Bacterial tyrosine kinases (BY-kinases) comprise a family of protein tyrosine kinases that are structurally distinct from their functional counterparts in eukaryotes and are highly conserved across the bacterial kingdom. BY-kinases act in concert with their counteracting phosphatases to regulate a variety of cellular processes, most notably the synthesis and export of polysaccharides involved in biofilm and capsule biogenesis. Biochemical data suggest that BY-kinase function involves the cyclic assembly and disassembly of oligomeric states coupled to the overall phosphorylation levels of a C-terminal tyrosine cluster. This process is driven by the opposing effects of intermolecular autophosphorylation, and dephosphorylation catalyzed by tyrosine phosphatases. In the absence of structural insight into the interactions between a BY-kinase and its phosphatase partner in atomic detail, the precise mechanism of this regulatory process has remained poorly defined. To address this gap in knowledge, we have determined the structure of the transiently assembled complex between the catalytic core of the Escherichia coli (K-12) BY-kinase Wzc and its counteracting low–molecular weight tyrosine protein phosphatase (LMW-PTP) Wzb using solution NMR techniques. Unambiguous distance restraints from paramagnetic relaxation effects were supplemented with ambiguous interaction restraints from static spectral perturbations and transient chemical shift changes inferred from relaxation dispersion measurements and used in a computational docking protocol for structure determination. This structure presents an atomic picture of the mode of interaction between an LMW-PTP and its BY-kinase substrate, and provides mechanistic insight into the phosphorylation-coupled assembly/disassembly process proposed to drive BY-kinase function.

**Significance**

This study provides a structural view of the recognition of a bacterial tyrosine kinase (BY-kinase) by its counteracting phosphatase. BY-kinase function has been proposed to entail the cyclic assembly/disassembly of an oligomer coupled to the phosphorylation level of a C-terminal tyrosine cluster (Y-cluster). The structure of the BY-kinase–phosphatase complex highlights the use of the same conserved element on the kinase for both oligomerization and phosphatase docking. This mode of interaction prevents phosphatase engagement and dephosphorylation before exceeding a critical threshold of Y-cluster phosphorylation necessary for oligomer disassembly. Reassembly occurs once the Y-cluster is sufficiently dephosphorylated by the phosphatase that is then excluded, thereby repriming the cycle. This phosphorylation-coupled assembly/disassembly process likely serves to temporally coordinate intra- and extracytoplasmic events.
importance of maintaining the appropriate balance between self-interactions, to enable in trans autophosphorylation in BY-kinases, and those with a PTP, to facilitate dephosphorylation. While the atomic details of the self-interactions in BY-kinases have been highlighted by several structures (8, 12, 13), no such structural information is available for the interactions of the BY-kinases with their counteracting PTPs.

In prior studies, we had utilized NMR spectral perturbations to analyze the interactions between the CD of the \textit{Escherichia coli} (K-12) BY-kinase, Wzc (Wzc CD), and its cognate low–molecular weight PTP (LMW-PTP), Wzb (SI Appendix, Fig. S1B) (24). These studies suggested a conserved motif, 508Glu-[Xxx]2-Arg-[Xxx]2-Arg514, on the WzcCD α2-helix (SI Appendix, Fig. S1A), necessary for oligomerization (12), to also be involved in binding Wzb (24). Since those studies relied solely on NMR spectral perturbations, they did not resolve atomic details of the interactions that define the WzcCD–Wzb interface. Here, we utilize solution NMR methods to derive a variety of structural restraints to generate an atomic model of the Wzb–WzcCD complex that is validated through functional studies in vitro. This structure represents a detailed atomic view of the interaction between a BY-kinase and its cognate PTP, and indeed of any eukaryotic or prokaryotic LMW-PTP with its kinase partner, and provides critical insight into the role of this interaction in driving BY-kinase function.

**Results**

We have previously shown that the disordered C-terminal tail of WzcCD, that includes the Y-cluster (residues Tyr705 to Lys720), is dispensable for its interaction with Wzb (24). A nonhydrolyzable phosphorysine mimic was also found to make minimal contributions to the binding free energy (24).

**Table 1. Apparent binding affinities of various Wzc\textsubscript{CD\_ΔC} constructs for wild-type Wzb**

| Construct                          | \(K_D\), μM |
|------------------------------------|-------------|
| Wzc\textsubscript{CD\_ΔC}\*       | 8.6 ± 0.6   |
| Wzc\textsubscript{CD\_ΔC,Cys}     | 15.2 ± 0.6  |
| Wzc\textsubscript{CD\_ΔC,Cys,Val466Cys} | 8.9 ± 0.3 |
| Wzc\textsubscript{CD\_ΔC,Cys,Leu5OS5Cys} | 43.8 ± 1.3 |
| Wzc\textsubscript{CD\_ΔC,Cys,Ser516Cys} | 11.3 ± 0.7 |
| Wzc\textsubscript{CD\_ΔC,Cys,Gln523Cys} | 16.9 ± 0.5 |
| Wzc\textsubscript{CD\_ΔC,Cys,Lys556Cys} | 7.3 ± 0.4 |

Determined through two-dimensional line-shape analyses of \(^13\text{C},^1\text{H}\) HMQC spectra assuming that the fast-relaxing component can be neglected.

* Determined by ITC, 10.7 ± 1.3 μM, or SPR, 5.1 ± 0.2 μM.
Therefore, we utilized a construct, WzcCDAC, encoding the catalytic core of Wzc (Ser447 to Ala704) for our structural studies. We relied on two-dimensional line-shape analyses of resonances (25) in methyl \(^{13}\)C,\(^{1}\)H heteronuclear multiple quantum coherence (HMQC) spectra of Wzb, labeled at the isoleucine (B1), leucine and valine methyl positions (ILV-labeled), in the presence of increasing amounts of WzcCDAC (and its variants; described below) to quantify their mutual affinities. In the case of wild-type WzcCDAC, this approach yields a \(K_D\) of 8.6 ± 0.6 \(\mu\)M (Table 1), comparable to that obtained from isothermal titration calorimetry (ITC) (10.7 ± 1.3 \(\mu\)M; SI Appendix, Fig. S2) and that previously obtained using surface plasmon resonance (SPR) measurements (5.1 ± 0.2 \(\mu\)M) (24).

We had previously noted extensive line broadening in \(^{15}\)N,\(^{1}\)H transverse relaxation optimized spectroscopy (TROSY) spectra of Wzb in the presence of an equimolar amount of WzcCDAC (24), suggesting substantial exchange contributions attributable to the binding/unbinding process. Indeed, these effects prevented the use of spectral perturbations to precisely identify the WzcCDAC-binding residues on Wzb. To identify Wzb “seed” residues that facilitate WzcCDAC recognition, we analyzed the transient chemical shift changes induced on Wzb by the presence of substoichiometric amounts of WzcCDAC that manifest as exchange contributions to transverse relaxation (\(R_2\)) in Carr-Purcell-Meiboom-Gill (CPMG) based \(^{15}\)N relaxation dispersion measurements (26). While Wzb does not exhibit significant exchange in the absence of WzcCDAC (SI Appendix, Fig. S3A), specific regions display exchange effects in the presence of WzcCDAC, and the corresponding \(R_2\) values are enhanced by increasing concentrations of the latter (and the fraction of Wzb within the Wzb–WzcCDAC complex). That these \(R_2\) values are induced by binding/unbinding processes involving WzcCDAC is reinforced by the fact that the limiting relaxation rates, \(R_{2,\text{eff}}(\infty)\) (inferred from the corresponding values at the highest field used, 1,000 Hz), in the fast pulsing regime for almost all residues (SI Appendix, Fig. S4 for representative examples) are linearly dependent on the bound fraction (\(f_b\)).

As shown in Fig. 1A–E, both the number of Wzb residues that display significant \(R_2\) values, and the values themselves, are enhanced with increasing \(f_b\). The distribution of residues with substantial \(R_2\) values plotted on the Wzb surface for \(f_b = 0.37\) largely mirrors the spatial pattern of attenuations seen for the amide resonances of Wzb in the presence of an equimolar ratio of WzcCDAC (Fig. 1F) reported previously (24). At the lowest \(f_b\) value (0.05), F2, the catalytic C9 (on \(\beta^1\); SI Appendix, Fig. S1B), and R74 (\(\alpha^3\)) show statistically significant \(R_2\) values >5 s\(^{-1}\). At \(f_b = 0.24\), the \(R_2\) values for F2, C9, and R74, in addition to N75 and D77 (on the \(\alpha^3\)–\(\beta^3\) loop), I79 (on \(\beta^3\)), and M82 (on the \(\beta^3\)–\(\alpha^4\) loop), all exceed 15 s\(^{-1}\) (SI Appendix, Fig. S3). Within this group of residues, F2, R74, N75, and D77 are all significantly solvent-exposed and their substantial \(R_2\) values suggest that they are proximal to, if not in direct contact, with WzcCDAC within the Wzb–WzcCDAC complex. These transient chemical shift changes (as reflected by the significant \(R_2\) values) on Wzb were utilized (SI Appendix, Materials and Methods and Table S2 for details) to generate ambiguous interaction restraints (AIRs) in our structure calculation protocol discussed below.

To obtain a direct measure of distances between the interacting partners, we relied on the measurement of paramagnetic relaxation enhancement (PRE) effects induced on Wzb by spin-labeled WzcCDAC variants. First, the two native cysteines on WzcCDAC were mutated to Ser (Cys544Ser/Cys563Ser) to generate a cysteine-less background (WzcCDAC\(\Delta C\)) to which nonnative cysteines could be introduced, one at a time, for the subsequent attachment of a spin label (SI Appendix, Fig. S5). WzcCDAC\(\Delta C\) expressed well and can be purified similarly as WzcCDAC. The \(^{15}\)N,\(^{1}\)H TROSY spectrum of this mutant displays well-dispersed resonances indicative of a folded protein and suggests the absence of any significant structural reorganization compared with wild-type WzcCDAC (SI Appendix, Fig. S6A) with chemical shift perturbations (average ± SD over all resonances excluding those of residues within a 6-Å radius of the mutation sites) of 0.04 ± 0.03 parts per million (ppm). That the structural changes caused by the double mutation are modest is further confirmed by an almost unchanged affinity of WzcCDAC\(\Delta C\) toward Wzb (Table 1).

Given the substantial number of Wzb resonances that are broadened in the presence of equimolar amounts of WzcCDAC, substoichiometric concentrations of spin-labeled WzcCDAC variants would be necessary for precise PRE measurements requiring thoughtful placement of the spin labels. The spin labels would have to be proximal to the \(\alpha^2\)-helix harboring the important Glu102–Arg104 motif, were introduced into the WzcCDAC\(\Delta C\) background (SI Appendix, Fig. S5). All variants express well and are properly folded as indicated by the corresponding \(^{15}\)N,\(^{1}\)H TROSY spectra (SI Appendix, Fig. S6 B–P). The Wzb affinities of all variants, with the exception of WzcCDAC\(\Delta C\),Leu505Cys (Table 1), remain largely unchanged in comparison to WzcCDAC. The \(^{13}\)C,\(^{1}\)H HMQC spectra of Wzb display fast exchange in the presence of increasing amounts of WzcCDAC\(\Delta C\),Leu505Cys (Table 1), contrasting the slow-exchange regime seen for WzcCDAC and all other variants (SI Appendix, Fig. S7). Given the ∼5-fold reduction in affinity, and the likely related modification in its association dynamics, the WzcCDAC\(\Delta C\),Leu505Cys mutant was excluded from further analysis.

The patterns of chemical shift perturbations of methyl resonances (SI Appendix, Fig. S8) of Wzb in the presence of the WzcCDAC variants suggest no significant modification in the overall modes of interaction compared with the wild-type species.

To generate spin-labeled single-cysteine variants of WzcCDAC, we utilized a 3-(2-iodoacetamido)-proxyl spin label (IPSL) that covalently attaches to cysteine via a thioster bond. The efficiency of the spin labeling of \(^{1}\)H-labeled WzcCDAC variants (for use in the NMR experiments) was confirmed by mass spectrometry; a successful labeling reaction is indicated by a mass increase of ∼198 Da for the IPSL-modified species (SI Appendix, Fig. S9 for a representative example). Four IPSL-labeled variants (Val466Cys, Ser516Cys, Gln523Cys, and Lys556Cys), distributed around the Glu102–Arg104 motif, were introduced into the WzcCDAC\(\Delta C\) background (SI Appendix, Fig. S5). All variants express well and are properly folded as indicated by the corresponding \(^{15}\)N,\(^{1}\)H TROSY spectra (SI Appendix, Fig. S6 B–P). The Wzb affinities of all variants, with the exception of WzcCDAC\(\Delta C\),Leu505Cys (Table 1), remain largely unchanged in comparison to WzcCDAC. The \(^{13}\)C,\(^{1}\)H HMQC spectra of Wzb display fast exchange in the presence of increasing amounts of WzcCDAC\(\Delta C\),Leu505Cys (Table 1), contrasting the slow-exchange regime seen for WzcCDAC and all other variants (SI Appendix, Fig. S7). Given the ∼5-fold reduction in affinity, and the likely related modification in its association dynamics, the WzcCDAC\(\Delta C\),Leu505Cys mutant was excluded from further analysis.

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Ratios of intensities of Wzb resonances in the presence of each of the four spin-labeled WzcCDAC variants in their paramagnetic (\(I_{para}\)) and diamagnetic (\(I_{dia}\)) states (SI Appendix, Fig. S12; refer to SI Appendix, Materials and Methods for...
experimental details) show distinct patterns, suggesting that the measured PREs encode significant nonredundant distance information. Residues with statistically significant reduction in intensity ($I_{\text{para}}/I_{\text{dia}} < 0.7$) mapped onto the surface of Wzb (Fig. 2) are localized within well-defined patches, implying that the transiently assembled complex is restricted to a single overall orientation, or a set of very similar orientations.

A set of 59 unambiguous distance restraints (listed in SI Appendix, Table S3 and illustrated in SI Appendix, Fig. S13) arising from the four spin-labeled sites on WzcCDAC, Ser516Cys, Gln523Cys, and Lys556Cys—was supplemented with the AIRs (SI Appendix, Table S2) obtained from the relaxation dispersion measurements on Wzb (described above) and our previously measured spectral perturbations on WzcCDAC (SI Appendix, Fig. S14) (24) and utilized in docking calculations using the HADDOCK suite (27, 28) to obtain the structure of the Wzb–WzcCDAC complex. The importance of the PRE-based unambiguous distance restraints is highlighted by the fact that their inclusion is crucial in obtaining a single dominant structural cluster (ClusterALL,1 and ClusterPRE,1 with ~53 and ~59%, respectively, of the 400 refined structures; compare SI Appendix, Tables S4 and S5 with SI Appendix, Table S6). The overall characteristics of the protein interfaces calculated with or without AIRs are quite similar (SI Appendix, Fig. S15) but with the former set of calculations leading to more favorable HADDOCK scores. Therefore, unless otherwise stated, we focus exclusively on the structures that constitute ClusterALL,1 in the discussion below.

As previously noted, the α2-helix of WzcCDAC, and the invariant Glu508/Arg511/Arg514 triad therein, has been predicted to play a crucial role in the recognition of Wzb (24). A statistical analysis of the 212 structures that constitute ClusterALL,1 suggests that WzcCDAC residues that comprise the interface (defined as those that lie within 4 Å of any Wzb residue; SI Appendix, Fig. S16) includes all three residues of the triad, together with the intervening Ser512 in all constituent structures. His518, Phe519, and Met522, all on α2, and Asn555 and Arg557, on the α3–β2 loop, are found at the interface in >80% models. The complementary interface on Wzb involves residues (found in >80% of the structures; SI Appendix, Fig. S16) of the N terminus (F2), the α3–β3 loop (D77), the α4–β4 loop (P95, R98, G99, K100), the β4-strand (M102), the D loop (β3′–α1′ loop; H106, W107), and the C terminus (E145, Q146, V147). These same residues are also part of the protein–protein interface in the other major clusters, ClusterALL,2, ClusterALL,3, and ClusterALL,4 (SI Appendix, Fig. S16).

The interaction results in the burial of relatively modest surface areas of 757 and 747.2 Å² for WzcCDAC and Wzb, respectively, a likely explanation for their relatively weak interaction. Despite the presence of a significant number of polar residues at the interface, only seven intermolecular hydrogen bonds/salt bridges are consistently detected, of which only two involve Wzb side chains, D77 and R98, that form salt bridges with Asn555 and Glu508, respectively. The position corresponding to D77 is invariant in LMW-PTPs. The R98 position, though somewhat less conserved, is generally a basic (histidine or lysine) residue when substituted (SI Appendix, Fig. S17). Hydrogen bonds of the G99 and V147 main chains with Arg514 and Asp637 side chains, respectively, provide additional stability at the interface. These interactions are supplemented by those involving hydrophobic clusters on the two partners comprising H106 and W107 on Wzb and His518, Phe519, and Met522 on WzcCDAC. In LMW-PTPs, the position corresponding to H106 is somewhat less conserved, though it is a histidine in a large number of cases; the W107 position is largely conserved or contains an aromatic or a large hydrophobic residue in cases of deviations (SI Appendix, Fig. S17). Key interfacial interactions are illustrated in Fig. 3.

To test whether the structure of the Wzb–WzcCDAC complex (missing the Y-cluster) is compatible with Y-cluster dephosphorylation, we utilized the MODELLER suite (29) to attach the C-terminal tail fragment (Tyr705 to Lys720) onto the docked structures (SI Appendix, Materials and Methods for details). Five of the six Y-cluster tyrosine residues (Tyr708, Tyr710, Tyr711, Tyr713, and Tyr715) can access the Wzb active site in optimal fashion within the complex. The distances of the -OH moieties of these residues from the thiol group of C9 and the guanidino group of R15, both part of the catalytic C-[X]5–R-[S/T] motif (30), are within (or very close to) their optimal distances of 3.3 Å.
and 2.6 Å (31, 32), respectively (SI Appendix, Fig. S18). In contrast, these distances deviate significantly for Tyr705, suggesting that the dephosphorylation of this residue, if it occurs at all, is expected to be highly inefficient. It is notable that Tyr705 is the only position on the Y-cluster that has been shown not to be auto-phosphorylated in Wzc (17).

We have previously demonstrated that a Glu508Ala/Arg511Ala/Arg514Ala mutant, where the conserved residues of the $508\text{Glu}^\text{[Xxx]}_2\text{Arg}^\text{[Xxx]}_2\text{Arg}^{514}$ motif of WzcCD have been replaced by alanine, is severely compromised in its ability to interact with and be dephosphorylated by Wzb (24). Those results support the involvement of these residues in recognizing Wzb in line with our current structure. To test the complementary surface on Wzb, we mutated D77 and R98 (discussed above) individually to alanine. While the Wzb$_{R98A}$ mutant expresses well and can be purified to homogeneity, the expression of the Wzb$_{D77A}$ is suboptimal, suggesting that this mutation likely destabilizes the protein. We attributed this decreased stability to the loss of a salt bridge (Fig. 3) between

![Diagram of key interaction](image)

Fig. 3. Key interaction at the protein–protein interface in the Wzb–WzcCD$_\Delta$C complex. The structure of the lowest-energy model (from Cluster$_\text{ALL}_1$) is shown in surface (Top) and ribbon (Middle) representation. Interacting elements on each partner that comprise the interface are indicated. Expansions of specific regions (indicated by the colored rectangles) illustrating key interactions that stabilize the interface are shown (Bottom). The cyan dashed lines indicate intermolecular hydrogen bonds; the green dashed line indicates the intramolecular D77–K100 salt bridge discussed in the text.

![Graph of dephosphorylation](image)

Fig. 4. Dephosphorylation of WzcCD by Wzb and variants. Reaction mixtures containing fully phosphorylated WzcCD were incubated for various time intervals (0, 1, 3, 5, and 10 min) with either wild-type (WT) Wzb or forms carrying double (D77K/K100D) or single (R98A) mutations at key sites at the protein–protein interface. Representative immunoblots for the time courses are shown in A and the corresponding fits to extract apparent dephosphorylation rates (normalized to 100%) are shown in B. Also shown are the positions of the molecular weight markers (in kDa). Note that the same lane corresponding to the molecular weight markers has been shown on the left hand sides of the time traces for each of the D77K/K100D and R98A mutants of Wzb on the bottom plots in panel A. This has been done to better guide the eye.
Fig. 5. Schematic illustration of the phosphorylation-coupled assembly/disassembly that drives BY-kinase function. At low levels of Y-cluster phosphorylation (Y-low assembled; represented as fully dephosphorylated for illustrative purposes), the cytoplasmic catalytic domain (WzcCD) of Wzc (green) forms an octameric ring. This assembly allows each WzcCD subunit to simultaneously act as an enzyme, accommodating the Y-cluster of the preceding subunit (l-1) at its active site, and as a substrate, inserting its own Y-cluster into the active site of the following subunit (l+1). Oligomerization enables exchange of ATP Mg\(^{2+}\) (dark red) for the bound ADP (purple), facilitating the progression to chemistry. At this stage, the diffusive binding of cytosolic Wzb (orange) is rendered inefficient due to the enhanced local concentration of WzcCD monomers in the context of the membrane-associated species. Following several rounds of chemistry (phosphorylated tyrosine residues are indicated by blue spheres) and enhanced Y-cluster phosphorylation (Y-high assembled), the electrostatic repulsion (red arrows) increases above a critical threshold, making homotypic interactions between WzcCD monomers disfavored over heterotypic interactions. After several rounds of Wzb-mediated dephosphorylation, the electrostatic repulsion is once again lowered below the critical threshold (Y-low disassembled), allowing the successful exclusion of Wzb and regeneration of the oligomeric state of WzcCD (Y-low assembled), thus reinitiating the cycle.

Discussion

The structure of the Wzb-WzcCD\(_{AC}\) complex determined here not only reveals key structural features that enable molecular recognition between these two species but also provides a mechanistic framework to understand how the assembly/disassembly cycle correlates to Y-cluster phosphorylation states (8). As shown above, the \(\alpha\)2-helix of WzcCD plays a key role in the recognition of Wzb. Indeed, interactions involving this helix, both polar and hydrophobic, appear to be conserved within BY-kinase/LMW-PTP pairs by comparing predicted structural interfaces between several related BY-kinase/LMW-PTP pairs from gram-negative species (SI Appendix, Fig. S19). As has been mentioned before, the \(\alpha\)2-helix is also central to the formation of the oligomeric assembly (12) necessary to enable the active site of an enzyme-acting protomer to receive and phosphorylate the C-terminal Y-cluster of an adjoining substrate-acting subunit. Therefore, a competition for the \(\alpha\)2-helix site between the homotypic and heterotypic interactions involving WzcCD monomers and Wzb, respectively, is expected. Thus, the processes of phosphorylation and dephosphorylation would need to be temporally separated for maximal efficiency. We suggest that this is achieved through the intricate mechanism illustrated in Fig. 5. As mentioned before, WzcCD is the cytosolic domain of a protein that is embedded in the bacterial inner membrane (8) (SI Appendix, Fig. S1A). Indeed, all BY-kinases need to form a membrane-associated species for activity (5). This arrangement results in the formation of an octamer through interactions between WzcCD monomers (8), supplemented by extracytoplasmic interactions, in the absence of Y-cluster phosphorylation. This membrane-bound assemblage greatly enhances the local concentration of WzcCD monomers and shields the \(\alpha\)2-site from diffusion-controlled access by the wholly cytosolic Wzb. This allows phosphorylation to proceed [likely in sequence from the C terminus to the N terminus (8) of the Y-cluster] without the counteracting effects of premature Wzb-driven dephosphorylation. Once the Y-cluster is sufficiently phosphorylated, the homotypic affinity between WzcCD monomers is reduced due to strong electrostatic repulsion resulting from the significant accumulation of negative charge on the Y-cluster. When this repulsion exceeds a certain critical threshold, predicted to be at about four phosphorylated tyrosine residues based on functional data (8), the WzcCD monomers are pushed apart, allowing Wzb to successfully outcompete the intra-WzcCD interactions and dock at the \(\alpha\)2-site to initiate dephosphorylation. After the dephosphorylation of a sufficient number of Y-cluster tyrosines, the repulsion between WzcCD monomers decreases below the critical threshold, allowing their reassociation and the successful exclusion of Wzb, thereby reinitiating the cycle. It has also been suggested that oligomerization of WzcCD monomers is necessary to enable the exchange of product adenosine diphosphate (ADP) for substrate adenosine triphosphate (ATP) (33). Given that WzcCD possesses significant ATPase activity (34), oligomerization would have to precede ATP binding to prevent futile hydrolysis. Specific features within the \(\alpha\)2-helix have also been suggested to be coupled to that
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Materials and Methods

Protein Expression and Purification. All constructs (and variants) of the catalytic domain of E. coli (K-12) Wzc (447 to 725; WzcCD), its truncated form (lacking the C-terminal tail, 447 to 704; WzcCDΔC), and full-length E. coli (K-12) Wzb were expressed and purified using protocols described at length in SI Appendix, Materials and Methods. A list of constructs used in this study is given in SI Appendix, Table S1.

NMR Experiments. All NMR experiments were recorded at 25 °C using Bruker Avance III spectrometers operating at 600, 700, or 800 MHz, all equipped with cryogenic probes capable of applying pulsed-field gradients along the z axis. Additional details of specific experiments and methodology may be found in SI Appendix, Materials and Methods.

Computational Protocols. Structure calculations were performed using ambiguous interaction restraints (SI Appendix, Table S2) from the measurement of transient chemical shifts (Wzb) and static spectral perturbations (WzcCDΔC) in combination with PRE-based unambiguous restraints (SI Appendix, Table S3) using the HADDOCK suite (27). Procedures used for the preparation of starting structures for WzcCD and Wzb, generation of experimental restraints, and computational protocols used are described in detail in SI Appendix, Materials and Methods.

Protein Dephosphorylation Assays. The ability of phosphorylated WzcCD to be dephosphorylated by wild-type Wzb and corresponding D77K/K100D and R98A mutants was tested by immunoblotting using the monoclonal anti-phosphotyrosine antibody PY20 (dilution 1:5,000; Invitrogen). Experimental details are provided in SI Appendix, Materials and Methods.

Data Availability. The HADDOCK-generated structural ensembles are available through the Open Science Framework (OSF). (https://osf.io/fmwda/) (45). All other study data are included in the article and/or SI Appendix.

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