Phenol sensing in nature is modulated via a conformational switch governed by dynamic allostery

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Jayanti Singh\(^1\)\(^*,\) Mohammad Sahil\(^2,4\)\(^,\) Shamayeeta Ray\(^,\) Criss Dcosta\(^,\) Santosh Panjikar\(^3,4\)\(^,\) G. Krishnamoorthy\(^7\)\(^,\) Jagannath Mondal\(^1,5\)\(^*,\) and Ruchi Anand\(^1,6\)\(^*\).‡

From the \(^1\)Department of Chemistry, Indian Institute of Technology Bombay, Mumbai, Maharashtra, India; \(^2\)Tata Institute of Fundamental Research, Hyderabad, Telangana, India; \(^3\)Department of Biochemistry and Molecular Biology, Monash University, Clayton, Victoria, Australia; \(^4\)Australian Synchrotron, Clayton, Victoria, Australia; and \(^5\)Department of Biotechnology, Anna University, Chennai, Tamil Nadu, India

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The NtrC family of proteins senses external stimuli and accordingly stimulates stress and virulence pathways via activation of associated σ\(^{54}\)-dependent RNA polymerases. However, the structural determinants that mediate this activation are not well understood. Here, we establish using computational, structural, biochemical, and biophysical studies that MopR, an NtrC protein, harbors a dynamic bidirectional electrostatic network that connects the phenol pocket to two distal regions, namely the “G-hinge” and the “allosteric linker.” While the G-hinge influences the entry of phenol into the pocket, the allosteric linker passes the signal to the downstream ATPase domain. We show that phenol binding induces a rewiring of the electrostatic connections by eliciting dynamic allostery and demonstrates that perturbation of the core relay residues results in a complete loss of ATPase stimulation. Furthermore, we found a mutation of the G-hinge, ∼20 Å from the phenol pocket, promotes altered flexibility by shifting the pattern of conformational states accessed, leading to a protein with 7-fold enhanced phenol binding ability and enhanced transcriptional activation. Finally, we conducted a global analysis that illustrates that dynamic allosterically-driven conserved community networks are universal and evolutionarily conserved across species. Taken together, these results provide insights into the mechanisms of dynamic allosterically-mediated conformational changes in NtrC sensor proteins.

Proteins are involved in several biological processes where they undergo various conformational fluctuations, modulated by diverse signals such as ligand binding, environmental factors, etc. (1–3) Protein breathes through complex multidimensional landscape, which constitutes various conformational states (4, 5). For instance, allostery in proteins where binding of a ligand/effector at a distal site can stabilize a particular conformation can be one of the strategies adopted by nature to enable function (6). Here, long-distance communication can either be governed by conformational allostery, which is accompanied by a significant structural change within the protein structure, or it can be dynamic, where no detectable conformational change is observed (7–12). While in the case of conformational allostery, enthalpic contributions play a central role, in dynamic allostery, entropy effects are considered to be predominant (8). This is due to the fact that energetic perturbations, which are generally caused by small-scale internal side chain changes, can result in reorganization of the interaction network of the whole protein (7, 13). Most commonly used techniques to understand allosteric mechanism are by combining structure determination tools such as X-ray crystallography, NMR, and more recently cryo-EM with biophysical and computational tools. The structural tools partly provide snapshots of selective conformations of a protein, within its functional cycle, the supporting techniques help in delineating associated conformational heterogeneity that together help in unraveling the mechanism of complex allosteric regulation.

NtrC family of protein belongs to the broader bacterial enhancer binding proteins that assemble into ATPase motors. They serve as mechano-enzymes by triggering activation of σ\(^{54}\)-dependent RNA polymerase (RNAP) (14, 15). They respond to external stimuli and via either phosphorylation of their signal sensing domain or by entry of a particular ligand and in some cases via protein–protein interaction activate the RNAP for downstream transcription of select pathways (15). MopR, an NtrC family protein from *Acinetobacter calcoaceticus* NCIB8250, is a multidomain protein where binding of phenol to its N-terminally located ligand binding [A] domain results in derepression of ATP hydrolysis via the centrally located AAA+ [C] domain (14, 16). It also harbors a C-terminal DNA binding [D] domain, which assists MopR to latch onto a specific DNA (Fig. 1A, S1A) (14, 16) segment, upstream activation site, that lies 100 to 200 bases upstream of the RNAP-binding region. It is hypothesized that in presence of phenol and ATP, MopR assembles into an oligomeric structure and executes its mechanofunction by activating the σ\(^{54}\)-RNAP, which then triggers the downstream phenol degradation pathway. This ability of MopR to sense phenol and elicit a

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8 Both authors have contributed equally to the manuscript.

* For correspondence: Ruchi Anand, ruchi@chem.iitb.ac.in; Jagannath Mondal, jmondal@tifrh.res.in.

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Response has been used to create a plethora of sensors for monitoring concentrations of several xenobiotics in polluted water (14, 17–19). The ligand-binding domain of MopR is connected to the AAA+ domain via a connector B-linker helix, and it is envisioned that binding of phenol brings about a global allosteric change, transmitted via the B-linker, that is crucial for activating ATP hydrolysis, thereby allowing MopR to elicit its function (14, 20, 21).

There has been paucity in obtaining structural information in this family of enzymes. The structure of the signal sensing domain [A] and a portion of the tandem B-linker of MopR (MopRAB) was determined in complex with its cognate ligand, phenol (Fig. 1B, PDB ID: 5KBE), and a few of its derivatives in 2016 (14). The structure revealed that the ligand-binding domain of MopR forms a homodimer and belongs to the nitric oxide signaling fold that encompasses closely related bacterial homologs such as dimethylphenol regulator (DmpR) that responds to 2,3-dimethylphenol and XylR that senses benzene. Recent studies have further revealed that for sensor activity, MopR exhibits a concentration-dependent oligomerization with dimers converting to hexamers via a tetrameric intermediate (22). MopR also showed structural similarity to evolutionarily distant eukaryotic proteins that bind fatty acids and are part of a large complex that partakes in transport across the golgi (14, 22). The [A] domain of MopR harbors a zinc-binding site, which is ~10 Å away from the phenol-binding region, whose function is attributed to maintain structural integrity (14). The structure was especially

**Figure 1. Structural properties of MopRAB.** A, domain organization of MopR. B, MopRAB (residues 1–229) (PDB ID: 5KBE) dimer with one protomer in cartoon representation and other in surface representation (deep salmon) showing phenol completely buried. C, phenol-binding pocket of MopRAB. D, superimposition of a representative snapshots, 900 ns and 950 ns of the apo (pale blue) and bound (cyan) MopRAB structures. The G-hinge region is shown in pale magenta and magenta for apo and bound MopRAB, respectively. The allosteric linker region is shown in light pink and wine red for apo and bound MopRAB. E, RMSD plot of apo and bound MopRAB depicting conformational changes in the α5 (G-hinge) and allosteric linker region. PDB, Protein Data Bank.
instrumental in understanding the key features that lead to selective binding of phenol. In brief, the binding pocket of phenol is mostly composed of hydrophobic residues that stabilize the aromatic ring of phenol via π-stacking interactions (14). In particular, the phenolic OH moiety is anchored in the active site via hydrogen bonding interactions by key sensor residues, H106, and W134 (Fig. 1C). The snug nature of the pocket imparts its selectivity such that monosubstituted phenols bind with slight reduction in binding affinity, whereas bulkier phenols are unable to bind (14, 17–19). The OH anchor ensures that non-phenolic aromatic hydrocarbons are unable to occupy the phenol pocket (18, 19).

More recently the X-ray crystal structure of a homolog of MopR, DmpR, that encompasses both the sensor along with the ATPase domain has also been determined (23). In this structure, the relative orientations of the [A] and the central AAA+[C] domains when phenol is bound to the [A] domain could be visualized. However, since most AAA+[C] motors assemble as hexamers or heptamers, it is difficult to discern if the captured tetrameric form is an inactive or an activated conformation of the enzyme (15, 24, 25). The structure of the unliganded form of MopR and its homologs has not been determined till date. Since MopR is very selective toward its ligands, here we explore conformational changes that enable the enzyme to bind its ligand. Moreover, how ligand binding initiates derepression of the downstream ATPase domain, the associated signal transduction pathway that connects the two domains, is also not well understood, resulting in the overall mechanism to remain elusive. Therefore, as a first step to understand how the binding of ligand (phenol) induces significant conformational changes both within the [A] domain of the protein as well as those that percolate outside to the adjacent B-linker region, we carried out a combination of experimental and molecular dynamics (MD) simulation studies. Further, regions of the enzyme that reside outside the phenol pocket that may influence and play a critical role in facilitating function were also explored. The study in particular helped in unraveling how dynamic networks that run within the protein play a critical role in communication. More importantly, it establishes a conserved community network relay that also exists in other aromatic ligand binding subclass of NtrC family of proteins. It especially entails how distal regions far from the binding or active sites play a significant role in flow of information and thereby control overall protein function.

Results

Dynamic networks in MopR

Analysis of the crystal structure of the MopRAB in the phenol-bound form shows that the binding pocket is buried inside the protein and there is no direct path to facilitate ligand entry (Fig. 1B). Various attempts to crystallize the unliganded form of MopRAB (Fig. S1B) were unsuccessful, probably owing to the increased flexibility of the protein in absence of phenol. Therefore, we undertook MD simulations to achieve a conformation of the MopR sensor domain (MopRAB) (using PDB ID: 5KBE) in its apo state.

Comparison of the equilibrated structure of apo with that of phenol-bound states of MopRAB reveals that although the phenol pocket still remains buried, there are several other regions of the enzyme that show localized changes in structure. Maximum change was observed in two regions (Fig. 1D): first, near the vicinity of the α2 (residues 48–62) and α5 (residue 138–147) and its following loop (residue 148–154) and the other being at the end of the sensor domain from α7 and region preceding it (residues 195–225). The RMSD (Fig. 1E) and RMS fluctuation plots (Fig. S2) confirm that these changes are consistent through the simulation.

In the first region, upon binding of phenol, the helix α5 rotate away from α2 by about 20°, pulling the connector loops with it (Fig. 1D). To gauge the significance of the associate motions, this region was further subjected to a bioinformatic analysis, which revealed that the NtrC subset of proteins that bind aromatic ligands (Fig. S3, Table S1) contain a conserved glycine residue in the flexible ‘GAS’ motif (constituting of amino acids G148, A149, and S150 in MopR) (Fig. S3, 1D), which resides at the tip of α5. Previous in silico studies have identified such small flexible residues like glycine, serine, and alanine as hinge regions, and reports have attributed these motifs to be present where flexibility of motion is a prerequisite (26–28). The movements about strategic hinge regions have also been shown to play an important role in facilitating protein–ligand interactions or in domain movements to bring about allosteric communication (27, 29). Using this premise, we referred to this region as the “G-hinge region” (Fig. 1, D and E). The second region where substantial motion was observed is the helix-loop motif that connects to the B-linker (Fig. 1, D and E). In the NtrC family of proteins, the B-linker serves as a connection point between the N-terminal sensor and the tandemly located ATPase domain (14, 15, 20). The sensing activity is attributed to be turned on in the presence of the sensor molecule, phenol, and the information is passed through this connector region (15, 21). Comparison of MD snapshots of apo versus bound forms of MopRAB (Fig. S1B) shows that the secondary structure in this region gets reorganized upon phenol binding (Fig. 1, D and E). Since this region is ~20 Å away from the phenol-binding site and a conformational change was nevertheless observed in response to phenol binding, the region has been referred henceforth as the “allosteric linker region” (Fig. 1D).

To gain deeper insights into the process of how ligand binding propagates downstream signal relay, in-depth analysis of the apo and bound form of the structures was undertaken. It is a well-established fact that signal transduction in protein from one point to another propagates by gain and loss of a set of the interactions and several proteins operate via a large interconnected network of interactions (30). In order to unravel such an associated network of residues that bring about the communication of the allosteric linker and the G-hinge region, hydrogen bond propensities were calculated from the simulation trajectory (Figs. 2 and 3, S4, Tables S2–S4). MD results show that binding of phenol creates space in the
binding pocket and drives a conformational change such that W134 and H106 shift outward (Table S5). For instance, H106 imidazole sidechain reorients to form hydrogen bonds with phenol and V112 (Fig. 2, A and B), and this motion in turn leads to decreased hydrogen bond propensities for the adjacent Q113(β4)-L135(β5) and M111(β4)-F138(α5) pairs that otherwise formed stable hydrogen bonds in the apo state (Fig. 2, A and B). This expansion of the binding pocket, upon phenol binding, percolates to α5 and results in its separation from α2 and α6 as represented by a rotation of ~20° of α5 with respect to α2 in the apo versus the phenol binding network, percolates to α5 and results in its separation from α2 and α6 as represented by a rotation of ~20° of α5 with respect to α2 in the apo versus the phenol binding network. This is also exemplified by decrease in overall hydrogen bond propensities between residues on α5 with that of α2 and α6. For example, E139 of α5 with R55 of α2 and W156, M157, and L158 of α6 (Fig. 2, C–F) all show a decrease in hydrogen bond propensity. E139 of α5 with R55 of α2 and W156, M157, and L158 of α6 (Fig. 2, C–F) all show a decrease in hydrogen bond propensity. E139 of α5 with R55 of α2 and W156, M157, and L158 of α6 (Fig. 2, C–F) all show a decrease in hydrogen bond propensity. E139 of α5 with R55 of α2 and W156, M157, and L158 of α6 (Fig. 2, C–F) all show a decrease in hydrogen bond propensity.
formed. Y176 also gets reoriented via its backbone hydrogen bonding with amide group of Q177. It is noteworthy that Y176 acts as a bridging residue and upon phenol binding, Y176 additionally interacts with phenol through aromatic stacking interactions (Table S6). The conformational rewiring in the aforementioned binding pocket residues percolated to other adjacent residues, specifically a decrease in hydrogen bond propensities of W199 sidechain with I175 and K194 was observed (Fig. 3, A and B). The W199 consequently get released, hence relaxing the adjacent loop (202–210 of allosteric linker region), which then restructures into a helix ($\alpha7'$) as predicted by patterned change in the hydrogen bond propensities of C=O of ith residue and NH of (i+4)$^{th}$ residue (Fig. 3, A and B). Thus, the MD strongly suggests that sequential changes in the hydrogen bonding patterns allosterically connect the binding pocket and the allosteric linker region.

To summarize, the presence of phenol is percolated to the two extreme ends of the structure which is exemplified via changes in the hydrogen bond propensities. Apart from the predominant electrostatic effect that seems to govern the relay network in MopR, it was also noticed that an underlining subtle hydrophobic effect may play an integral role in effecting the relay. For instance, residue Y165 and Y176 that have strong
Dynamic allostery in phenol sensing protein, MopR

shift in hydrogen bonding propensities upon phenol binding also form the wall of the phenol pocket and provide stabilizing stacking interactions that assist in phenol binding (Table S6). Hence, these residues support the relay network via dual contributions.

Structural and thermodynamic probing of G-hinge region

To validate the MD results and discern the functional role of the G-hinge in ligand binding, we decided to mutate this region. It has been established that proline when introduced at the end of a helix has a destabilizing effect on the helix integrity (31, 32). Moreover, in proteins that regulate the biosynthesis of aromatic compounds, the glycine residue has been shown to play an integral role in establishing a communication network between the active site and the distal amino acid binding regulatory site (33). Hence, with the 2-fold objective shown to be obliterated upon introduction of the G to P mutation in this system (30). This network was a destabilizing one, a G148P mutation was performed. The change the capping residue on the α5 from a stabilizing glycine to a capping one, a G148P mutation was performed. The G148P mutant was expressed and purified and a comparison of the CD spectra of the G148P mutant (MopRG148P) with the WT protein (MopRAB) shows that the mutation does not cause perturbation in the overall secondary structure (Fig. S5, A and B). Tm, studies reveal that the thermal stability of MopRG148P is approximately 5 °C less as compared to that of the native protein. Phenol binding leads to an overall enhancement of the thermal stability for both the native as well as MopRG148P with the Tm for both proteins increasing to ~82°C (Fig. S5, C and D). Additional, size-exclusion chromatography studies confirm that oligomeric state of the MopRG148P is maintained as a dimer form in solution, similar to that observed for the WT (Fig. S6); thus, this mutation does not perturb the structure in an adverse fashion. To discern if this mutation has any functional relevance, isothermal titration calorimetry (ITC) studies were carried out. Surprisingly, ITC studies show that MopRG148P exhibits a 7-fold higher binding affinity compared to MopRAB toward phenol (Fig. 4, A and B; Table S7) with a Kd value of (0.07 ± 0.02 μM). The data suggest that the effect of G148P substitution that is situated at a distance ~20 Å percolates to the MopRAB pocket, making the protein more conducive to phenol binding. The ITC corroborate our MD simulation results and reassert that there is indeed some communication between the phenol pocket and the G-hinge region. The G148P mutation likely results in a rearrangement of the network, which is the dominant reason for the 7-fold increase in phenol affinity. Since the G-hinge dynamics seems to be correlated with the pocket, we speculate that this region plays a role in either facilitating entry of the ligand or coordinated motions in this region that assist in phenol pocket formation.

In order to further validate the role of the α5 region, we additionally attempted to crystallize the apo and phenol-bound form of the hinge mutant MopRG148P; however, the crystals were only obtained in the presence of phenol. The structure of the sensor domain (Fig. S1B) of MopRG148P in complex with phenol was determined to a resolution of 2.3 Å (Protein Data Bank [PDB] ID: 7VQF) by molecular replacement (MR) method using the native MopRAB-phenol structure as the search model (PDB ID: 5KBE). The data processing and refinement statistics are provided in Table 1. Analysis reveals that the overall structure of the sensor domain in the phenol-bound form is similar with a RMSD of 0.63 Å aligning 180 residues (alpha carbon atoms) (Fig. 4D). Comparative analysis shows that the introduction of a proline residue in G148P mutation indeed destabilizes α5 and induces its shortening such that F147 that was previously part of α5 now resides on a loop (Fig. S7). Another notable observation from the crystal structure was a marked conformational change in α7 region that forms the allosteric hinge. Here, it was observed that residues 212 to 225 (α7) in the native protein, are no longer visible. Rather this region becomes disordered and instead forms the allosteric hinge to the AAA+ domain. Thus, it was surprising to observe that perturbation in the G-hinge, which is separated spatially by more than 20 Å, causes a major rearrangement in the allosteric linker region. Since crystallography can only provide static snapshots, this result intrigued us to further probe protein dynamics in both apo and phenol-bound forms.

Fluorescence studies to gauge protein dynamics

In order to understand the conformational heterogeneity in the apo versus the phenol-bound forms of MopRAB, (Fig. S1B) as well as changes induced on introduction of the G148P mutation, tryptophan fluorescence studies were undertaken. Three tryptophan residues naturally located respectively in the three strategic regions of the protein, that is, phenol pocket cap region, G-hinge region, and allosteric linker region were used as markers to study the local environment for both the native and G148P mutant (Fig. 4C).

For generating each single tryptophan construct, all other tryptophan residues were mutated to alanine or phenylalanine. The following constructs were made: MopRW156, MopRW134, MopRW199, MopRW156, MopRG148P, MopRW134, G148P, and MopRW199, G148P (Experimental procedure and Table 2, superscript refers to the tryptophan residue retained). The structural integrity of all the mutants was tested and it was confirmed that it was maintained (CD spectra shown in Fig. S9).

Analysis of the steady-state fluorescence data, under saturating concentrations of phenol, reveals that for the native constructs, MopRW134 shows maximum fractional decrease in fluorescence intensity between the apo and phenol-bound forms (Figs. 4G, S10). This was not surprising as W134 is a key phenol pocket residue. This change in fluorescence intensity of MopRW134 could be attributed either to quenching...
observed as a result of phenol binding or due to restructuring of the phenol-binding pocket. Steady-state studies were also performed with the hinge mutant MopRG148P mutant that exhibits 7-fold enhanced phenol binding. Surprisingly, we observe that addition of phenol does not cause significant change in the fluorescence intensity of MopR^W134^G148P (Fig. 4G). Thus, if the quenching in MopR^W134^ was due to phenol binding, a similar effect should be observed in both the MopR^W134^ and MopR^W134^G148P, which was though not the case. Consequently, in MopR^W134^, the change in fluorescence can be attributed to a restructuring of the sensor pocket, which is now primed for phenol binding and not an effect of direct quenching by phenol. Further, ITC-derived thermodynamic parameters (Table S7) also show that the measured change in enthalpy (ΔH) upon phenol binding is lower in G148P mutant relative to the native MopR^AB^, reflecting lesser reorganization.
required for the pocket to accommodate phenol in the case of the mutant. In addition, the total entropy change associated with binding, $\Delta S$ shows a net increase in the case of G148P as compared to native MopRAB, suggesting a preformed pocket for MopRG148P.

To reiterate, the fluorescence and the ITC data point to a more compact structure of the phenol pocket in the hinge mutant as compared to the native. To further corroborate the changes observed via our initial steady state studies, supporting fluorescence lifetime studies were also performed. The observations from the steady-state experiments (Figs. 4G, S10), at all the three tryptophan positions in both native and G148P mutants, are largely reproduced in the mean fluorescence lifetime ($\tau_m$) (Fig. S11 and Table S8). For instance, we observe a consistent decrease in $\tau_m$ upon binding to phenol in the MopR134 variant, whereas this trend was not observed in the MopR134G148P mutant, reasserting that there has been an arrangement in the phenol pocket region upon introduction of the G148P mutation.

To obtain better insight into the conformational dynamics in the apo versus the phenol-bound forms, lifetime studies were combined with potassium iodide (KI) quenching for both MopRAB and MopR148P proteins (Fig. 4, H and I, S12). Due to the larger size as well as the charged nature of the iodide ion, it exhibits high polarizability and hence it cannot penetrate into the hydrophobic core. Thus, KI quenching studies provide a direct measure of the surface accessibility of a particular tryptophan residue via measurement of its $k_q$. Table 3 and Fig. 4, H and I list the values of $k_q$ estimated for all the proteins with and without phenol. A significant decrease in the value of $k_q$ was observed for MopR134G148P in comparison to MopR134 variant. This reasserts that the apo form of MopR148P is more compact in the vicinity of the phenol-binding region and this pocket has become less accessible to solvent. Evidence that the pocket is indeed preformed in the MopR148P is strengthened as $k_q$ for both the apo and phenol-bound forms were found to be similar in this mutant. Furthermore, the $k_q$ was comparable to the phenol-bound form of MopRAB, corroborating the fact that

| Table 1 | Crystallographic data |
|---------|-----------------------|
| Processing | MopR148P + phenol (PDB ID: 7VQF) |
| Wavelength ($\AA$) | 1.54 |
| Resolution ($\AA$) | 33.30–2.30 |
| Space group | C2221 |
| Unit cell | $a=62.20$, $b=118.30$, $c=56.70$ |
| Total reflections | 58,593 (6957) |
| Unique reflections | 9592 (1131) |
| Multiplicity | 6.10 (6.15) |
| Completeness (%) | 99.6 (99.9) |
| $I/\sigma(I)$ | 9.26 (2.29) |
| CC1/2 | 0.99 (0.77) |
| $R_{merge}$ (%) | 13.5 (88.2) |

Refinement

| Total no. of nonhydrogen atoms | 1580 |
| Total no. of protein atoms | 1547 |
| Total no. of water atoms | 21 |
| Total no. of ligand atoms | 11 |
| Total no. of metal atoms | 1 |
| No. of reflections in refinement | 9589 |
| No. of reflections in test set | 480 |
| $R_{factor}$ (%) | 22.1 |
| $R_{free}$ (%) | 26.0 |
| Bonds ($\AA$) | 0.004 |
| Angles (deg) | 0.657 |
| Most favored region (%) | 96.3 |
| Additional allowed region (%) | 3.74 |
| Disallowed region (%) | 0.00 |
| Mean B factors ($\AA^2$) for overall structure | 38.0 |

Statistics for MopR148P in the presence of phenol. Values ( ) are for the highest resolution shell.

$R_{merge} = \sum_{hkl} \sum_{i} |I_i(hkl)| - (\langle I(hkl) \rangle)^2 / \sum_{hkl} \sum_{i} I_i(hkl)$, where $I_i(hkl)$ is the $i$th intensity measurement of reflection $hkl$, $\langle I(hkl) \rangle$ its average.

| Table 2 | Tryptophan mutants |
|---------|--------------------|
| Mutations made | Final constructs |
| W37F_W199F_W156F | MopR134 (Active site) |
| W134A_W199F_W37F | MopR134W199F (Zinc site) |
| W37F_W156F_W134A | MopR134W199F (Linker site) |
| W37F_W199F_W156F_G148P | MopR134G148P (Active site with hinge) |
| W134A_W199F_W37F_G148P | MopR134G148P (Zinc site with hinge) |
| W37F_W156F_W134A_G148P | MopR134G148P (Linker site with hinge) |

Description of various single tryptophan mutants of MopRAB.
very little change in local environment occurs upon phenol binding.

The other tryptophan residues such as W156, which is in the vicinity of the α5 hinge region, were also probed via KI quenching experiments. It was observed that W156 does not show any local environmental changes between the apo and phenol-bound forms in the native protein. However, the G148P mutant has a different profile from that of the native where W156 is now more exposed to the solvent, as a result of the mutation. This observation was corroborated by the simulations performed on the MopRG148P mutations. Here, it was observed that presence of a proline at G148 results in disruption of the stacking interaction between Y59 and F147, which results in an increase accessibility of W156 in the mutant protein (Fig. S13, Table S9). This, we believe, is a consequence of the helix α5 being destabilized in the G148P mutant, as observed in the MopR G148P crystal structure. Overall, it is important to note that the changes in accessibility are observed only as the function of mutation and not due to phenol binding, suggesting that the mutation may have a global effect in altering the overall relay network.

The third tryptophan W199 resides in the allosteric hinge region. Snapshots from MD trajectory show that allosteric linker region, on which W199 resides, shows a significant conformational change upon phenol binding. Moreover, since this helix adopts different conformations in both the phenol-bound MopR AB and MopR G148P crystal structures, we envision this region to be dynamic. Since in other NtrC family proteins, B-linker is implicated in controlling ATPase activity via transmission of information between the signal sensing domain to the ATPase domain, via the B-linker, we believe that motions in the B-linker may have bearing on the derepression mechanism (14, 20, 21). In MopR, we have exploited W199 as a reporter residue to gauge the conformational dynamics of this region. By observing the solvent accessibility of W199 in various states along with MD and structural data, we aim to provide insights into the mechanism of communication. Comparison of kq values obtained from KI quenching studies shows that W199 region is indeed dynamic and reorients upon phenol binding. However, the maximum difference in accessibility was observed between the phenol-bound states of MopRAB and MopRG148P. Comparison of the crystal structure of the MopR AB versus MopR G148P clearly shows that this region has a dramatic reorganization of the linker in both cases. While in MopRAB, the W199 is partially shielded by residues lining α7 in the G148P mutant, this helix is disordered exposing the W199 to the solvent (Fig. 5). This is supported by enhanced KI quenching as exhibited by a 3-fold increase in kq in this mutant. The MD simulations on native and MopRG148P additionally indicated a reorientation of the allosteric linker region (Fig. S14). The relative orientation of the α7 as well as overall allosteric linker region with respect to the β6 (residues 174–180) reflected the dynamic nature of the allosteric linker region and the long-distance impact of G148P mutation. Thus, fluorescence studies highlight that both the phenol pocket and the allosteric linker region are dynamic in nature and show different accessibilities both in the apo versus bound form. The changes in the G-hinge region are more subtle in

### Table 3

| Protein         | τm (ns) ±0.07 | Ksv (M⁻¹) (KI) | kq (10⁹ M⁻¹ s⁻¹) (KI) |
|-----------------|---------------|----------------|-----------------------|
| MopR W199       | 3.24 ±0.07    | 4.36 ±0.03     | 1.34 ±0.06            |
| MopR W199 + phenol | 3.15 ±0.06    | 2.39 ±0.08     | 0.76 ±0.11            |
| MopR W156       | 3.59 ±0.05    | 4.60 ±0.08     | 1.27 ±0.06            |
| MopR W156 + phenol | 3.47 ±0.08    | 4.21 ±0.06     | 1.21 ±0.07            |
| MopR W134       | 2.70 ±0.02    | 6.21 ±0.09     | 2.30 ±0.05            |
| MopR W134 + phenol | 2.62 ±0.03    | 3.90 ±0.08     | 1.49 ±0.06            |
| MopR W199 G148P | 2.68 ±0.02    | 2.98 ±0.04     | 1.11 ±0.04            |
| MopR W199 G148P + phenol | 2.41 ±0.11    | 6.09 ±0.11     | 2.86 ±0.12            |
| MopR W156 G148P | 2.03 ±0.05    | 4.57 ±0.08     | 2.24 ±0.08            |
| MopR W156 G148P + phenol | 2.09 ±0.06    | 4.21 ±0.07     | 2.01 ±0.09            |
| MopR W134 G148P | 2.79 ±0.03    | 3.64 ±0.06     | 1.30 ±0.05            |
| MopR W134 G148P + phenol | 2.80 ±0.06    | 3.62 ±0.07     | 1.29 ±0.08            |

Bimolecular quenching constant (kq) values for the single tryptophan mutants estimated from mean lifetime (τm) and Stern–Volmer constant (Ksv).

**Figure 5. Flexibility of the allosteric linker.** Environment of W199 (shown as orange sticks) in (A) bound MopR AB and (B) bound MopR G148P. W199 is less accessible in native MopR AB as compared to MopR G148P.
nature, and mutations in this region do not affect local environment, rather distal regions show differences if the G-hinge is perturbed.

**Allosteric network and its implications to transcription activation**

Mapping of the hydrogen bonding propensities obtained via MD simulations on the MopR\textsuperscript{AB} (Fig. S1B) crystal structure helped create a communication network which connected the three strategic regions; phenol cap, G-hinge and allosteric linker. The electrostatic connections indicate that the G-hinge region likely communicates with the allosteric linker region via the phenol pocket cap (Fig. 6A). The reliability of the hydrogen bond network was tested by creating mutations that disrupt the existing network. To access the role of this network in regulating the activity of the tandemly located ATPase motor, that is to understand the effect of these mutation on ATP hydrolysis, the mutations were introduced on the longer version of the MopR protein, MopR\textsuperscript{A+C} (Fig. S1C), which comprised of both the phenol binding and ATP hydrolysis domain. All phenol affinities were measured only using the MopR\textsuperscript{AB} construct (Fig. S1C). It was observed that any perturbation of the network results in marginal decrease in phenol affinity but leads to complete loss of the downstream ATPase activity. For instance, mutation of H106 and W134, where the G-hinge network originates, shows that although the H106A and W134A mutants are able to bind phenol with moderate affinity, a complete loss in ATPase activity occurs (Fig. 6B). The observation highlights that it is not just the binding of phenol to the ligand pocket that determines downstream signaling, rather the contacts between the

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**Figure 6. Allosteric communication network.** A, overall allosteric communication network in MopR sensory domain mapped using hydrogen bond propensities. B, bar graph representing the dissociation constant ($K_d$) and percent stimulation of ATPase activity of various mutants upon phenol binding. C, whole cell biosensor design of MopR based on a luciferase expression chemiluminescent readout. D, whole cell biosensing assay depicting enhanced luciferase activity for the MopR\textsuperscript{G148P} mutant (MopR\textsuperscript{G148P}) as compared to the WT.
intermediary residues are crucial for correct passage of the binding event. A similar scenario was observed for the phenol pocket-allosteric linker network, here also any perturbation emanating from the phenol pocket, completely abrogates ATPase activity as was observed for the proteins where the S166, Y165, and Y176 network has been individually disrupted via mutations of these residues (Fig. 6B). It is noteworthy that in these mutants, the phenol affinity is only marginally affected but ATPases activity shows a dramatic loss (Figs. 6B, S15).

The G148P mutation is at the end of the G-hinge network, and here, we observed that phenol binding had increased by 7-fold with no significant effect on the ATPase activity. Hence, to understand the effect and relevance of the mutation and its implications toward downstream transcriptional activity of MopR, an in vivo transcriptional assay was performed for both the native and G148P variant. A whole-cell setup (Figs. 6C, S1D), where the entire upstream activation cassette, which includes the upstream activation site, full-length MopR gene under its σ70 promoter, and the downstream σ54 promoter was constitutively cloned in the pGL3basic vector (Fig. 6C) such that the luciferase expression could serve as a reporter of the transcription activity (described in detail in experimental procedures). In A. calcoaceticus, expression of MopR is essential for activation of the σ54-dependent RNAp, which then transcribes the phenol degradation cassette. Here, this gene cassette was replaced by luciferase to gauge the transcription ability of MopR G148P luciferase signal of MopR G148P LUC was always higher (25%) than the native MopR LUC, indicating better transcription ability of MopR G148P LUC (Fig. 6D). Thus, analyzing the results, it becomes apparent that the G-hinge mutant state is indeed effectively primed for downstream activation. It is not certain why the G-hinge mutant exhibits higher transcriptional activity or phenol binding. One reason we speculate is that G148P mutation facilitates the opening of the sensor domain and allows the phenol moiety to enter with more ease that subsequently expedites the downstream relay.

Discussion

Proteins elicit their function by shuffling between conformational states that allow for various important events such as binding of ligand to catalysis and in some instances facilitate transfer of information between active sites (34). In several of these systems, underlying allosteric regulation governs a switch between distinct states that allows for efficient function (35). Here, it is important to assert that allosteracy can operate both via inducing a major structural change, such as in conformational allostery, or in some instances, subtle changes in side chain of certain amino acids, without any apparent structural change can contribute to function, like seen in systems that exhibit dynamic allostery (9, 36, 37). A common observation in both types of allosteric systems is a shift in hydrogen bonding network, where perturbation has resulted in dramatic change in the functional properties. For example, systems where conformational allosteric is predominant, such as Hsp70, a complicated bidirectional hydrogen bonding network has been observed where allosteric cues between the nucleotide-binding domain and the substrate-binding domain are passed via rearrangement of this crucial connection (38).

In this work, by employing MopR as a model system, we explore the importance of allosteric networks that connect different segments of a protein domain and decipher how these connections, which sometimes are elusive and seemingly silent, can cause changes in the protein network that have a profound effect on both affinity and activity. Two distinct networks, one that is primarily governed by dynamic allosteric, the phenol pocket-G-hinge network, and the other which exhibits aspects of both dynamic and conformational allosteric, namely the phenol pocket- allosteric linker network, was unearthed.

The phenol pocket-G-hinge network role was deciphered by examining the crystal structure of the phenol-bound form and comparing it with the MD trajectory of the apo form. The X-ray structure shows that binding of phenol results in a compact structure where it binds to the interior of the protein, in a snug pocket. Further, MD simulations of the apo form shows that the structure in the absence of phenol is more or less similar and only few notable conformational changes were observed. However, a noteworthy feature was a distinct shift in the electrostatic network that respectively connects the G-hinge and the allosteric linker regions with the phenol pocket. For instance, it was observed that phenol binding induces a shift in the hydrogen bonding propensities of W134 and H106; this effect percolates to the next residue in the relay and finally the network that terminates at the G-hinge, which is different from the apo state. What was most fascinating is that an analysis of the residues that partake in the relay network reveals that the residues are more or less conserved in close homologs of MopR such as PoxR, DmpR, XylR, etc (Fig. 7). It was observed that the side chain residues that form an integral part of the network were completely conserved, whereas residues that contribute through their backbone show some variation. This highlights the fact that the network is more like a community network, which has co-evolved in these members such as to allow a more orchestrated shift between distinct allosteric states that are functionally relevant. Perturbation of this network by introducing a G148P mutation experimentally confirmed that indeed the phenol pocket is influenced by changes in the G-hinge region. Here, we think that the G-hinge network most likely assists in entry of phenol, as MD simulations hint that synchronous fluctuations between the G-hinge region and the pocket cap may facilitate phenol binding. In the apo state, the protein likely accesses multiple conformational states, and it is possible that the state conducive for phenol entry, where coordinated motion between the pocket cap and the G-hinge occurs, allows for effective phenol access. The observation that the G148P mutation affects phenol affinity and corroborating fluorescence studies that show that
the phenol pocket is partially preformed in G148P, confirms that perturbation in the G-hinge is noticed at the phenol-binding site. These observations reassert that G-hinge and phenol pocket region are strongly connected. However, introduction of G148P does not cause any perturbation in the transmission network, as relay is not disrupted. Supporting transcription activity studies for this mutant further corroborate that this mutant not only aids in phenol binding but also exhibits an enhanced overall transcriptional rate of around 25%. Thus, it appears that by introducing the G148P mutation, we have shifted the conformational landscape of MopR, and it predominantly now accesses states that are more conducive to allow for quick passage of signal that favor downstream function.

The network established for MopR protein is primarily governed by rearrangement of the electrostatic interactions and only subtle structural changes are observed; therefore, it appears that dynamic allostery is predominantly at play. Comparative analysis of thermodynamic parameters of MopR in both the native and G148P mutant shows that both the enthalpic and entropic contributions influence binding. It was earlier envisioned that in systems that exhibit dynamic allostery, such as catabolite-activating protein (CAP) and PDZ domains, entropy-driven allostery solely governs conformational fluctuation (7, 39). However, recent reports by Kumawat et al. clearly show that dynamic allostery can encompass enthalpic effects (9, 40). In their work, they show that restructuring of the electrostatic network in the PDZ protein has an enthalpic contribution. Deletion of a helix that lies at the periphery of the PDZ domain does not perturb the overall structure but it results in 21-fold reduction in affinity of the ligand (39). This is because this helix assists in a rearrangement of the electrostatic network upon ligand binding. On a similar vein, in MopR, introducing a G148P mutation likely alters the overall phenol-binding dynamics of MopRAB, resulting in a 7-fold increase in affinity for this mutant. Here, also both entropic and enthalpic contributions were found to be key players that determine the shift. Thus, dynamic allostery in the MopR system governs the communication between strategically located hotspots in this protein. Such type of changes in affinity via the dynamic allosteric pathway are also observed in cold adaptation of enzymes, as observed in adenylate kinase, where specific distal surface mutation that increase the propensity of the enzyme to unfold result in marked change in ligand affinity (41).

Apart from the phenol pocket-G-hinge network, the other network that also originates at the phenol pocket and terminates near the end of the ligand-binding domain, near the B-linker region also showed interesting features. This phenol pocket-allosteric hinge communication network is an example of a complex interplay of conformational and dynamic allostery. The B-linker helix at the end of the phenol-binding domain connects this domain with the tandemly located ATPase domain, and it is envisioned that entry of phenol stimulates ATPase activity of the AAA+ domain via conformational changes in the B-linker that are passed on (20, 21). Therefore, dynamic nature of the B-linker as corroborated by fluorescence and X-ray crystallographic studies was expected. What is more interesting is the presence of dynamic allostery that is facilitated via a conserved network of connections that run between the phenol pocket and tip of the B-linker that helps communicate the progress of phenol binding. Comparison of the apo and bound forms of ligand-binding domain shows that a rewiring of the electrostatic connections between the phenol pocket and the B-linker in the presence of phenol enables passage of information. Thus, the ATPase domain is informed of phenol binding via dynamic allostery within the internal network that spans within the phenol-binding domain.
and expressed as conformational allostery via structural motion in the B-linker. Both, ITC binding studies, and ATP hydrolysis studies with variants of MopR, where this network has been disrupted show the importance of conserved interactions for effective relay. Almost complete loss of ATP stimulation is observed whenever the network is disrupted. In some of these cases, such as mutations at the periphery of the phenol pocket, as in W134A and H106A, we observe that these point mutations are still capable of binding phenol, although with somewhat decreased affinity. However, the ATP stimulation is completely obliterated due to these mutations. Other mutations in the relay that are not directly involved in phenol binding leave the phenol binding almost unaltered but again complete loss in ATPase activity occurs. This highlights the importance and precision with which the relay network has to operate to elicit function and asserts the idea that phenol binding is not the rate determining step that controls MopR function, rather it is the integrity of the relay that is paramount to maintain function. The importance of this network is again highlighted as it was observed that like in the case of the G-hinge network, the phenol pocket-allosteric hinge relay network is also conserved in other proteins such as PoxR, XylR, and DmpR that bind similar aromatic compounds.

The MD simulations indicated independent relay networks emanating from the phenol pocket to the G-hinge and allosteric linker. However, a striking observation that emerged from the crystal structure of the ligand-binding domain of G148P mutant was a rearrangement of the B-linker. This was unexpected as MD analysis does not reveal any direct connection of the G-hinge with the allosteric linker region. Thus, this experimental observation seeded the idea that the phenol pocket, G-hinge, and allosteric linker are all connected and the networks operate in an interdependent fashion. It can be envisioned that the networks run via the central phenol pocket and are perceived by other functionally important parts of the protein. ITC and transcriptional activity assay also confirm this idea; otherwise, a mutation in the G-hinge would not have such far reaching functional effects. The study emphasize that it is not only the local active site region that partakes in function rather seemingly distal and benign parts of protein domains have a profound influence on functional and binding properties of proteins. It reasserts that apart from the obvious active site residues, several other portions of the protein also play an integral role in determining the functional outcome of an enzyme.

In conclusion, here we show via a combination of MD and experimental approaches that MopR has distinct regions that form a highly involved bidirectional network. The central phenol pocket within the ligand-binding domain connects to two other regions G-hinge, which likely influence entry of phenol, and allosteric hinge, which passes the signal to the downstream ATPase domain. Phenol binding induces a rewiring of the electrostatic connections that brings about function by eliciting dynamic allostery. It was established that it is not the ligand binding that determines downstream ATP hydrolysis, rather the order in which the rearrangement of interactions that occur upon ligand binding is more crucial in maintaining the integrity of the relay. Further the flexibility of the G-hinge was probed by undertaking a G148P mutation, and it was determined that G-hinge influences both phenol binding as well as downstream transcription. Changes in this region are communicated via the bidirectional network that spans across the ligand-binding domain. The work highlights the importance of long-distance communication and shows how relay networks in proteins are highly sensitive to disruption and an important backbone that support protein function.

**Experimental procedures**

**MD simulations**

The computational structural model for the phenol-bound WT (Bound MopR<sup>AB</sup>) was the crystal structure of a dimer of MopR sensor domain (PDB ID: 5KBE) (Fig. 1B, S1B). For apo simulations, the coordinates of phenol ligand were removed from this crystal structure (Apo MopR<sup>AB</sup>). For MD simulations of the G148P mutant (MopR<sup>G148P</sup>), the apo and bound states were modeled by *in silico* substitution of glycine 148 to proline in apo and bound MopR<sup>AB</sup>, respectively. All the crystal water molecules were removed. The missing crystal residues were modeled using CHARMM-GUI (42). The N and C terminus of the protein were capped by NH<sup>3+</sup> and COO<sup>-</sup>. The protonation state of all amino acid residues corresponds to neutral pH except zinc coordinated cysteines (C155, C181, C189), which were modeled in their deprotonated states (43). The proteins, the ligand (phenol), and the ions were modeled using Charmm36m forcefield (44). The systems were solvated with using the TIP3P water model (45) where for each of the system state, the starting models were placed at center of truncated octahedron box with empty box volume filled by TIP3P-charmm water (~18,500 molecules) and neutralized by 200 mM NaCl ions. The system was energy minimized using steepest descent algorithm with position restraint on all protein heavy atoms. After minimization, 500 ps position restrained MD simulation is performed in NVT ensemble at 298 K temperature maintained by Nose–Hoover thermostat (46, 47) with relaxation time of 1 ps. The system was subsequently subjected to unrestrained MD simulation for a duration of 1 μs in NPT ensemble at 298 K temperature (using Nose–Hoover thermostat) and 1 bar pressure using Parrinello–Rahman barostat (48) with a coupling constant of 5 ps. All simulations were performed with 2 fs time step using Leap-frog integrator with-in GROMACS 20XX simulation package. Periodic boundary condition was implemented in all three dimensions. The Verlet cut-off scheme (49) was employed for Lennard Jones interaction and short-range electrostatic interactions. Long range electrostatic interactions were treated by Particle Mesh Ewald summation method (50). All hydrogen bonds were constrained using LINCS algorithm (51). The bonds and angles of TIP3P water molecules were constrained using SETTLE algorithm (52). The simulations with apo protein were replicated in three independent trajectories. The final 500 ns of each of the 1 μs long trajectories were considered for atomistic analysis.
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**MD simulations analysis**

For each of the system state, hydrogen bond profiles were analyzed using a criterion of distance ≤ 3.5 Å and angle ≤ 30° (53, 54). The propensity of hydrogen bond formation is calculated for individual hydrogen bonds as time average over all the sampling for each system. The propensity of hydrogen bond formation is basically the probability of finding a hydrogen bond between two residues and ranges between 0 and 1. The hydrogen bond propensities showing difference ≥0.5 were considered significant. The significant difference is considered if it follows either or all of the criteria. The first criteria considered was if hydrogen bond propensity showing difference ≥0.5. The second criteria was if a residue has decreased hydrogen bond propensity with one residue and simultaneous increase in hydrogen bond propensity with other residue as system go between different system states, with total difference ≥0.5. The third criteria being a residue has decreased/increased hydrogen bond propensity with multiple residues with total difference ≥0.5.

π-π aromatic stacking was considered for residues tyrosine, tryptophan, phenylalanine, and histidine (55) if the stacking pairs are within 7 Å distance (54). In order to calculate the pocket volumes, MD pocket utility is used (56). The pairwise distance between α2 (residues 48–62) and α5 (residues 138–147), β6 and α7/allosteric linker region were determined by taking the average distance of all possible pairs of main chain atoms.

**Bioinformatic analysis**

For mapping the sequence conservation, sequences of NtrC family proteins that are specific to the aromatic hydrocarbons were chosen. Details of the proteins are given in Table S1. The sequence alignment was performed using ClustalW (57). The sequence logo was generated using WebLogo (58). The sequence conservation was estimated as bits score on y-axis and residue number on x-axis. A value of ~4.3 corresponds to strict identity of the amino acid residue (58).

**Site-directed mutagenesis and protein purification**

The recombinant pET vector construct of MopR<sub>AB</sub> (Fig. S1B) was used as template (14) to make the glycine to proline (G148P) hinge mutant, MopR<sub>G148P</sub>. For fluorescence spectroscopy studies, the following mutations were made to generate single tryptophan constructs using MopR<sub>AB</sub> (Fig. S1B) as the starting template: W134A_W199F_W37F (MopR<sub>W156G</sub>), W37F_W156F_W134A (MopR<sub>W199</sub>), W37F_W199F_W156F (MopR<sub>W134G</sub>), W134A_W199F_W37F_G148P (MopR<sub>W156G</sub> G148P), W37F_W156F_W134A_G148P (MopR<sub>W199G</sub> G148P), W37F_W199F_W156F_G148P (MopR<sub>W134G</sub> G148P) (Table 2). For mutants Y165L, Y176L, and S166G, the sensor domain constructs were generated using MopR<sub>AB</sub> (Fig. S1B), as template and construct MopR<sub>Ac</sub> (Fig. S1C) constituting of both sensor and ATPase domain using a template that encodes 1 to 500 residues of the mopR gene.

All the mutants were made by employing standard site-directed mutagenesis protocol using the Phusion DNA polymerase from New England Biolabs. The mutant expression constructs were subsequently transformed into *Escherichia coli* BL21(DE3) plysS cells, overexpressed with 1 mM IPTG as six-his tag fusion proteins and cultured at 16 °C for 16 h. All the mutated proteins were purified using Ni-NTA resin by standard His-tagged affinity purification protocol. The composition of the buffers used in subsequent purification steps was as follows: lysis buffer (50 mM Tris–HCl buffer, pH 7.5; 2 mM imidazole; 200 mM NaCl), wash buffer (50 mM Tris–HCl buffer, pH 7.5; 30 mM imidazole; 200 mM NaCl), and elution buffer (50 mM Tris–HCl buffer, pH 7.5; 350 mM imidazole; 100 mM NaCl). The eluted fractions were desalted using an Econo-Pac 10DG (Bio-Rad) column that was pre-equilibrated with a desalting buffer containing 25 mM Tris–HCl buffer, pH 7.5; 80 mM NaCl, 5% glycerol, and 0.5 mM DTT. The desalted protein fractions were pooled and concentrated up to 5 to 8 mg/ml. The fractions were then flash-frozen in liquid N2 and stored at ~80 °C until they were used. The purity of the protein was verified by running a 10% SDS-PAGE followed by Coomassie blue (HiMedia) staining.

**Ligand-binding experiments using ITC**

All the ITC experiments were carried out on the constructs of MopR<sub>AB</sub> constituting of residues 1 to 229 (Fig. S1B). All the protein and ligand samples were prepared in a buffer that contained 25 mM HEPES (pH 7.5) and 80 mM NaCl. In the ITC experiment, phenol was titrated against buffer and subtracted from the raw data prior to model fitting, in order to nullify the heat of dilution. About 40 μM of MopR<sub>AB</sub> and 10 μM of MopR<sub>G148P</sub> was titrated with 400 μM and 200 μM of phenol, respectively. For the Y165L, S166G, and Y176L mutants, 25 μM of protein and 400 μM of phenol were used. The value of K<sub>d</sub> for W134A and H106A were taken from the previously reported results (14). The volume of the titrant (ligand) added at each injection into the sample cell was 2 μl for 5s. A set of 20 injections were performed for each experiment with an interval of 120 s between each successive injection. The temperature was maintained at 25 °C. The stirring rate was kept constant at 1000 rpm throughout all the ITC experiments. The data obtained were fitted and analyzed with Origin 2019b software (OriginLab) using ‘one set of site model’ (59). The curve fitting was done in the acceptable experimental window of c values of 10 ≤ c ≤ 500, where c-value = n[Protein]/K<sub>d</sub> for n noninteracting identical sites (K<sub>d</sub> is the dissociation constant) (60).

**Size-exclusion chromatography**

A Superdex 75 10/300 GL column installed on an NGC Chromatographic System (BioRad) and equilibrated with the sample buffer (25 mM Tris–HCl [pH 7.5], 80 mM NaCl, 5% glycerol, and 0.5 mM DTT). About 100 μl of ~4 mg/ml MopR<sub>AB</sub> and ~2 mg/ml of MopR<sub>G148P</sub> in sample buffer were injected onto equilibrated column at 4 °C at a flow rate of 0.3 ml/min. The column was calibrated with the following
standards, carbonic anhydrase (29 kDa), Ovalbumin (44 kDa), and Conalbumin (75 kDa).

**Crystallization of MopR<sup>G148P</sup>**

The purified his-tagged MopR<sup>G148P</sup> (10 mg/ml) was first screened for crystallization using several commercially available crystallization screens such as Crystal screen, PEG/Ion (Hampton Research) and JCSG suite, PACT suite (Qiagen) using a Crystal Phoenix crystallization robot, and the sitting-drop vapor diffusion technique at the crystallization facility at IIT Bombay; however, no notable crystal hits were found in any of the screens. Attempts were then made to cocrystallize MopR<sup>G148P</sup> with phenol. MopR<sup>G148P</sup> was incubated at 4 °C for around 30 min with 5 mM of phenol and was then subjected to crystallization trials in a manner like that of apo protein, using the screens that are commercially available at Hampton and Qiagen. All the protein–ligand complex solutions were filtered before crystallization trials. Crystals were obtained within around 30 min with 5 mM of phenol and was then subjected to crystallization trials in a manner like that of apo protein, using the screens that are commercially available at Hampton and Qiagen. All the protein–ligand complex solutions were filtered before crystallization trials. Crystals were obtained within 7 days of setup in the following condition, 0.2 M magnesium acetate.4H<sub>2</sub>O, 15% w/v PEG 3350. The crystals were further optimized at 20 °C using the hanging-drop vapor-diffusion method, with 1 μl of a protein solution, 1 μl of a precipitant solution, and a 500 μl reservoir volume. The trays were monitored in a temperature-controlled cabinet. Diffraction quality isolated crystals of the maximum size grew (100 × 70 × 70 μm) over a period of 10 to 12 days. Under optimized conditions, MopR<sup>G148P</sup> in complex with phenol crystallized in the C-centered orthorhombic space group C222<sub>1</sub>, with unit cell dimensions of a = 62.20, b = 118.30, c = 56.70, and α=β=γ = 90°.

**Data collection and processing**

X-ray diffraction experiments were performed at the home source of Indian Institute of Technology (IIT) Bombay using a Rigaku MicroMax-007HF X-ray diffractometer. A single crystal of the ligand complex was cryo-protected with 20% (v/v) ethylene glycol (prepared using mother liquor) prior to data collection. The crystal was then flash cooled in liquid nitrogen and transferred to a stream of nitrogen gas at 100K. X-ray data were collected at a wavelength of 1.5418 Å on a Rigaku R-Axis IV++ detector. The dataset was indexed, integrated, and transferred to a stream of nitrogen gas at 100K. X-ray data collection and processing

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X-ray crystal structure solution

The structure of MopR<sup>G148P</sup> cocomplexed with phenol was determined at 2.3 Å by the MR method using the MR module of the PHASER program (62) and native monomeric unit of MopR<sup>AB</sup>–phenol complex structure as a search model (PDB ID: 5KBE). Manual model building of the partially refined structures was carried out using the graphics program COOT (63) and they were further refined using REFMAC5 (64). All figures were made in PyMOL (65).

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**CD studies**

The CD spectra of the MopR<sup>AB</sup>, MopR<sup>G148P</sup>, and single tryptophan mutants were recorded on a Jasco J-815 CD spectrometer. The protein concentration used was 0.5 mg/ml and phenol used was 0.2 mM, respectively. All the protein and ligand samples were prepared in phosphate buffer (25 mM sodium phosphate pH 7.5, 80 mM NaCl). Using same method, the CD spectra were recorded for all single tryptophan mutants (Data shown in Fig. S9). Scans were performed at 20 °C using 0.1 cm path length quartz cuvettes with 8 s differential integration time at a scan rate of 50 nm/sec. The mean residual ellipticity (MRE) in units of deg.cm<sup>2</sup>.dmol<sup>-1</sup> was determined using the formula (66),

\[
\text{MRE} = \frac{(\text{MRW} \cdot \theta)}{10(d \cdot c)}
\]

where θ is the observed ellipticity (degrees), d is the path length (cm), and c is the concentration (in units of g/ml). The MRW (mean residue weight) is obtained by dividing the molecular mass by N - 1, where N is the number of amino acids. The CD-based thermal denaturation experiment was performed for MopR<sup>AB</sup> and MopR<sup>G148P</sup> to determine the stability of these proteins. Scans were performed at a temperature range of 20 °C to 95 °C using 0.1 cm path length quartz cuvettes with 16 s differential integration time at a scan rate of 100 nm/sec with 3 min delay time per temperature change.

**Steady state fluorescence measurements**

Absolute fluorescence intensities were measured on FluoroMax-4 spectrofluorometer with 4 μM of protein samples in buffer (25 mM HEPES, 80 mM NaCl). The quenching experiments were performed on a Varian Cary Eclipse spectrofluorometer with 12 μM of protein in buffer (25 mM HEPES, 80 mM NaCl) titrated with increasing concentration (0–300 mM) of potassium iodide (KI). For the phenol-bound form, the mutants were incubated with 5 mM phenol for 30 min prior to study. All the experiments were performed in quartz cuvette of 1 cm path length and the samples were excited at 295 nm and the emission spectra were recorded for 300 to 450 nm wavelength range. All measurements were carried out in triplicates and mean ± standard error have been reported.

**Time-resolved fluorescence measurements**

The time-resolved fluorescence decay was recorded using Ti-sapphire laser (Mai Tai HP, Spectra Physics) pumped by an Nd: YVO4 laser (Millennia X, Spectra Physics) generating the 885 nm pulses of width ~1 ps. A flexible second- and third-harmonic generator (GWU, Spectra Physics) was used to obtain the frequency-tripled laser of 295 m for excitation. Fluorescence emission was collected through a 305 nm cut-off filter to exclude scattered photons completely when the monochromator was set at 335 nm. To obtain fluorescence lifetimes, a polarizer oriented at the magic angle (54.7°) was used to eliminate anisotropy decay artifact in the fluorescence
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decay data. All the measurements were made in 1 cm path length with 10 μM concentration of protein. All measurements were carried out in triplicates and mean ± standard error have been reported.

Fluorescence data analysis

The obtained decay curves collected at the magic angle were deconvoluted with the instrument response factor by using nonlinear least-square iterative deconvolution method based on the Levenberg–Marquardt algorithm and expressed as a sum of exponentials with equation,

\[ I(t) = \sum a_i \exp \left( -t/\tau_i \right) \]  

(2)

where, \( I(t) \) is the fluorescence intensity at time \( t \) with \( a_i \) being the amplitude of the \( i^{th} \) lifetime \( \tau_i \) such that \( \sum a_i = 1 \). The average fluorescence yield was estimated by calculating the mean lifetime using the equation: \( \tau_m = \sum a_i \tau_i \).

The goodness of fitting was assessed from the reduced chi square (\( \chi^2 \)) values as well as from randomness of the residuals.

For monitoring the solvent accessibility of the four tryptophans, the KI quenching experiments were done and fitted to Stern–Volmer equation (67):

\[ \frac{F_0}{F} = 1 + k_q \tau_m |Q| \]  

(3)

where, \( F_0 \) and \( F \) are the fluorescence intensity in the absence and presence of quencher KI, respectively, \( |Q| \) is the concentration of quencher and \( k_q \) is the bimolecular quenching rate constant (M\(^{-1}\) s\(^{-1}\)). \( \tau_m \) is the mean lifetime in the absence of Q. \( K_w \) is the Stern Volmer constant given by \( k_q \tau_m \) and is obtained from slope of Stern Volmer plots.

Colorimetric ATPase assay

The Malachite green assay was used for determining the ATPase activities. The assay was carried out on the constructs of MopRA\(^{AC} \) constituting of residues 1 to 500 (Fig. S1C). About 2 μM of protein sample was incubated with 1 μM phenol. About 2 mM of ATP was then added to the reaction mixture, and the reaction was allowed to proceed for 15 min followed by quenching with 0.5 M EDTA. The dye reagent (consisting of Malachite green, ammonium heptamolybdate and Tween20) was added to the reaction mixture to form a phosphomolybdate complex, which gave a green color and for which absorbance was measured at 630 nm. All the data were collected in triplicates to estimate the errors.

Construction of the whole cell MopR\(_{LUC}\) and MopR\(_{G148P,LUC}\) construct

The whole cell construct of native MopR was first designed (MopR\(_{LUC}\)), which consists of the full-length mopR gene under the control of \( P_m' \) promoter with a luc reporter module attached upstream, cloned into a pGL3basic expression vector that has been purchased from Promega (Fig. S1D). \( P_m \) is the \( \alpha^{73} \)-based promoter that controls the mopR gene transcription activity, and \( P_{mop} \) is the \( \sigma^{34} \)-based promoter, which triggers downstream catabolic pathways on activation of MopR with suitable pollutants. For cloning experiment, firstly, the purified genomic DNA of \( A.\) calcoaceticus NCIB8250 (2 μg/μl) (16) was used as a template for the PCR amplification of mopR - \( P_m/P_{mop} \) gene sequence. PCR was performed by using forward primer 5’-ACCGAGGTACCATTTAAGCCCGATAATTTA A-3’ and a reverse primer, 5’-ATTGACTCGAGATTCCGCTC ACCAGTAATAC-3’, with XhoI and KpnI sites at the ends. The amplified gene was then cloned into the pGL3basic vector using standard cloning techniques (Fig. 6C). This native whole cell clone was further used as a template to make the G148P mutant (MopR\(_{G148P,LUC}\)) in mopR - \( P_m/P_{mop} \) by employing standard ‘site-directed mutagenesis’ protocol using the “site-directed mutagenesis kit” from Kapa biosystems. The cloned MopR\(_{G148P,LUC}\) construct was then transformed into \( E.\) coli DH5α cells and preinoculum was setup at 37 °C overnight with the transformed colonies for performing the luciferase assay.

Luciferase assay design on MopR\(_{LUC}\) and MopR\(_{G148P,LUC}\)

The luciferase assay described in this work is based on the luciferase assay system kit protocol from Promega, which was performed to monitor the downstream transcription activation potential of the native and mutated MopR (68). The overnight grown cultures of \( E.\) coli DH5α cells harboring the recombinant MopR\(_{LUC}\) and MopR\(_{G148P,LUC}\) plasmids were subcultured and grown to the log phase till the optical density at 600 nm reached ~0.3. This was followed by supplementation with phenol at a gradient concentration of 0.1 to 1000 μM. After 1 h of induction, 50 μl of the cells were withdrawn and frozen at −70 °C with the addition of 5 μl of 1 M KH₂PO₄ and 20 mM EDTA (pH 7.8). The cells were lysed by the addition of 150 μl of lysis solution (1.25 mg/ml lysozyme, 2.5 mg/ml bovine serum albumin, 1X CCLR) (Promega) and incubating at room temperature for 10 min. Supernatants were obtained by centrifugation. For the luciferase activity, 20 μl of supernatant was mixed with 50 μl of firefly luciferin solution (Promega). The bioluminescence was measured for 30 s by Luminometer Berthold detection system. Induction of the MopR\(_{LUC}\) and MopR\(_{G148P,LUC}\) by phenol was expressed as normalized luminescence (NL) calculated as follows:

\[ NL = \frac{S_{L} - S_{B} \times CF}{S_{L} - S_{B}} \]  

(4)

where \( S_{L} \) is the luminescence of the biosensor in the dilution of the aromatic pollutant, \( S_{B} \) is the background luminescence of the sensor bacteria, and CF is the correction factor.

\[ CF = \frac{L_{B}}{L_{L}} \]

where \( L_{B} \) is the background luminescence of the control bacteria, and \( L_{L} \) is the luminescence of the control bacteria in the presence of an aromatic compound. The detection limit for the sensor bacteria for different pollutants was defined as a
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Abbreviations—The abbreviations used are: DmpR, dimethylphenol regulator; ITC, isothermal titration calorimetry; MD, molecular dynamics; MR, molecular replacement; PDB, protein data bank; RNAP, RNA polymerase.

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