Evolving New Protein-Protein Interaction Specificity through Promiscuous Intermediates

Graphical Abstract

Highlights
- ParD-ParE toxin-antitoxin systems interact in a highly specific manner
- Toxin-antitoxin systems can coevolve without ever disrupting their interaction
- Promiscuous variants can serve as mutational intermediates during coevolution
- Promiscuous variants are abundant in sequence space and connected to specific variants

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In Brief
Interacting proteins can coevolve through the generation of promiscuous variants, which serve as mutational intermediates that preserve the ability of the two proteins to functionally interact while they evolve.
Evolving New Protein-Protein Interaction Specificity through Promiscuous Intermediates

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SUMMARY

Interacting proteins typically coevolve, and the identification of coevolving amino acids can pinpoint residues required for interaction specificity. This approach often assumes that an interface-disrupting mutation in one protein drives selection of a compensatory mutation in its partner during evolution. However, this model requires a non-functional intermediate state prior to the compensatory change. Alternatively, a mutation in one protein could first broaden its specificity, allowing changes in its partner, followed by a specificity-restricting mutation. Using bacterial toxin-antitoxin systems, we demonstrate the plausibility of this second, promiscuity-based model. By screening large libraries of interface mutants, we show that toxins and antitoxins with high specificity are frequently connected in sequence space to more promiscuous variants that can serve as intermediates during a reprogramming of interaction specificity. We propose that the abundance of promiscuous variants promotes the expansion and diversification of toxin-antitoxin systems and other paralogous protein families during evolution.

INTRODUCTION

Many interacting proteins within the same cell, particularly signaling proteins, are members of large paralogous families that have expanded through duplication and divergence. To expand in number, paralogous interacting proteins typically must become specific after duplication to avoid unwanted cross-talk (Capra et al., 2012; Zarrinpar et al., 2003). The specificity determinants of protein-protein interactions remain poorly defined in most systems. Even in the cases where they have been identified, we lack a detailed understanding of how a new, insulated protein-protein interaction emerges during the course of evolution and, more generally, the mutational paths followed during protein evolution (DePristo et al., 2005).

Computational studies demonstrate that interacting proteins often coevolve. Indeed, identification of coevolving residues has helped guide identification of the specificity determinants of many protein-protein interfaces (Ovchinnikov et al., 2014; Skerker et al., 2008). The implicit notion or underlying model behind these analyses is usually that an interaction-disrupting mutation in one protein can be rescued by a mutation in its partner (Figure 1A). This model, which we call the compensatory mutation model, implies that the system passes through a non-functional or non-interacting state. However, such a state is highly unlikely, particularly for a protein-protein interaction that is critical for the viability of an organism. Alternatively, the specificity of a given protein-protein interaction could change, and become insulated from other paralogous systems, if one of the proteins passes through a promiscuous intermediate (Figure 1B). In this model, an initial mutation in protein A would broaden its specificity, enabling its partner, protein B, to accumulate a mutation that would have disrupted its interaction with the original, ancestral form of protein A. A subsequent mutation in protein A would then narrow its specificity to include the derived, but not the ancestral, form of protein B. In this promiscuous intermediate model, the specificities of the interacting proteins change without ever transitioning through a non-functional intermediate state. Note that in both models, A and B continue to interact through the same set of interfacial residues and do not evolve an alternative interface de novo (Kuriyan and Eisenberg, 2007).

Which of the two models in Figure 1 applies to most pairs of interacting proteins is unclear. In each case, the mutational trajectory involved would produce a signature of pairwise amino-acid coevolution in the phylogenetic record. However, only the latter, promiscuous intermediate model invokes the existence of mutations that are transiently introduced to broaden the specificity of one of the two proteins. The prevalence of such promiscuous states is unknown, as is whether they are easily reached from more specific, extant states.

Bacterial toxin-antitoxin (TA) systems provide an excellent model system for dissecting the coevolutionary dynamics of protein-protein interactions. Originally identified on plasmids, these systems are widely found in bacterial chromosomes, with many species encoding multiple, paralogous copies that share extensive similarity at the sequence and structural levels (Leplae et al., 2011). The biological function of TA systems is unclear, but they have been implicated in stress responses, resistance to phage, formation of persister cells, and bacterial pathogenicity (Yamaguchi et al., 2011). Typically, the toxin is a stable, globular protein that can inhibit cell growth or viability unless antagonized by a cognate antitoxin that directly binds and sequesters the toxin. Changes in the degradation rate or synthesis of the antitoxin can trigger release of the toxin. A toxin is typically encoded in...
the same operon as an antitoxin, and toxin-antitoxin paralogs frequently arise through operon duplications. An unresolved question is whether toxin-antitoxin systems interact in an exclusive one-to-one manner. Genetic data suggest that these interactions may be specific (Fiebig et al., 2010), and the growth inhibitory effects of a toxin are usually rescued only by expressing its co-operonic antitoxin (Hallez et al., 2010; Ramage et al., 2009). However, interaction specificity has only been directly tested in a limited number of cases, and some groups have suggested that toxins and antitoxins encoded in different operons are capable of interacting in vivo and in vitro, possibly forming large, promiscuous networks (Yang et al., 2010; Zhu et al., 2010).

Here, we systematically measure the binding preferences of 20 ParD-ParE TA family members and find that these toxins and antitoxins are highly specific, interacting almost exclusively with their partner from the same operon. This specificity is encoded by a small set of coevolving residues at the toxin-antitoxin interface, and mutations in these residues are sufficient to reprogram a ParD antitoxin to interact with non-cognate ParE toxins. Guided by these findings, we generated a library with ~10^8 variants of the key, specificity-determining residues in a ParD antitoxin and selected mutants that antagonize the cognate toxin, a non-cognate toxin, or both. Strikingly, we find that promiscuous variants that antagonize multiple toxins are easily obtained and are also highly connected in sequence space to specific variants. These results suggest that mutational paths leading to changes in toxin-antitoxin specificity are likely to involve promiscuous intermediates. Such paths enable the reprogramming of toxin-antitoxin specificity through the pairwise coevolution of interfacial residues, but without passing through an intermediate state that disrupts the protein-protein interaction. The abundance of promiscuous states likely facilitates the evolutionary expansion of these and other paralogous protein families following operon and whole-genome duplications during evolution.

**RESULTS**

**Toxins and Antitoxins from the ParDE Family Exhibit High Interaction Specificity**

To systematically measure the interaction specificity of TA systems, we focused on the ParD-ParE family, which is often found in multiple copies on bacterial chromosomes (Fiebig et al., 2010; Leplae et al., 2011) (Figure S1A). We initially cloned the three chromosomally encoded ParD-ParE pairs from the α-proteobacterium *Mesorhizobium opportunismum* into vectors that allow for separate and inducible expression of the ParE toxin and ParD antitoxin. To measure the interaction specificity for these pairs, we then co-transformed all pairwise combinations of toxin and antitoxin plasmids into *E. coli* and assessed whether the induced expression of each ParD antitoxin rescues the growth arrest resulting from inducing each ParE. As a control, we first confirmed that inducing each ParE toxin inhibited growth of *E. coli* (Figure 2A). Then, plating on a medium that induces both ParD and ParE, we observed growth for each of the three cognate ParD-ParE pairings (Figure 2A). No growth was observed for the six non-cognate pairs, indicating that the ParD antitoxins from *M. opportunismum* can only neutralize their cognate ParE toxins.

We extended this analysis to the 20 chromosomally encoded ParDE pairs from eight different bacteria, including the three pairs from *M. opportunismum* (Figure S1B). For this 20 × 20 matrix of ParD and ParE pairs we observed strong interactions between all 20 co-operonic ParDE pairs, but only 11 of the 380 (or 3%) other possible pairings (Figure 2B). Importantly, these cross-reactions were only observed between ParD and ParE proteins not encoded in the same species, indicating that the ParDE pairs within a given organism are typically insulated from one another. These results indicate that ParD antitoxins are highly specific for their cognate ParE toxins.

**Identification of Covarying Residues in ParD and ParE**

As a first step in understanding the molecular basis of specificity in ParD-ParE complexes, we solved a 1.59-Å cocrystal structure of the *M. opportunismum* ParD3 antitoxin bound to ParE3, its cognate toxin. This structure revealed a heterotetrameric asymmetric unit composed of ParD3 and ParE3 dimers (Figure S2A), similar to a *C. crescentus* ParD-ParE structure (Dalton and Crosston, 2010). Crystal packing and an estimated mass of ~87 kDa in solution indicate that the biological assembly is composed of two tetramers (Figures S2B and S2C). Within this complex, each ParD3 subunit makes extensive contacts with a
neighboring ParE3 subunit primarily through its second and third alpha helices, with a total buried surface area of 1,624 Å² (Figure 3A).

Previous work with bacterial two-component signaling systems demonstrated that their interaction specificity is controlled by a subset of residues at the protein-protein interface formed by a histidine kinase and response regulator (Skerker et al., 2008). These specificity-determining residues coevolve to maintain the interaction between cognate signaling proteins. Thus, to pinpoint the residues that contribute to the specificity of ParD-ParE interactions, we used GREMLIN, a pseudo-likelihood-based model for coevolution (Kamisetty et al., 2013; Ovchinnikov et al., 2014), to search for residues that strongly covary in a multiple sequence alignment of concatenated, co-operonic ParD and ParE proteins. This analysis identified 10 residues in ParD and 11 residues in ParE that coevolve most strongly. Hereafter, we call these 21 amino acids “specificity” residues, as our work below indicates that they play the dominant role in determining partner specificity. Mapping these specificity residues onto the ParD3-ParE3 crystal structure indicated that they cluster into two groups at the primary molecular interface formed by these proteins (Figures 3B and 3C). The first group sits at the base of the second alpha helix in ParD3 and covaries with residues in the three-stranded beta sheet in ParE3. The second group clusters in the third alpha helix in ParD3 and covaries with residues in the first and second alpha helices of ParE3. We also used GREMLIN to identify residues within each protein (four in ParD and six in ParE) that coevolve with the specificity residues (Figure 3C and S3A). These “supporting” residues may indirectly contribute to ParD-ParE interaction specificity by influencing the orientation or packing of the interfacial specificity residues.

**Covarying Residues Dictate Interaction Specificity in the ParD-ParE Family**

To determine whether the coevolving residues identified are sufficient to dictate interaction specificity of the ParD-ParE family, we constructed a series of chimeric proteins in which different regions of the M. opportunistum ParD3 were replaced with the corresponding regions of ParD1 or ParD2 (Figure S3B). Replacing the entire C-terminal region of ParD3 with the corresponding
region of ParD1 or ParD2 produced a chimera that lost its ability to interact with ParE3 but gained the ability to interact with ParE1 or ParE2 (Figure 3D). These chimeras involved both clusters of interfacial residues identified as coevolving between ParD and ParE proteins. Replacing only one of these clusters in the ParD3 C terminus was sometimes sufficient to reprogram specificity, but depended on the toxin tested (Figure S3C). These results indicate that the C-terminal region of ParD, which contains the specificity and supporting residues, is sufficient to dictate interaction specificity.

To pinpoint the residues required for interaction specificity, we focused additional mutagenesis on the coevolving residues identified computationally. We generated variants of ParD3 in which all of the specificity and supporting residues were replaced with the corresponding residues in ParD1 or ParD2, for a total of 8 or 9 substitutions, respectively. In each case, we found that these mutations were sufficient to reprogram ParD3 to interact with ParE1 or ParE2 and lose its ability to interact with ParE3 (Figure 3D). Interestingly, ParD3 could be reprogrammed to interact with ParE1 or ParE2 with fewer substitutions. For example, we found sets of four substitutions that were sufficient to reprogram ParD3 to interact with ParE1 or ParE2 (Figure 3D). Taken together, our results indicate that mutating the most highly coevolving residues in an antitoxin can be sufficient to reprogram its interaction specificity, and, in some cases, mutating only a subset of these residues allows a complete switch in partner specificity.

High-Throughput Mapping of Interface Mutant Fitness

The results presented above indicate that antitoxin interaction specificity can be reprogrammed by changing just four residues. But how does specificity change as these four individual substitutions are introduced and does the substitution order matter? Does the specificity of antagonizing one ParE toxin to another change abruptly, or are there promiscuous mutational intermediates? To answer these questions, we sought to generate a large
library of ParD3 variants that included combinations of residues shown to be specific for antagonizing ParE3 or ParE2, as well as the mutational intermediates separating these specific states. To this end, we generated a library of mutants at four of the key interfacial positions in the ParD3 antitoxin, Leu69, Trp60, Asp61, and Lys64 (LWDK). To reduce the complexity of our library, we only allowed residues at each library position that are commonly found in naturally occurring ParD homologs (see Experimental Procedures). The resulting library has a theoretical diversity of 9,360 variants, with 12, 6, 13, and 10 possible residues encoded at the four respective positions of the library (Figure 4A). Deep-sequencing of the relevant region in parD3 in the initial library revealed that >98% of the predicted variants were represented by at least 10 reads and >94% had at least 100 reads (Figure S4A). Measurements of read numbers were highly reproducible between replicates ($R^2 > 0.99$, Figure S4B).

To assess the ability of each ParD3 variant to bind and antagonize ParE3, we co-transformed E. coli with the ParD3 library and an inducible ParE3 vector. When cultured in conditions that do not induce ParD3, cell growth arrested within 200 min after inducing the ParE3 toxin (Figure 4B). In contrast, when the ParD3 library was expressed, growth slowed after inducing the toxin but eventually resumed, suggesting that some fraction of the population could neutralize ParE3 toxicity (Figure 4B). To determine which mutants neutralized ParE3 and hence were enriched during the course of this experiment, we harvested samples every 100 min and deep-sequenced the relevant region of parD3. We observed large changes in the frequency of individual variants over this time course (Figure S4C). For example, the variant containing the wild-type ParD3 residues (LWDK) was enriched ~6-fold, whereas variants with frameshift mutations in parD3, which are presumably non-functional, were depleted ~7-fold (Figure S4C). To validate the functionality of variants inferred from this competitive growth assay, we isolated six mutants that exhibited different frequency dynamics following toxin induction (Figure 4C). We tested these six mutants individually using our toxicity-rescue assay and found clear agreement between the change in the frequency of each variant in the library and its individual plating efficiency (Figure 4D).

To quantify differences in variant behavior during competitive growth, we generated a linear fit to the frequencies of each mutant as a function of time, and then calculated the log-fold expansion of each mutant relative to the rest of the population, producing a raw fitness value ($W_{raw}$) for each mutant. We then transformed these raw fitness values such that the $W$ value for frameshift variants was 0 and the $W$ value for the wild-type (LWDK) sequence was 1; the resulting distribution of $W$ values ranged from $-0.04$ to 1.13 and was highly reproducible between biological replicates (Figure 4E, $R^2 = 0.98$). We found a total of 252 variants with $W$ values > 0.5, representing 2.7% of the total (Figure 4F). This set included the wild-type combination of residues (LWDK) and 31 single, 189 double, and 31 triple mutants relative to the wild-type sequence (Figure S4D). There were no quadruple mutants, as position 60 was invariably tryptophan. The most common residues in this set as a whole were wild-type. However, the identification of 252 variants that can effectively antagonize ParE3 indicates a substantial degree of functional degeneracy in the ParD3 interfacial residues.

Next, to assess the ability of each ParD3 variant to antagonize the non-cognate toxin ParE2, we repeated the competitive growth experiment but co-transformed E. coli with our ParD3 library and an inducible ParE2 vector. As before, we observed growth rescue following ParD3 library expression with large changes in the frequency of individual variants over time (Figures 4G and S4E). However, the frequency changes observed here differed from those observed against the cognate toxin ParE3. For example, a variant containing the specificity residues found in the native ParD2 antitoxin, AWIL, was enriched in the ParD3 library screened against ParE2 but was depleted when screened against ParE3 (Figures 4C and 4H). We quantified variant fitness as before and found a total of 151 variants (1.6% of the total) capable of antagonizing ParE2 with $W$ values > 0.5 (Figures 4I and S4E). The most common residues were Ala, Trp, Leu, and Lys. However, we noted important differences between variants reactive against ParE2 and ParE3, particularly at the last two variable positions in our library. ParE2-specific variants tended to have small hydrophobic or positively charged residues at position 61, whereas ParE3-specific variants favored negatively charged residues at this position (Figures 4F and 4I). Additionally, ParE2-specific variants were more likely to contain small hydrophobic residues at position 64, whereas ParE3-specific variants tended to have positively charged residues (Figures 4F and 4I).

**Mutational Paths That Reprogram Specificity Tend to Involve Promiscuous Variants**

To more systematically probe the sequence space governing the specificity of ParD3, we generated a scatterplot of ParD3 variant fitness when screened against the ParE2 or ParE3 toxin (Figure 5A). This analysis revealed variants spanning all ranges of fitness, including those capable of antagonizing ParE2, ParE3, or both toxins simultaneously. We identified a total of 31 promiscuous variants ($W > 0.5$ for both toxins), which represents a subset of the 252 ParE3-reactive and 151 ParE2-reactive variants (Figure 5B). We then grouped variants by specificity class (Figure S5A) and found that the promiscuous variants, such as LWE, tended to harbor sequence elements from both ParD3 and ParD2, often with negatively charged residues at position 61 (ParD3-like) and aliphatic residues at position 64 (ParD2-like) (Figure 5C).

To visualize the connectivity of functional variants in sequence space, we created a force-directed graph where individual nodes represent functional variants with lines connecting variants that differ by a single amino acid (Figure 5D). Node sizes increase with greater connectivity and node colors represent the specificity class of a given variant (Figure 5D). The resulting graph was densely interconnected but generally grouped variants based on their specificity. The average number of edges per node, or degree, was 17.8 and ranged from 7 to 31. However, we noted that the average number of edges per node was 23% higher for promiscuous variants than for variants specific for ParE2 or ParE3 (Figure 5E). We also generated a force-directed graph in which edges represent variants that differ by a single-nucleotide substitution, following the standard genetic
code (Figure S5B). For this graph, promiscuous variants were, on average, 31% more connected to other nodes than their ParE2- or ParE3-specific counterparts (Figure 5E). This increased connectivity of promiscuous variants was highly significant for both amino acid and nucleotide graphs, as it was lost when the edges of each graph were randomly shuffled (p < 10^{-4}).
Figure 5. Specificity-Reprogramming Paths Are Highly Enriched for Promiscuous Variants

(A) Fitness of ParD3 variants against ParE2 and ParE3. Green, specific for ParE3; blue, capable of antagonizing both ParE2 and ParE3; red, specific for ParE2. Histograms of fitness values against ParE2 and ParE3 are shown.

(B) Venn diagram of ParD3 variants reactive against ParE3, ParE2, or both.

(C) Frequency logo of promiscuous ParD3 variants (WE2 > 0.5, WE3 > 0.5).

(D) Force-directed graph of all ParD3 variants reactive against ParE3 or ParE2 (W > 0.5). Nodes represent individual variants and edges represent single amino-acid substitutions. Node size scales with increasing degree and color corresponds to the specificity classes in (A).

(E) Average number of edges per node for the indicated categories of ParD3 variants. Error bars indicate SEM.

(F) Examples of “switch-like” and “promiscuity-based” mutational paths from an E3-specific variant to an E2-specific variant with the fitness against each variant color-coded based on the scale shown.

(G) Left, percentage of “switch-like” and “promiscuity-based” paths from the wild-type ParD3 sequence (LWDK) to each of the 66 ParE2-specific variants (WE2 > 0.5, WE3 < 0.1). Right, same as left panel but for 10,000 simulations in which the graph edges were randomly shuffled while keeping the total edge count and degree distribution constant. Error bars represent SEM.

(H) Histogram representing percentage of “promiscuity-based” paths in 10,000 edge shuffling simulations; red line indicates percentage for the observed amino acid graph.

Also, see Figure S5.
The dense connectivity of promiscuous variants suggested that mutational paths that change ParD3 specificity from ParE3-specific to ParE2-specific, or vice versa) tend to travel through promiscuous intermediates. To test this hypothesis, we first defined two types of specificity-reprogramming paths. Note that for the following analysis, we exclude paths in which ParD3 fails to interact with both ParE3 and ParE2 (also see Discussion). The first class of paths are “switch-like” and only involve intermediates that are specific for ParE2 or ParE3, whereas the second class of paths are “promiscuity-based” and travel through at least one intermediate that can inhibit both ParE2 and ParE3 (Figure 5F). To determine whether paths that change the interaction specificity of ParD3 tend to be switch-like or promiscuity-based, we identified all shortest mutational paths from the wild-type ParD3 variant (LWDK) to each of the 66 variants that are highly specific for ParE2 ($W_{E2} > 0.5, W_{E3} < 0.1$; Figure S5A); for this analysis, each mutational step involved a single amino-acid substitution. We found a total of 370 shortest paths, of which 40% involved a promiscuous intermediate (Figure 5G). The percentage of paths via promiscuous intermediates increased to 61% when considering only paths that involve single-nucleotide substitutions (Figure 5G).

To determine whether the number of paths that involve promiscuous variants is greater than would be expected by chance, we generated graphs in which the edges were randomly shuffled, and again calculated the percentage of each class of paths from ParD3 (LWDK) to the ParE2 highly specific variants. For these graphs with randomized edges, the percentage of paths involving promiscuous intermediates dropped to 15% for the amino acid neighbor graph and 20% for the nucleotide neighbor graph (Figures 5G and 5H). Thus, the enrichment of promiscuity-based paths in the observed graphs is significant ($p < 0.005$) (Figures 5G, 5H, and S5F). Collectively, our results demonstrate the dense connectivity of functional variants in the sequence space governing ParD-ParE interaction specificity and reveal that specificity-reprogramming paths are highly enriched for those that involve promiscuous variants, which may facilitate the evolution of ParD-ParE systems with new specificities.

**Epistasis: Mutational Order Dictates Specificity Class of Intermediate Variants**

Inspection of the paths connecting ParD3 variants with different specificities indicated that the third and fourth library positions, residues 61 and 64 in ParD3, contribute significantly to the insulation of the ParD-ParE system. For instance, the wild-type residue combination in ParD3, LWDK, renders it specific for binding to ParE3, whereas the double-mutant variant LWKL is specific for ParE2. Strikingly, however, the two possible paths connecting LWDK and LWKL are in different classes (Figure 6A). A single ParD3 substitution (K64L in LWDL) resulted in promiscuous binding to ParE2 and ParE3, whereas a second substitution in this background (D61K in LWKL) resulted in specificity for ParE2 (Figure 6A). In contrast, incorporating these substitutions in the reverse order, D61K and then K64L, resulted in a switch-like change in specificity in which the initial D61K substitution retained specificity for ParE3, but then enabled the subsequent K64L substitution to produce a ParE2-specific antitoxin (Figure 6A). These results underscore how a small number of mutations can fully reprogram protein-protein interaction specificity and demonstrate that the order of mutations can strongly affect whether the path to a new specificity state involves a promiscuous intermediate or a rapid switch.

Our finding that changes in specificity can depend strongly on the order of substitutions represents a form of epistasis, broadly defined as cases where the functional effect of individual substitutions is context-dependent rather than additive and independent (Lehner, 2011). To more broadly quantify this epistasis for...
the ParD3 interface residues, we first defined six types of specificity-reprogramming paths that involve two amino-acid substitutions (Figure 6B). Three of the six path types are epistatic with the two intermediates having different specificities, implying that substitution order influences changes from ParE3 to ParE2 specificity. We quantified the path type for each case in which two substitutions reprogram ParD3 from being specific for ParE3 (\(W_{E3} > 0.5, W_{E2} < 0.5\)) to being specific for ParE2 (\(W_{E2} < 0.5, W_{E3} > 0.5\)) and found a total of 2,653 such cases, of which 92% were epistatic (Figure 6B). The percentage of epistatic paths was robust to the threshold used for defining positive interactions (Figures S6A and S6B). Taken together, our results highlight the pervasive effects of epistasis on ParD function. Although studies of epistasis typically consider the interdependence of individual substitutions with respect to protein folding or a single-protein function (Kondrashov and Kondrashov, 2015; Lehner, 2011), our findings indicate that epistasis can also manifest at the level of interaction specificity. This form of epistasis may significantly impact the evolution of new ParD-ParE systems. Promiscuous intermediates enable a change in protein-protein interaction specificity without passing through a non-functional state, in which a liberated toxin would suppress growth and proliferation (Figure 1A). Thus, the epistasis documented here may fundamentally restrict mutational trajectories during evolution to those involving promiscuous intermediates.

**Mutational Trajectories to an Orthogonal ParD3-ParE3 Pair****

Thus far, we have considered changes to one side of the ParD-ParE interface. To probe how the interaction specificity of a ParD-ParE protein pair coevolves, we sought to generate a variant of the toxin ParE3 that does not interact with ParD3, and then select ParD3 variants from our library that can neutralize this novel toxin. To this end, we generated a variant of the toxin, called ParE3*, that retains toxicity but is incapable of binding to the ParD3 antitoxin. In particular, we mutated five ParE3 positions (Arg54, Arg58, Ala61, Met63, and Leu72, or RRAML) that strongly covary with the specificity residues in ParD3. We mutated RRAML→VEIRF, as each individual variant residue was frequently observed in ParE3 homologs and was chemically different from the corresponding wild-type residue (Figure S7A). As expected, we found that ParE3* retained toxicity but was no longer neutralized by ParD3 (Figure 7A).

To determine whether variants in the ParD3 library neutralized ParE3*, we performed a competitive growth experiment following co-transformation. As before, we converted changes in variant frequencies to fitness values, which were highly reproducible (\(R^2 = 0.96, \text{Figure S7B}\)). Sequence analysis of the high-fitness mutants (\(V > 0.5\)) revealed large differences in amino-acid preferences at positions 60 and 61 relative to those shown above (Figures 4F and 7B). In particular, for the ParD3 variants that neutralized ParE3*, the invariant Trp60 was replaced by Ile/Val/Leu and the strong preference for a negatively charged residue at position 61 was replaced by positively charged or neutral residues (Figures 4F and 7B). One of the high-fitness variants with specificity residues LIAK, renamed ParD3*, no longer neutralized ParE3 but robustly interacted with ParE3* (Figure 7C). Taken together, our results indicate that mutations in the specificity residues of ParD3 and ParE3 are sufficient to create an orthogonal, interacting protein pair.

Our results indicate that mutational paths leading to a change in ParD specificity tend to pass through promiscuous intermediates (Figure 5). Thus, we wanted to determine whether mutational paths between the wild-type ParD3-ParE3 and the orthogonal ParD3*-ParE3* systems also pass through promiscuous intermediates, thereby changing the specificity of both proteins without disrupting their interaction. We therefore generated variants of ParE3 containing all possible subsets of the substitutions in ParE3* (32 mutants) and variants of ParD3 containing all possible subsets of the substitutions in ParD3* (4 mutants). We then co-transformed each possible pairing of ParD3 and ParE3 variants (128 pairs total) into E. coli and assessed interaction using the toxicity-rescue assay (Figure 7D). Interestingly, 90 of the 128 pairs of ParD3 and ParE3 variants were capable of interacting, likely because most (17 of 32) of the ParE3 variants were promiscuous, which we define as interacting strongly with both ParD3 and ParD3* (Figure 7D).

To determine whether paths between the wild-type and insulated ParD-ParE pairs tend to pass through promiscuous intermediates, we first enumerated the total number of trajectories between these systems. Assuming one residue is changed per step and no reversions are considered, there are 5,040 paths from ParD3-ParE3 to the orthogonal ParD3*-ParE3* pair; of these paths, 1,030 retain functionality at each intermediate step. Strikingly, we found that all of these 1,030 functional paths passed through at least one promiscuous intermediate of ParE3 with an average of five promiscuous ParE3 intermediates per path (Figure S7C). The prevalence of these promiscuous states may enable the ParD-ParE system to readily evolve a new interaction specificity. An initial broadening of ParE3 specificity enables the movement of ParD3 in sequence space, followed by a narrowing of ParE3 specificity in the final step (Figure 7E). By contrast, mutational paths in which a substitution in either ParD or ParE yields a “switch-like” change in specificity would, by definition, be broken until a second substitution restores the interaction. Thus, our results support the notion that the coevolution and expansion of the ParD-ParE family occurs through promiscuous intermediates.

**DISCUSSION**

**Mutational Trajectories and the Coevolution of Protein-Protein Interactions**

Interacting proteins coevolve, and the identification of coevolving amino acids in two proteins can often help to pinpoint the residues that mediate their interaction. Such analyses are typically predicated on the idea that a mutation in one protein that disrupts an interaction then drives selection of a compensatory mutation in the partner, thereby restoring the interaction (Figure 1A). However, this model implies that organisms tolerate (at least transiently) a non-functional, or less functional, interaction, which seems unlikely if the protein-protein interaction is essential for viability. Our results provide a solution to this conundrum, demonstrating experimentally how interacting proteins can coevolve and acquire new specificity by having one of the proteins pass through a promiscuous intermediate (Figure 1B). For
instance, a mutation in an antitoxin can initially broaden its specificity; the toxin can then accumulate a mutation that moves it in sequence space but retains its interaction with the antitoxin. A subsequent substitution in the antitoxin can then narrow its specificity to include the mutated toxin and exclude the original form. The net result is a change in specificity without disruption of the protein-protein interaction, which is critical as a disruption at any step would liberate a toxin that prevents growth and proliferation. This model for protein coevolution involves a minimum of three instead of two mutations but means that the protein-protein interaction is functional at each step. Thus, such mutational trajectories could be entirely neutral but importantly would retain a pairwise-coevolution signature in multiple sequence alignments.

Our systematic identification of ParD3 variants that can antagonize ParE3, ParE2, or both revealed an abundance of promiscuous variants in sequence space that are, on average, more highly connected to other functional variants than are specific variants. Consequently, the mutational trajectories that reprogram the specificity of ParD3 frequently involve promiscuous intermediates (Figures 5F and 5G). The high frequency of mutational paths involving promiscuous intermediates was seen when
considering transitions in ParD3 from being specific for ParE3 to specific for ParE2, and even more so when considering mutations on both sides of the interface. We assessed the complete set of mutational trajectories between the wild-type ParD3-ParE3 and the orthogonal ParD3*-ParE3* by testing 128 pairwise interactions between all possible ParD3 and ParE3 mutational intermediates. Strikingly, 17 of the 32 ParE3 intermediate variants were promiscuous, or capable of interacting with both the ParD3 and ParD3* variants (Figure 7). Consequently, all of the functional paths between ParD3-ParE3 and ParD3*-ParE3* involved at least one promiscuous intermediate, with most involving more than five (Figure 7). Our results thus suggest that promiscuous variants of ParD and ParE are abundant in sequence space and that promiscuity-enabling mutations can facilitate the evolution of new interaction specificities while still using the same set of interfacial residues.

A similar principle may apply to other protein-protein interactions throughout biology, even those not involving toxic proteins. The disruption of a given protein-protein interaction could prevent the execution of an essential cellular function or lead to an unwanted, detrimental interaction with another protein, thus favoring coevolutionary trajectories that retain function at each step. This same principle may also underlie the coevolution of transcription factors and their DNA binding sites. The evolutionary history of a steroid hormone receptor and its recognition repertoire is largely determined by a small number of interfacial residues (Anderson et al., 2015). Several of the intermediates were promiscuous and may have facilitated coevolution of the receptor and its recognition element toward a new specificity without disrupting the interaction. However, that study only considered mutational intermediates containing residues present in the ancestral or derived states, and our analyses of the ParD-ParE interface suggest that promiscuous intermediates can also involve substitutions that appear in neither the ancestral nor the derived states.

Like many protein families, toxin-antitoxin systems can expand through duplication and divergence. The duplication of a toxin-antitoxin system could allow one of the protein pairs to wander unconstrained in sequence space toward a new interaction specificity via switch-like paths that involve non-functional intermediates. After a duplication, one antitoxin could accumulate interaction-disrupting substitutions while its toxin is still inhibited by the other antitoxin. The toxin could then subsequently mutate to restore an interaction with the derived antitoxin. However, this scenario assumes that the evolving antitoxin does not, in the intermediate state, interact inappropriately with other proteins, and it assumes that the other antitoxin is produced at sufficiently high levels to inhibit 2-fold more toxin, i.e., that there is normally a significant excess of free antitoxin, which may not be the case. Determining whether and when switch-like or promiscuous paths are followed will require careful reconstructions of toxin-antitoxin evolution.

High-Throughput Mapping of Protein Interaction Specificity

Deep mutational scanning via next-generation sequencing is a relatively new approach for interrogating the relationship between protein sequence and function, including folding, enzymatic activity, or the binding of a target protein or RNA (Fowler and Fields, 2014). These studies have begun to reveal the functional degeneracy of proteins by examining all, or nearly all, possible single mutants of a given protein. Similar approaches have also been used to probe subsets of all possible double and higher-order mutants (Melamed et al., 2013) or to systematically probe all possible mutants at a limited set of positions (Podgornaia and Laub, 2015).

Deep mutational scans have been focused primarily on how mutations alter a single function or protein interaction. One study examined the ability of a PDZ domain to interact with both a cognate and non-cognate peptide ligand (McLaughlin et al., 2012), but only queried single-point mutants. However, the interaction specificity of a protein is a distributed property of multiple amino acids, and the prevalence of epistasis means that the behavior of multiple mutations is difficult to infer from the properties of the corresponding single mutants. We queried a diverse library of ParD3 variants harboring multiple mutations of key specificity residues against two separate proteins: the cognate toxin ParE3 and the non-cognate toxin ParE2. This focused library approach was possible as the specificity of ParD3 is largely determined by a small number of interfacial residues (Figure 3). Our approach yielded a high-density map of the sequence space of ParD3 that underpins its substrate interaction specificity (Figures 5A–5D). From these data, we uncovered the residues in ParD3 most responsible for its selective binding of one toxin over another (Figures 4F and 4I). We found that three positions (60, 61, and 64) primarily dictate specificity, with substitutions at two sites (61 and 64) sufficient to switch ParD3 from antagonizing ParE3 to ParE2, and substitutions at an overlapping set of sites (60 and 61) sufficient to switch ParD3 from antagonizing ParE3 to ParE3*. As noted, our results also demonstrated the existence of many residue combinations that promote a promiscuous state of ParD3 or ParE3. Mutations that render proteins more promiscuous, with respect to catalytic activities or binding partners, has been noted anecdotally (Aharoni et al., 2005; Bloom and Arnold, 2009), but the prevalence of such states and, importantly, their accessibility from more specific, wild-type states has never been mapped in a comprehensive manner.

By building and screening libraries harboring multiple mutations, our work also sheds new light on protein epistasis and the non-additive relationship of individual substitutions. Epistasis has been well documented but is typically assessed with respect to a single-protein function. By contrast, the epistasis documented here for ParD3 pertains to its specificity and interaction with two different proteins, revealing interdependencies that would be missed when considering only a single function. For instance, consider the example in Figure 6A where ParD3 transitions from the E3-specific residues LWDK to the E2-specific residues LWKL. With respect to antagonizing the toxin ParE3, the two single mutants, LWDL and LWKK, are each functional. However, with respect to toxin ParE2, LWDL is functional whereas LWKK is not, reflecting a non-additive relationship between the two substitutions leading to the double mutant LWKL. This type of epistasis may, like other forms of epistasis, restrict the evolution of ParD-ParE systems, which likely follows
mutational paths that involve promiscuous states, as discussed above.

**Interaction Specificity of Toxin-Antitoxin Systems**

The specificity of interactions in bacterial toxin-antitoxin systems had previously been unclear, with some reports indicating that these protein-protein interactions are specific (Fiebig et al., 2010) and others suggesting that TA systems form large, cross-reactive networks (Yang et al., 2010; Zhu et al., 2010). Here, by performing a systematic assessment of interaction specificity for a TA family, we found that ParD antitoxins typically exhibit an exquisite preference for binding to their co-transcribed ParE toxins, forming exclusive, cognate pairs. Of 180 non-cognate pairings tested, we found cross-talk in only 11 cases (Figure 2) and, importantly, no cross-talk was observed for non-cognate pairs present in the same species.

The high degree of protein-protein interaction specificity observed for the ParD-ParE family is similar to that observed for other large, paralogous protein families (Newman and Keating, 2003; Skerker et al., 2008; Stöffler et al., 2007; Zarrinpar et al., 2003). The specificity of many of these paralogous families has been attributed to selection against detrimental cross-talk (Capra et al., 2012; Zarrinpar et al., 2003), raising the possibility that the ParD-ParE family may be under similar selective pressures. However, the biological rationale for maintaining the specificity of TA systems is unclear, and will require a deeper understanding of the function of these systems in bacterial physiology.

**Final Perspective**

In sum, our work provides a rationale and molecular basis for how protein interaction specificity can change and how two proteins can coevolve without involving non-functional intermediates. Mutations that produce promiscuity have been described for a variety of proteins, but the frequency of such mutations and their accessibility from more specific states had been unclear. Our results indicate that, at least for ParD3 and likely other proteins, promiscuous mutants are prevalent and easily reached from the wild-type sequence through a single mutation. The prevalence of promiscuous intermediates may facilitate the expansion of toxin-antitoxin systems and, more broadly, other paralogous protein families.

**EXPERIMENTAL PROCEDURES**

**ParD3-ParE Structure Analysis**

For details on the structural analysis of *M. opportunistum* ParD3 and ParE3, see Supplemental Experimental Procedures.

**Identification of Coevolving Residues**

Coevolving residues in the ParDE family were identified using GREMLIN at http://gremlin.bakerlab.org. Input sequences were ParD3 and ParE3 from *M. opportunistum*, and we set the number of iterations to four and the E-value cutoff to 1E-04. To identify specificity residues, we isolated all residue pairings that had a scaled coupling score greater than 1.25. To identify supporting residues, we performed the following iterative procedure using a score cutoff of 1.25: (1) identify residues within ParD or ParE that covary with the specificity residues; (2) identify residues within ParD or ParE that covary with either the specificity residues or the supporting residues identified in step (1); (3) repeat step (2) until no new supporting residues are identified.

**ParD3 Library Construction and Analysis**

For details on construction of the ParD3 library, see the Supplemental Experimental Procedures. To assess the ability of each ParD3 variant to antagonize different ParE toxins, *E. coli* cells harboring the ParD3 plasmid library were electroporated with a plasmid containing an arabinose-inducible copy of the ParE toxin. Cells were grown overnight in 200 ml M9L supplemented with 0.4% glucose and antibiotics. The following day, cells were spun down, washed in 50 ml of M9L, and re-suspended at an OD of 0.03 in 500 ml of M9L supplemented with 100 μM IPTG (to induce the ParD3 library) and antibiotics. Cells were grown out at 37°C with shaking for 100 min, and then ParD3 toxin expression was induced by the addition of 0.2% arabinose. Cell density was measured every 20 min and samples (50 ml) were taken every 100 min, pelleted, and frozen at −20°C. Competitive liquid growth assays were performed in duplicate. Plasmid DNA was extracted and used as template for PCR (20 cycles) with custom barcoded primers containing Illumina flowcell adaptor sequences. Samples were sequenced on an Illumina HiSeq and then filtered, counted, and converted to fitness values as described in the Supplemental Experimental Procedures.

**SUPPLEMENTAL INFORMATION**

Supplemental Information includes Supplemental Experimental Procedures, seven figures, and one table and can be found with this article online at http://dx.doi.org/10.1016/j.cell.2015.09.055.

**ACKNOWLEDGMENTS**

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**REFERENCES**


Supplemental Figures

(A) Homologs of *Mesorhizobium opportunistum* ParD3 and ParE3 were identified independently using an iterative jackhammer search. ParD and ParE homologs were then paired if they were neighboring genes in a given genome. The number of bacterial genomes that have the indicated number of ParD-ParE homologs is shown.

(B) Name and genome accession numbers for the twenty ParD-ParE systems examined in Figure 2.

**Figure S1.** TA Systems Are Widely Present on Bacterial Genomes, Related to Figure 2

1. *Methylobacterium populi* BJ001
   - ParD: Mbp parDE1, GeneID: 188584325
   - ParE: Mbp parDE2, GeneID: 188584324
2. *Methylobacterium populi* BJ001
   - ParD: Mbp parDE1, GeneID: 188579862
   - ParE: Mbp parDE2, GeneID: 188579861
3. *Methylobacterium populi* BJ001
   - ParD: Mbp parDE1, GeneID: 188584614
   - ParE: Mbp parDE2, GeneID: 188584613
4. *Methylobacterium populi* BJ001
   - ParD: Mbp parDE1, GeneID: 188584366
   - ParE: Mbp parDE2, GeneID: 188584365
5. *Methylobacterium populi* BJ001
   - ParD: Mbp parDE1, GeneID: 188583979
   - ParE: Mbp parDE2, GeneID: 188583978
6. *Sphingobium chlorophenolicum* L-1
   - ParD: Sc parDE1, GeneID: 334344591
   - ParE: Sc parDE2, GeneID: 334344592
7. *Sphingobium chlorophenolicum* L-1
   - ParD: Sc parDE1, GeneID: 334346364
   - ParE: Sc parDE2, GeneID: 334346365
8. *Sphingobium chlorophenolicum* L-1
   - ParD: Sc parDE1, GeneID: 334346363
   - ParE: Sc parDE2, GeneID: 334346362
9. *Mesorhizobium opportunistum* WSM2075
   - ParD: Mo parDE1, GeneID: 337268996
   - ParE: Mo parDE2, GeneID: 337268995
10. *Mesorhizobium opportunistum* WSM2075
    - ParD: Mo parDE1, GeneID: 337268962
    - ParE: Mo parDE2, GeneID: 337268961
11. *Mesorhizobium opportunistum* WSM2075
    - ParD: Mo parDE1, GeneID: 337270049
    - ParE: Mo parDE2, GeneID: 337270048
12. *C. crescentus* NA1000
    - ParD: Cc parDE1, GeneID: 221233854
    - ParE: Cc parDE2, GeneID: 221233853
13. *Sinorhizobium meliloti* 1021
    - ParD: Sm parDE1, GeneID: 16263810
    - ParE: Sm parDE2, GeneID: 16263811
14. *Sinorhizobium meliloti* 1021
    - ParD: Sm parDE1, GeneID: 15966432
    - ParE: Sm parDE2, GeneID: 15966431
15. *Paracoccus denitrificans* PD1222
    - ParD: Pd parDE1, GeneID: 119386537
    - ParE: Pd parDE2, GeneID: 119386538
16. *Paracoccus denitrificans* PD1222
    - ParD: Pd parDE1, GeneID: 119386578
    - ParE: Pd parDE2, GeneID: 119386787
17. *Rhodopseudomonas palustris* Bt818
    - ParD: Rp parDE1, GeneID: 90423718
    - ParE: Rp parDE2, GeneID: 90423719
18. *Rhodopseudomonas palustris* Bt818
    - ParD: Rp parDE1, GeneID: 90421684
    - ParE: Rp parDE2, GeneID: 90421683
    - ParD: Rl parDE1, GeneID: 209552210
    - ParE: Rl parDE2, GeneID: 209552209
    - ParD: Rl parDE1, GeneID: 209552404
    - ParE: Rl parDE2, GeneID: 209552403

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Figure S2. *M. opportunistum* ParD3-ParE3 Is a Dimer of Tetramers, Related to Figure 3

(A) Structure of the tetrameric *M. opportunistum* ParD3-ParE3 complex. Light orange, ParE3 dimer; light blue, ParD3 dimer. The full complex is a dimer of tetramers (not shown).

(B) Estimation of the oligomeric state of the ParD3-ParE3 complex using size exclusion chromatography. Comparison of ParD3-ParE3 elution profile (left panel) to molecular weight standards (right panel) yields an estimated molecular weight of ~87 kDa, which is consistent with a dimer of tetramers.

(C) Structure of the octameric ParD3-ParE3 complex, shown as in left panel of (A). Light orange, ParE3 dimers; light blue, ParD3 dimer of dimers.
Figure S3. Mapping of Supporting Residues onto the ParDE3 Crystal Structure and List of ParD3 Mutants Tested, Related to Figure 3

(A) Structure of *M. opportunistum* ParD3-ParE3 with the supporting residues (Figure 3C) shown in gray space-filling representation.

(B) Sequences of ParD3 mutants tested in Figure 3D and in (C). Substitutions relative to the ParD3 wild-type sequence are shown in red.

(C) Mutants in the ParD3 C terminus were tested against each ParE homolog from *M. opportunistum*, as in Figure 3D.
Figure S4. Statistics on High-Throughput Sequencing of ParD3 Library against ParE3 and ParE2 Toxin, Related to Figure 4

(A) Number of library variants present with at least the indicated number of reads.

<table>
<thead>
<tr>
<th></th>
<th>&gt;3 reads</th>
<th>&gt;10 reads</th>
<th>&gt;100 reads</th>
</tr>
</thead>
<tbody>
<tr>
<td>replicate 1</td>
<td>9290 (99.3%)</td>
<td>9215 (98.5%)</td>
<td>8876 (94.8%)</td>
</tr>
<tr>
<td>replicate 2</td>
<td>9292 (99.3%)</td>
<td>9211 (98.4%)</td>
<td>8860 (94.7%)</td>
</tr>
<tr>
<td>both</td>
<td>9267 (99.0%)</td>
<td>9200 (98.3%)</td>
<td>8835 (94.4%)</td>
</tr>
</tbody>
</table>

(B) Read counts are highly reproducible between replicates. The ParD3 library was independently transformed into *E. coli*, grown to saturation overnight, subjected to deep sequencing and read counts compared.

(C) Changes in library variant frequencies following induction of the ParE3 toxin. Scatterplot of variant frequency pre-toxin induction (200 min.) and after 500 min of toxin induction (700 min.). Grey point, wild-type LWDK sequence; yellow points, frameshift mutations predicted to be non-functional.

(D) Number of functional variants with the indicated number of mutations relative to the wild-type, LWDK.

(E) Changes in library variant frequencies following induction of the ParE2 toxin. Scatterplot of variant frequency pre-toxin induction (200 min.) and after 500 min of toxin induction (700 min.). Grey point, wild-type LWDK sequence; yellow points, frameshift mutations predicted to be non-functional.

(F) Scatterplot indicating reproducibility of fitness measurements made for ParD3 library screened against ParE2.
Figure S5. Sequence Composition by Specificity Class; Connectivity of Nucleotide Neighbors, Related to Figure 5
(A) Amino acid composition of variants in each specificity class, defined based on the thresholds indicated.
(B) Force-directed graph of all ParD3 variants with $W > 0.5$ against ParE3 or ParE2. Nodes represent individual variants and edges represent single-nucleotide substitutions. Node size scales with increasing degree and color corresponds to specificity classes in Figure 5A.
(C) The greater connectivity of promiscuous variants is highly significant. We randomized the edge connectivity in the amino-acid substitution graph while keeping the number of edges constant. We then calculated the percentage greater connectivity of promiscuous variants versus non-promiscuous variants. Grey, result of 10,000 simulations; red line, observed value.
(D) Same as (C), except calculated on the nucleotide substitution graph.
(E) Promiscuous variants have a greater number of average edges per node than ParE2 highly specific variants ($W_{E2} > 0.5, W_{E3} < 0.1$) or ParE3 highly specific variants ($W_{E3} > 0.5, W_{E2} < 0.1$). Grey dotted line indicates average for ParE2/ParE3 highly specific variants.
(F) Enrichment of “promiscuity-based” paths is highly significant. Same calculations as Figure S5H, except for the nucleotide substitution graph.
Figure S6. The Prevalence of Epistasis Is Not Sensitive to Fitness Thresholds, Related to Figure 6

(A) The percentage of epistatic paths (as defined in Figure 6B) is plotted for different fitness thresholds used to define functional variants.

(B) Shown are the six path types that reprogram ParD3 specificity from ParE3 to ParE2 in two mutational steps, as in Figure 6B. Below each path type is the percentage of mutational paths in each category when using a threshold of 0.25 to define a positive interaction.
Figure S7. Generation and Testing of ParE3* Variant against ParD3 Library; Mutational Paths between ParD3-ParE3 and ParD3*-ParE3*, Related to Figure 7

(A) An alignment of ParE homologs was analyzed for the most commonly occurring residues at positions 54, 58, 61, 63, and 72 in ParE3 (middle column). The residue combinations were then scored based on their chemical similarity to the residues in ParE3, RRAML, using BLOSUM62 (right column). We chose to incorporate the residue combination VEIRF given that it was the most distant from RRAML by BLOSUM62.

(B) Fitness measurements of the ParD3 library against the ParE3* toxin are reproducible between biological replicates.

(C) Most mutational paths between ParD3-ParE3 and ParD3*-ParE3* pass through multiple promiscuous ParE3 intermediates. Promiscuous ParE3 intermediates are defined as those capable of interacting with both ParD3 and ParD3*. Only functional paths are scored.
Evolving New Protein-Protein Interaction

Specificity through Promiscuous Intermediates

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Supplemental Experimental Procedures

Bacterial strains and media

*Escherichia coli* strains were grown in M9L medium (M9 minimal medium supplemented with 5% LB (v/v) and 0.4% glycerol) at 37°C, unless otherwise indicated. To induce expression from the $P_{BAD}$ and $P_{lac}$ promoters, media was supplemented with 0.2% arabinose or 100 µM IPTG, respectively. All toxins were cloned into the SalI and HindIII sites of the arabinose-inducible pBAD33 vector, and all antitoxins were cloned into the SalI and HindIII sites of the IPTG-inducible pEXT20 vector. Toxin and antitoxin plasmids were cotransformed into *E. coli* TOP10 cells and plated on LB medium with 0.4% glucose and appropriate antibiotics. Single colonies were grown to saturation overnight in M9L medium with 0.4% glucose and antibiotics. The following morning, cultures were serially diluted and spotted onto M9L plates supplemented with antibiotics and 0.4% glucose, 0.2% arabinose, or 0.2% arabinose and 100 µM IPTG. Plates were then incubated at 37°C for 24 hours. Positive interactions yielded single colonies on M9L with 0.2% arabinose and 100 µM IPTG after 24 hours of growth. Intermediate interactions yielded modest growth on plates but no visible single colonies. No intermediate growth phenotypes were observed for the 20x20 matrix (Fig. 2B).

ParD3-ParE3 expression and purification

Recombinant *Mesorhizobium opportunistum* ParDE3 protein complex was expressed in *E. coli* Rosetta(DE3)pLysS (Novagen). A 50 mL overnight culture in LB medium supplemented with 50 µg/ml kanamycin (LB-Kan50) was used to inoculate 2 L of LB-Kan50; this culture was incubated at 37°C in a rotary shaker at 220 rpm. Transcription of recombinant *parDE3*
was induced at an OD$_{660}$ of 0.8 by adding 1 mM isopropyl β-D-1-thiogalactopyranoside (IPTG). After 4 h of induction, the cells were harvested by centrifugation at 12,000g for 20 min at 4°C. Cell pellets were resuspended in 30 ml of lysing/binding buffer (10 mM Tris (pH 7.4), 150 mM NaCl, 10 mM imidazole with 5 μg/ml of DNase I (Sigma-Aldrich) and half a tablet of cOmplet EDTA free protease inhibitor cocktail (Roche Life Science). Cells were disrupted by one passage through an LV1 microfluidizer (Microfluidics, Westwood, MA) and the cell debris was removed by centrifugation for 20 min at 25,000 g. The supernatant was loaded onto a Ni$^{2+}$ Sepharose affinity column (GE Life Sciences) pre-equilibrated with the binding buffer. Two washing steps were performed using 10 mM and 75 mM of imidazole followed by two elution steps with 200 mM and 1 M imidazole in the binding buffer. After purity of the different fractions was assessed by SDS-PAGE, the protein solution was dialyzed against 10 mM Tris (pH 7.4), 150 mM NaCl, 200 mM imidazole buffer.

**Crystallization of ParD3-ParE3**

Purified ParDE3 was purified and concentrated using a centrifugal filter (3 kDa MWCO, Amicon-Millipore). Protein purity was estimated to be 95% as assessed by 14% SDS-PAGE stained with Coomassie brilliant blue. Initial crystallization screening was carried out using the sitting-drop, vapor-diffusion technique in 96-well microplates (Nunc). Trays were set up using a Mosquito robot (TTP LabTech) and commercial crystallization kits (Nextal-Qiagen). The drops were set up by mixing equal volumes (0.1 μl) of the protein and the precipitant solutions equilibrated against 75 μl of the precipitant solution. In all trials, the protein concentration was ~ 40 mg/mL. In approximately five days, needle-like crystals
appeared in condition 15 of the Pro-complex Suite crystallization kit (Qiagen). After manual refinement of the crystallization condition, the best crystals were obtained at 19°C with the following crystallization solution: 400 mM Sodium Acetate, 100 mM Sodium Citrate pH5.5, 20% PEG 4000, 20 % glycerol. All manual crystallization attempts were carried out using the hanging-drop, vapor-diffusion technique in 24-well plates (Hampton). Prior to flash freezing in liquid nitrogen, drops containing the crystals were mixed with 1 µl of a crystallization solution containing 100 mM sodium iodide and incubated for 4 hours. Crystals were then cryo-protected by soaking them in the crystallization solution containing 25% glycerol and 100 mM sodium iodide.

**Crystallographic data collection and data processing**

Crystal diffraction was measured at a temperature of 100 K using a 1 degree oscillation range on beamline 21-ID-D (LS-CAT, Advanced Photon Source, Argonne, Illinois); diffraction images were collected on a MAR Mosaic 300 detector. Diffraction images were processed using the Xia². Geometric refinement and examination of the scaled amplitudes revealed that the ParDE3 crystals belong to orthorhombic space group I222, with cell dimensions a=43.18, b=118.84, c=211.42 (α=β=γ=90˚) (see Table S1).

Diffraction from a single ParDE3 protein crystal was measured to 1.53 Å at an energy of 12.66 keV (0.979 Å). The anomalous signal in the data was used to locate iodide atoms in the lattice, and the structure was phased by single wavelength anomalous dispersion (Dauter, 2002) using the Autosol SAD routine in Phenix (Adams et al., 2010). Two ParDE3 complexes are present in the asymmetric unit. Eight iodine sites were located within the asymmetric unit. A preliminary ParDE3 structural model was built de novo from the initial
experimental, solvent-flattened maps using the AutoBuild routine and phenix.refine. This initial model was then manually examined and corrected; solvent addition and refinement of the structure was conducted iteratively using Coot (Emsley and Cowtan, 2004) and phenix.refine (Adams et al., 2010). The final structural model was refined to an R<sub>work</sub> of 16.85% and R<sub>free</sub> of 19.54%. Coordinates of ParDE3 were deposited in the Protein Data Bank (PDB ID 5CEG). Crystallographic data and refined model statistics are in Table S1.

**Size exclusion chromatography**

A purified sample of ParDE3 (10 mg/ml-300 µl) was injected on a GE Healthcare Superdex 200 10/300 GL column (flow rate 0.5 ml/min) and fractions of 500 µl were collected. 10 mM Tris pH 7.4, 150 mM NaCl, 200 mM imidazole was used as a running buffer. Collected fractions were resolved on 14% SDS-PAGE gels and compared to the elution profile. To estimate the molecular weight and, hence, oligomeric state of the ParDE3 complex in solution, its elution volume was compared to molecular weight standards (blue dextan, aldolase, conalbumin and ovalbumin) resolved on the same column using the same buffer and flow protocol.

**ParD3 library construction**

Residues incorporated at each library position were chosen to closely resemble that of naturally occurring ParD homologs. Briefly, the software HMMER was used to identify and align homologs of *C. crescentus* ParD3 using an E-value cutoff of 0.0001, and then sequences greater than 95% identical were removed. For each library position, amino acid frequencies were extracted from the curated ParD3 alignment, and a library residue set was chosen that covered at least 95% of the sequence diversity in the ParD3 alignment.
This approach yielded a total of 12, 6, 13, and 10 residues at the four positions of the library.

The ParD3 library was generated using the ProxiMAX technique (Ashraf et al., 2013). The main advantage of this technique over traditional NNS libraries is that the amino acid composition at each position in the library can be specified, thus reducing the complexity of the library. To generate the ParD3 library, we began with an “acceptor” fragment that contained the region of parD3 upstream of the first position in the library (L59) and different hairpin “donor” fragments that contain (i) one of the codons we wish the incorporate and (ii) a downstream MlyI restriction site. We set up separate blunt-end ligations between the acceptor fragment (0.1 μM) and each codon donor fragment (0.5 μM) using T4 ligase and incubated at 22°C for 1 hour. These ligations were diluted 1:100 in water and used as a template for separate high-fidelity PCR (Phusion) using primers specific for the acceptor and donor fragments. These PCR reactions were gel purified, quantified, and then pooled in equimolar amounts. The pooled mixture was then cut using MlyI and PCR purified, resulting in a new acceptor fragment that contains the L59 position randomized. This acceptor fragment was then used for three additional rounds of ProxiMAX randomization as outlined above, to create a library in positions L59, W60, D61, and K64 in ParD3. The final fragment was sub-cloned into the Sacl and PvuI sites of pEXT20 and library composition was verified by Illumina sequencing of the relevant region of parD3.
Illumina-based sequencing and fitness calculations

Plasmid DNA was extracted from frozen cell samples (Qiagen) and used as a template for PCR reactions (20 cycles) with custom barcoded primers containing Illumina flowcell adaptor sequences. The samples were multiplexed and run on an Illumina HiSeq instrument. Multiplexed Illumina reads from a single lane were sorted based on an exact match to a four-letter barcode sequence. Reads were then filtered to remove sequences that (a) contained frameshift mutations, (b) encoded for a parD3 variant not in the planned library, or (c) lacked an exact match to six nucleotides before (AGGCAG) and after (GCAAGC) the randomized region. Sequences that passed these quality filters were then counted and frequency-normalized. We calculated the fitness of each variant as described previously (van Opijnen et al., 2009). Briefly, we generated a linear fit to the frequencies of each mutant as a function of time, and then calculated the log-fold expansion of each mutant relative to the rest of the population, yielding $W_{raw}$ for each variant:

$$W_i = \frac{\log(E(t_1/t_0))}{\log(E(1-t_1)/(1-t_0))}$$

where $t_0$ is the frequency of the mutant at 200 min, $t_1$ is the frequency of the mutant at 600 min, and $E$ is the expansion factor of the culture (OD at $t_0$ / OD at $t_1$). We then transformed these raw fitness values such that the $W$ value for frameshift variants was 0 and the $W$ value for the wild-type (LWDK) sequence was 1.
Creation of the orthogonal ParE3* toxin

To create a ParE3 toxin with a novel specificity profile, we focused on residues in ParE3 that covary with W60/D61 from ParD3. We found that residues R58/A61/L72 in ParE3 covary with W60/D61 from ParD3 with a GREMLIN scaled score greater than 1. We then searched for residues within ParE3 that covary with R58/A61/L72 (termed “supporting residues”) with a GREMLIN scaled score greater than 1. Repeating this search process iteratively produced two more supporting residues in ParE3, M63/R54, for a total of five specificity and supporting residues in ParE3: R54/R58/A61/M63/L72.

To identify which mutations to make in these five residues, we searched naturally existing ParE sequences for combinations of residues that often occur at these positions. We chose to incorporate the residue combination VEIRF because each residue was commonly observed at the corresponding position in naturally existing ParE sequences and because it was chemically dissimilar to the wild-type residue at that position, RRAML.

Generation of force-directed graphs

Graphs were generated using networkx and visualized using Gephi (Jacomy et al., 2014). For the graph in Figure 5C, edges were drawn for every two ParD3 variants that are separated by a single amino acid substitution; for the graph in Figure S5B, edges were drawn for every two ParD3 variants that are separated by single nucleotide substitutions. Nodes were colored based on specificity class using custom-written Python scripts. The layout was generated using the Force Atlas algorithm to completion.
Table S1. Data collection and refinement statistics; Related to Figure 3.

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<th>Description</th>
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<td>Unit cell</td>
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<td>Average B-factor</td>
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Statistics for the highest-resolution shell are shown in parentheses.
Supplemental References


