t*v*absorbance at 600 nm); t is time in minutes, and v is the volume of the reaction in milliliters.

Data availability

All data are contained within the article or available to be shared upon request (please contact the corresponding author).

Control of phosphatase activity and specificity

Liquid-based beta-galactosidase assays

Beta-galactosidase assays to measure β reporter activity were adapted from previously described protocols (39). In brief, cells were grown to desired absorbance, spun down, and frozen at ~80 °C. Thawed pellets were resuspended in Z-buffer (60 mM Na2HPO4 7•H2O, 40 mM NaH2PO4 H2O, 10 mM KCl, 1 mM MgSO4, 50 mM 2-mercaptoethanol) and kept on ice. Cells were arrayed in a 96-well plate, and the absorbance at 600 nm was measured to assess cell culture density. Cells were lyzed in the Z-buffer containing a final concentration of 10 mg/ml lysozyme for 30 min at 37 °C. β-D-galactoside was dissolved in Z-buffer (4 mg/ml) and added to lysed cells to a final concentration of 0.67 mg/ml. Absorbance at 420 nm (to measure β-o-nitrophenyl-β-D-galactoside cleavage) and 550 nm (to control for light scattering caused by cell debris) was measured over 40 min, and the rate of LacZ production was calculated from the slope of the linear phase of A420 − 1.75*A550 (between 300 and 1500 s). Miller units were calculated using the formula that 1 Miller Unit = 1000*(A420 − (1.75*A550))/(t*v*absorbance at 600 nm); t is time in minutes, and v is the volume of the reaction in milliliters.

Figure preparation

Figures from structural models were generated using PyMol (Version 2.0) and UCSF Chimera (40). The sequence alignment in Fig. S1 was constructed and validated using UCSF Chimera and rendered using Geneious R7.

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Abbreviations—The abbreviations used are: LB, lysogeny broth; PP2C, protein phosphatase 2C; X-gal, 5-bromo-4-chloro-3-indolyl-β-D-galactopyranoside.

References

Control of phosphatase activity and specificity


