Forcing the Issue: Aromatic Tuning Facilitates Stimulus-Independent Modulation of a Two-Component Signaling Circuit

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Supporting Information

ABSTRACT: Two-component signaling circuits allow bacteria to detect and respond to external stimuli. Unfortunately, the input stimulus remains unidentified for the majority of these circuits. Therefore, development of a synthetic method for stimulus-independent modulation of these circuits is highly desirable because particular physiological or developmental processes could be controlled for biotechnological purposes without the need to identify the stimulus itself. Here, we demonstrate that aromatic tuning, i.e., repositioning the aromatic residues commonly found at the cytoplasmic end of the receptor (EnvZ) transmembrane domain, facilitates stimulus-independent modulation of signal output from the EnvZ/OmpR osmosensing circuit of Escherichia coli. We found that these osmosensing circuits retained the ability to respond appropriately to increased external osmolarity, suggesting that the tuned receptors were not locked in a single conformation. We also noted that circuits containing aromatically tuned variants became more sensitive to changes in the receptor concentration than their wild-type counterpart, suggesting a new way to study mechanisms underpinning receptor concentration-dependent robustness. We believe that aromatic tuning has several advantages compared to previous methods aimed at stimulus-independent modulation of receptors and that it will be generally applicable to a wide-range of two-component circuits.

KEYWORDS: aromatic tuning, two-component circuit engineering, stimulus-independent modulation, concentration-dependent robustness

Two-component circuits are the most prevalent mechanism by which bacteria sense, respond, and adapt to external stimuli. These systems mediate responses to a wide range of environmental conditions such as nutrient availability, ambient temperature, or external osmolarity. They also facilitate multiorganism phenomena such as quorum sensing, biofilm formation, and host–pathogen interaction. In addition, they control essential environmental and agricultural processes such as chloroplast synthesis and root nodule formation.

Therefore, development of a synthetic method for stimulus-independent modulation of these circuits is highly desirable because particular physiological or developmental processes could be controlled and characterized for biotechnological purposes without the need to identify the stimulus itself.

A canonical circuit consists of a membrane-spanning sensor histidine kinase (SHK) and a cytoplasmic response regulator (RR). The largest group of SHKs possesses a periplasmic or extracellular domain responsible for stimulus perception. Subsequent signal transmission to the cell interior occurs via the adjacent transmembrane domain. Within the cytoplasm, most SHKs participate in both the phosphorylation (kinase activity) and dephosphorylation (phosphatase activity) of their cognate RR. For bifunctional SHKs, the extent of input stimulus controls the ratio of these activities, thereby governing the intracellular level of phosphorylated RR.

Phosphorylation of the RR modulates the activity of the covalently attached output domain, which usually interacts with DNA to control transcription of genes appropriate for mediating a response to the perceived stimulus.

A vast amount of genetic, biochemical, and structural information has been recently integrated into a “regulated unfolding” model of intraprotein signaling by modular proteins, including SHKs. This model proposes that modular proteins are composed of individual folding domains that contribute distinct functionalities. In the case of SHKs, it was suggested that the effector domain is maintained in an inactive conformation by a rigid connection between the stimulus perceiving and effector domains. Upon perception of stimulus, this structurally labile connection is disengaged, which, in turn, allows the effector domain to adopt an active conformation.

Therefore, the regulated unfolding model suggests that the transmembrane (TM)–HAMP junction would be a suitable region to target with site-directed mutagenesis with the aim of destabilizing the coupling between the periplasmic stimulus-perceiving domain and any downstream signaling domains (Figure 1). Other results more explicitly support targeting this...
phenotype was governed by the aromatically tuned SHK. If a particular SHK, then this would suggest that the desired phospho-OmpR regulates the transcription of a number of other membrane-spanning receptors.\(^{12,16}\) This allows intracellular levels of OmpR-P to be estimated by calculating the CFP/YFP ratio. Aromatic tuning of individual SHKs

**Figure 1.** Synthetically tuning signal output from SHKs. In a canonical SHK, stimulus is perceived by the periplasmic domain (peri) and transmitted through the transmembrane (TM) and HAMP domains to the catalytic ATPase (CA) and dimerization/histidylphosphotransfer (DHp) domains. One potential example of employing aromatic tuning, i.e., repositioning the aromatic residues commonly found at the TM–HAMP junction, would be to assign downstream phenotypes to particular SHKs. Within an organism of interest, each SHK could be individually subjected to aromatic tuning (red boxes) and subsequently monitored for the phenotype of interest. If the appearance of the phenotype (filled box) correlated with aromatic tuning of a particular SHK, then this would suggest that the desired phenotype was governed by the aromatically tuned SHK.

**Figure 2.** The EnvZ/OmpR osmosensing circuit of *E. coli* was subjected to aromatic tuning. The phosphorylated and unphosphorylated forms of EnvZ and OmpR are in equilibrium. EnvZ is a bifunctional SHK that phosphorylates and dephosphorylates its cognate RR, OmpR. Osmotic pressure (Osm), due to the presence of small inner-membrane-impermeable solutes, alters the ratio of these activities resulting in a net increase of intracellular OmpR-P. In this study, osmotic pressure (red) was induced by growing cells in the absence of stimulus identified. The easily controllable nature of the input stimulus and the well-characterized transcriptional output makes the EnvZ/OmpR osmosensing circuit an ideal choice for examining aromatic tuning within an SHK.

Aromatic tuning of EnvZ demonstrated that repositioning the Trp-178/Lue-179/Phe-180 triplet located at the TM–HAMP junction was sufficient to modulate signal output. We found that these tuned osmosensing circuits retained the ability to respond appropriately to additional external osmolarity, which demonstrates that the tuned EnvZ receptors possess altered steady-state signal output but were not locked in a single conformation. We also noted that osmosensing circuits containing aromatically tuned receptors became more sensitive to changes in EnvZ levels than their wild-type counterpart, pointing to a new way of studying the mechanisms underpinning receptor concentration-dependent robustness within two-component circuits. We conclude by discussing the general applicability of aromatic tuning to a wide-range of two-component circuits and the advantages of this strategy compared to those previously aimed at stimulus-independent modulation of signal output. This is highly desirable because particular biological processes could be controlled in the absence of stimulus identification.

**RESULTS AND DISCUSSION**

**Measurement of Steady-State Signal Output from the EnvZ/OmpR Osmosensing Circuit.** To analyze steady-state signal output from EnvZ/OmpR osmosensing circuits containing aromatically tuned receptors, the two-color fluorescent reporter strain MDG147\(^{-24}\) was used. MDG147 is a derivative of strain K-12 MG1655 that possesses transcriptional fusions of *cfp* to *ompC* and of *yfp* to *ompF* within its chromosome (Figure 2B). Quantifying the ratio of CFP to YFP fluorescence provides a rapid and sensitive measure of the ratio of *ompC* to *ompF* transcription, which estimates the intracellular level of phosphorylated OmpR. MDG147 cells harboring the control vector pEB5\(^{-25}\) were grown in glucose minimal medium containing increasing amounts of sucrose to increase signal output from the EnvZ/OmpR osmosensing circuit. As previously reported, MDG147 cells exhibited an increase in *ompC* transcription, as indicated by increased CFP fluorescence, and a decrease in *ompF* transcription, shown by decreased YFP fluorescence (Figure S1A)\(^{-24}\). These results confirm that the ratio of CFP to YFP fluorescence (CFP/YFP) can be used to estimate the intracellular phospho-OmpR levels (Figure S1B).

Strain EPB30\(^{-26}\) is an *envZ*\(^{-}\) derivative of MDG147 that is suitable to assess the effects of plasmid-based *envZ* expression. EPB30 cells were complemented with plasmid pEnvZ\(^{-27}\) or pRD400, a derivative expressing a V5-epitope tagged version of EnvZ. pRD400 maintains the IPTG-based induction of pEnvZ while adding a previously used heptadepsipeptide linker of Gly-Gly-Ser-Ser-Ala-Ala-Gly and the V5 epitope tag to the C-terminus of EnvZ.\(^{12,15,28-30}\) The wild-type and epitope-tagged versions of EnvZ were induced by addition of a wide range of IPTG concentrations, and the steady-state signal output of the various
osmosensing circuits was analyzed. Comparisons of CFP fluorescence, YFP fluorescence, or the CFP/YFP ratio of plasmid-complemented EPB30 cells grown under the low (0% sucrose) or high (15% sucrose) osmolarity regimes demonstrated that an intermediate range of IPTG concentrations was required to maintain steady-state signal output when either wild-type or the epitope-tagged version of EnvZ was present (Figure S2). Under either the low and high osmolarity regimes, steady-state signal output, as defined by CFP/YFP in EPB30/pRD400 cells, was similar to that of MDG147/pEBS cells when EnvZ-V5 was induced by addition of between roughly 10 and 50 μM IPTG (Figure S2).

Immunoblotting against the V5 epitope was performed to gain a quantitative understanding of the composition of osmosensing circuits containing EnvZ-V5 (Figure S3). When grown under either the low or high osmolarity regimes, osmosensing circuits within EPB30/pRD400 cells could tolerate a roughly 10-fold range in EnvZ-V5 levels while retaining steady-state signal output similar to MDG147/pEBS cells (Figure 3). It is important to note that EnvZ-V5 levels outside this range resulted in changes of CFP fluorescence but not YFP fluorescence, as previously reported (Figure S4).

**Figure 3.** Steady-state signal output from osmosensing circuits possessing increasing amounts of EnvZ-V5. Under the low (A) or high (B) osmolarity regimes, osmosensing circuits in EPB30/pRD400 (open circles) cells possess steady-state signal output similar to that of MDG147/pEBS over a broad range of receptor levels. The receptor level was determined by comparison to a control band within each lane on an immunoblot (see Figure S3). Error bars represent standard deviation of the mean with a sample size of n ≥ 3. The transparently shaded area represents the mean of the steady-state signal output within MDG147/pEBS cells with a range of one standard deviation of the mean (n ≥ 3).

**Figure 4.** Primary sequence of the C-terminal end of TM2 from the aromatically tuned EnvZ variants. A Trp-Leu-Phe triplet was repositioned, and the minus series of receptors has the triplet repositioned in the N-terminal direction while the plus series of receptors has the Trp-Leu-Phe triplet repositioned in the C-terminal direction. EnvZ WLF 0 is the wild-type receptor. Residue positions within EnvZ are provided above the primary sequences.

immunoblotting techniques similar to those described in Figure S3 were used to estimate the extent of receptor expression. During our analysis, we assessed whether osmosensing circuits containing the aromatically tuned variants possessed normal levels of CFP fluorescence, YFP fluorescence, and CFP/YFP ratios. We also assessed whether these values were constant regardless of the amount of tuned EnvZ-V5 present. When EPB30 (envZ) cells are grown under the low or high osmolarity regime and express untuned or aromatically tuned EnvZ-V5 from pRD400, the CFP/YFP ratio can be used to estimate the intracellular level of phosphorylated OmpR (Figure S1).

In the uppermost panels of Figure 5, we illustrate that osmosensing circuits tolerate a broad range of EnvZ-V5 levels (same data as that in Figures 3 and S4). When EPB30/pRD400 cells were grown under the low osmolarity regime, steady-state CFP fluorescence was attained between EnvZ-V5 levels of 0.1 and 1.0. Likewise, when these cells were grown under the high osmolarity regime, steady-state CFP fluorescence was maintained between EnvZ-V5 levels of 0.1 and 0.9. In contrast, steady-state YFP fluorescence was observed over the entire range of EnvZ-V5 levels examined. CFP/YFP ratios were dependent on attaining steady-state CFP fluorescence, so the range of EnvZ-V5 required for reaching steady-state was the same as that for CFP fluorescence alone. Trendlines are provided over the ranges of EnvZ-V5 levels where normal signal output was attained. These trendlines were subsequently replicated within the other panels to aid in comparison.

We began by comparing CFP/YFP ratios from circuits containing an aromatically tuned variant to the circuit containing the untuned EnvZ-V5 (left panels in Figure 5). It is noteworthy that circuits containing the aromatically tuned variants of EnvZ-V5 did not show the expected decrease in CFP/YFP at higher levels of EnvZ-V5; therefore, all data points above EnvZ-V5 levels of 0.1 were included during calculation of subsequent trendlines. Under the low osmolarity regime, circuits containing the minus series of receptors (WLF−5 through WLF−1) achieved steady-state output at least equal to that of circuits containing the untuned receptor, as indicated by a CFP/YFP ratio of approximately 0.30–0.35 (compare the light green and light gray trendlines in the left panels of Figure 5). However, within certain circuits, such levels of signal output were attained only at higher EnvZ-V5 levels. This can be observed as the light green trendline passing through the light gray trendline. Circuits containing the plus-series of receptors (WLF+1 and WLF+2) failed to attain normal signal output. In...
Figure 5. Steady-state signal output from osmosensing circuits containing the WLF series of tuned EnvZ receptors. CFP/YFP ratio (left panels), CFP fluorescence (center panels), or YFP fluorescence (right panels) are presented for osmosensing circuits containing one of the aromatically tuned receptors. The amount of receptor present is determined as described in Figure S3. Osmosensing circuits containing the untuned receptor are presented at the top of the figure (WLF 0). Data from EPB30/pRD400 cells grown under the low (open circles) and high (filled circles) osmolarity regimes are shown. These trendlines are present in all charts for comparison to the results from circuits containing the aromatically tuned variants. Trendlines for cells grown under the low osmolarity regime are presented as light gray lines, while those from cells grown under the high osmolarity regime are shown as dark lines. CFP fluorescence was steady between EnvZ-V5 levels of 0.1 and 1.0 when cells were grown under the low osmolarity regime and between EnvZ-V5 levels of 0.1 and 0.9 when grown under the high osmolarity regime. This is in contrast to YFP, which remains steady over the entire range of EnvZ-V5 levels. CFP/YFP was affected by the reduction of CFP fluorescence at either end of the spectrum. However, no reduction in CFP or CFP/YFP was observed at higher levels of the tuned EnvZ-V5 variants. Thus, when determining the trendlines for CFP fluorescence or CFP/YFP for circuits containing the tuned variants, only EnvZ-V5 levels above 0.1 were considered. For circuits containing the tuned variants, the light and dark green lines represent CFP/YFP ratios for EPB30/pRD400 cells grown under the low and high osmolarity regimes, respectively. Likewise, CFP and YFP fluorescence are represented as light and dark blue and yellow trendlines, respectively.
these cases, the light green trendline never passes through the light gray trendline. When EPB30/pRD400 cells were grown under the high osmolarity regime, we observed similar trends. All circuits, with the exception of those containing the WLF+1 or WLF+2 variant, possessed a CFP/YFP that equals or exceeds signal output from those containing the untuned variant (i.e., a CFP/YFP ratio of ~2.5–3.0). Again, this was usually observed at higher receptor levels (compare solid green and black trendlines in the left panels of Figure 5). The steady-state output of several circuits exhibited large changes in CFP/YFP ratio that were dependent on EnvZ-V5 level. To facilitate a more quantitative comparison between circuits, we have calculated the slope (m) of each trendline (Table S1). Of the circuits that attained normal steady state-signal output, changes on EnvZ-V5 level were notable for those containing the WLF-3 (m = +4.5), and WLF-1 (m = +18) variants and to a lesser extent for circuits containing the WLF-5 (m = +1.5) variant.

To gain a further understanding of steady-state signal output from these circuits, we compared changes in the extent of CFP or YFP fluorescence individually (center and right panels of Figure 5, respectively). When cells were grown under the low osmolarity regime, the absolute CFP fluorescence for circuits containing all of the minus series of receptors (WLF-5 through WLF-1) achieved steady-state signal output greater than that from circuits containing untuned EnvZ-V5, and again, this usually occurred at higher receptor levels (compare light cyan and gray trendlines in the center panels). Cells containing circuits with WLF+1 and WLF+2 did not attain normal steady-state CFP fluorescence, even at higher receptor levels. When cells were grown under the high osmolarity regime, only circuits containing the WLF+1 or WLF+2 variants did not approach normal steady-state levels of CFP (compare solid cyan and black trendlines in the center panels). We also observed that circuits containing several tuned variants appeared to be sensitive to the level of EnvZ-V5 present under the low osmolarity regime. This included circuits containing the WLF-5 (m = +35), WLF-4 (m = +32), WLF-3 (m = +62), and WLF-1 (m = +78) variants. Under the high osmolarity regime, the WLF-5 through WLF-2 variants resulted in levels of CFP fluorescence similar to that of circuits containing the untuned variant. Circuits containing WLF-1 remained elevated compared to that of the untuned variant. Interestingly, the WLF+1 variant resulted in a slight decrease in CFP fluorescence at high expression levels, and, in a similar manner to the low osmolarity regime, the WLF+2 variant never resulted in normal levels of CFP fluorescence.

When grown under the low osmolarity regime, circuits containing the WLF-5 variant were the only circuits that approximated YFP fluorescence from circuits containing the untuned receptor (compare the light yellow and gray trendlines in the right panels). Those containing the WLF-4 (m = −90), WLF-3 (m = −18), and WLF-1 (m = −91) variants exhibited a sharp decrease in YFP fluorescence as increasing levels of the tuned variant were present. This is in contrast to circuits containing the WLF-2 (m = +60) and WLF+1 (m = +77) variants, which produced slightly greater than normal YFP fluorescence as the receptor levels increased, while circuits containing WLF+2 never attained normal YFP fluorescence. The observed trends were similar when cells were grown under the high osmolarity regime, with the exception of circuits containing WLF+1 not exhibiting greater than normal levels of YFP fluorescence (compare solid yellow and black trendlines in the right panels).
Under the low osmolarity regime, most circuits containing little tuned EnvZ-V5, i.e., an [EnvZ-V5] of 0.2, possessed signal output similar to that of wild-type circuits. The exception is those containing the WLF+2 variant, which exhibits essentially no CFP or YFP fluorescence regardless of the amount of receptor present. Circuits containing the WLF-2 and WLF+1 variants exhibited decreasing signaling output as the amount of receptor present is increased, which manifested as increasing YFP fluorescence \((m = +0.57, +0.42, \text{respectively})\). The other tuned variants, e.g., WLF-5, WLF-4, WLF-3, and WLF-1, all result in increased signal output as the amount of receptor present is increased, i.e., a slope of CFP/YFP trendline > 0. In most cases, this was observed as increased levels of CFP fluorescence \((m > 0)\) and decreased levels of YFP fluorescence \((m < 0)\), with the WLF-1 variant exemplifying this phenotype (Table S1). Under the high osmolarity regime, a similar pattern is observed, except that the mutants are shifted toward the right end of the curve, as expected (Figure 6).

In summary, the majority of osmosensing circuits containing aromatically tuned receptors resulted in increased in signal output as EnvZ-V5 levels increased, with the exception of those containing the WLF-2 or WLF+1 variants, which resulted in decreased signal output. Circuits containing the WLF+2 variants always possessed the lowest signal output.

These results suggest that aromatic tuning is sufficient to modulate EnvZ signal output in a manner that correlates with the surface of TM2 that the residues are placed upon. One interpretation of this data would be to suggest that the movement of the aromatic residues destabilizes the TM–HAMP junction in manner that mimics signal output; however, additional experimentation would be required to confirm this hypothesis (Figure S5). Although the aromatic residues were repositioned, other residues substitutions occurred, which may also contribute to changes in signal output. One possible example would be the loss of the Arg-182 from the WLF+2 variant. We did not explicitly examine the change in the charge density, but we do address it indirectly below.

To ensure that aromatic tuning was not restricted to these particular residues (Trp-Leu-Phe), another series of aromatically tuned EnvZ receptors was created. However, this time a Trp-Tyr-Ala triplet was employed at the same initial residue positions of 178–180 (Figure S6). In this case, the Trp and Tyr residues were selected because they were previously moved within the aspartate chemoreceptor of *E. coli* (Tar). However, in order to keep the changes as similar as possible between these sets of aromatically tuned receptors, i.e., moving a triplet, an alanyl residue was also repositioned (Trp-Tyr-Ala). We employed the same techniques (Figure S7), and the data is consistent with the TM2 surface being critical (Figure S8), but additional experimentation is required to determine whether this is due to steric repulsion between individual helices at the cytoplasmic end of the TM domain. In addition, the data from the WYA+2 variant demonstrates that Arg-182 is not essential for EnvZ function, suggesting that maintenance of change density in this region is not critical.

**Advantages of Employing Aromatic Tuning To Modulate SHK Signal Output.** We believe that employing aromatic tuning to facilitate stimulus-independent modulation should be applicable to other SHKs because previously published alignments of primary sequences demonstrate that the majority of SHKs in *E. coli* possess aromatic residues at the cytoplasmic polar/hydrophobic interface. In addition, the majority of aromatically tuned EnvZ variants retain the ability to respond to stimulus (Figures 6 and S8), suggesting that their signal output is biased but not locked in either a stimulus-deprived or saturated conformation. In this regard, aromatic tuning is advantageous compared to deletion of entire SHKs or substitution of the conserved His residue involved in autophosphorylation and phosphotransfer because such methods may result in complete loss of kinase or phosphatase activity. Complete loss of activity has been shown to result in nonphysiological cross-talk between various two-component signaling pathways within a cell. On the basis of our results, we propose that aromatic tuning could be used to rapidly assign downstream physiological and developmental processes to particular SHKs (Figure 1).

**Using Aromatic Tuning To Study Receptor Concentration-Dependent Robustness.** Increased sensitivity to changes in SHK levels was seen for all osmosensing circuits containing an aromatically tuned EnvZ variant. A previous kinetic model predicted that the steady-state output of the EnvZ/OmpR osmosensing circuit should be insensitive to fluctuations in the concentration of EnvZ. A related model that predicts a stronger degree of robustness with respect to the regulatory proteins was also recently analyzed. In both cases, the steady-state signal output of the signaling circuit should be independent of the level of SHK, which was observed when wild-type EnvZ or EnvZ-V5 was present within the circuit (Figures 3 and S2–S4). This robustness was observed previously within the intact EnvZ/OmpR, PhoQ/PhoP, and CpxA/CpxR circuits. Here, for each aromatically tuned variant, a different relationship between steady-state signal output and receptor level was observed, apparent as a change in the slope of the CFP/YFP trendlines that are summarized in Table S1, suggesting that the ratio of kinase to phosphatase activities was different within each receptor and always different than wild-type EnvZ (Figures 5 and S6). Therefore, we propose that further biochemical examination of this series of aromatically tuned receptors will provide insight into the precise mechanisms underpinning receptor concentration-dependent robustness within two-component signaling circuits.

**METHODS**

**Bacterial Strains and Plasmids.** *E. coli* strain MC1061 [F− *araD139 Δ(ara-leu)7966 Δ(lac)X74 galU galK hisD25 (rK mC) mcrB1 rpsL] was used for all DNA manipulations. Strain MG1655 (F− *λ*− *iivG rfbS0 rph1*) was used to control for light scattering and cellular autofluorescence. Strains MDG147 [MG1655 (ΦompF*ompP*) (ΦompC*ompP*)24 and EPB30 (MDG147 *envZ*-kan)26] were used for analysis of the steady-state signal output from osmosensing circuits.

To analyze steady-state signal output from osmosensing circuits, plasmid pRD400 was made by adding an in-frame coding sequence for a seven-residue linker (GGSSAAAG) and a C-terminal V5 epitope tag (GKPIPNPLLGLDST). PCR amplification was employed to create a product with a 5′-terminus containing a BgII site corresponding to the internal site within *envZ* and a 3′-terminus encoding the linker, epitope tag, and a SalI restriction site. This product was subsequently cloned into pEnvZ with BgII and SalI, resulting in the removal of an approximately 800 bp region between the previous stop codon in *envZ* and the SalI site. This strategy was used to retain similar IPTG-based induction of EnvZ and EnvZ-V5 from pEnvZ and pRD400, respectively. A previously described plasmid, pEB5, served as an empty vector control that did not express *envZ*. 479
Analysis of Steady-State Signal Output from Osmosensing Circuits. Analysis was performed as described previously\(^{25}\) with slight modification. Briefly, MDG147\(^{24}\) or EPB30\(^{26}\) cells were transformed with pEB5\(^{25}\) pEnvZ\(^{27}\) or pRD400 as required. Fresh colonies were used to inoculate 2 mL overnight cultures of minimal medium A\(^{38}\) supplemented with 0.2% glucose. Ampicillin, sucrose, and IPTG were added where appropriate. Cells were grown overnight at 37 °C and diluted at least 1:1000 into 7 mL of fresh medium. Chloramphenicol was added to a final concentration of 170 \(\mu\)g/mL to inhibit protein synthesis when the cultures reached an OD\(_{600}\) nm \(\sim\) 0.3. Fluorescent analysis was immediately conducted with 2 mL of culture, while the remainder was centrifuged and stored at \(-80^\circ\)C for immunoblotting. All fluorescence measurements were performed with a Varian Cary Eclipse (Palo Alto, CA). CFP fluorescence was measured by using an excitation wavelength of 434 nm and an emission wavelength of 477 nm, while YFP fluorescence was measured by using an excitation wavelength of 505 nm and an emission wavelength of 527 nm. These values were corrected for cell density and sample loading.

**Protein Quantification of EnvZ-V5.** Pellets from cells expressing EnvZ-V5 were analyzed on 12% SDS/acylamide gels. Standard buffers and conditions were used for electrophoresis and immunoblotting.\(^{39}\) Anti-V5 (Inviogen) and anti-\(\beta\)-lactamase (Abcam) primary antibodies were used. Peroxidase-conjugated anti-mouse IgG (Sigma) was used as the secondary antibody. Bands were visualized with the ECL advance western blotting detection kit (GE Healthcare).

**ASSOCIATED CONTENT**

Supporting Information

Various control experiments described throughout the text (Figures S1–S4). A model proposing how the aromatic residues influence signal output (Figure S5). Data for the WYA (Trp-Tyr-Ala) series of aromatically tuned variants (Figures S6–S8). The slopes of the trendlines from Figures 5 and 7 (Table S1). This material is available free of charge via the Internet at http://pubs.acs.org.

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**Notes**

The authors declare no competing financial interest.

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