Evolutionary history expands the range of signaling interactions in hybrid multikinase networks

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Two-component systems (TCSs) are ubiquitous signaling pathways, typically comprising a sensory histidine kinase (HK) and a response regulator, which communicate via intermolecular kinase-to-receiver domain phosphotransfer. Hybrid HKs constitute non-canonical TCS signaling pathways, with transmitter and receiver domains within a single protein communicating via intramolecular phosphotransfer. Here, we report how evolutionary relationships between hybrid HKs can be used as predictors of potential intermolecular and intramolecular interactions ('phylogenetic promiscuity'). We used domain-swap genes chimeras to investigate the specificity of phosphotransfer within hybrid HKs of the GacS–GacA multikinase network of Pseudomonas brassicacearum. The receiver domain of GacS was replaced with those from nine donor hybrid HKs. Three chimeras with receivers from other hybrid HKs demonstrated correct functioning through complementation of a gacS mutant, which was dependent on strains having a functional gacA. Formation of functional chimeras was predictable on the basis of evolutionary heritage, and raises the possibility that HKs sharing a common ancestor with GacS might remain components of the contemporary GacS network. The results also demonstrate that understanding the evolutionary heritage of signaling domains in sophisticated networks allows their rational rewiring by simple domain transplantation, with implications for the creation of designer networks and inference of functional interactions.

Two-component signal transduction systems (TCSs) are mainly found in prokaryotes, where they control diverse and important adaptive responses to changes in environmental conditions1. The prototypical TCS comprises a pair of signal transduction proteins—a histidine kinase (HK) and cognate response regulator (RR)—both of which are multi-domain proteins. HKs contain sensor domains (input), which upon stimulation activate autophosphorylation of a histidine residue in the HK transmitter domain. The phosphorylated transmitter domain then binds to the receiver domain of the RR partner, leading to transfer of the phosphoryl group to an aspartate residue of the receiver domain. In the majority of cases, the RR phosphorylation changes the activity of an effector domain (output) within the RR, which brings about a physiological change, often through altered expression of specific genes. A subset of TCSs are non-prototypical systems, which include a class of TCSs designated ‘hybrid’ HKs2. Hybrid HKs have both transmitter and receiver domains, but variants are commonly found which possess additional signaling domains, for instance the ‘unorthodox’ sub-class of hybrid HKs have an additional histidine-containing phosphotransfer (HPt) domain.

The TCS gene sets in an organism can expand by duplication, divergence and lateral transfer, resulting in some species possessing several hundreds of such signaling proteins. This raises the question of how specificity is maintained between cognate partners despite the structural similarity common to all TCS proteins. It has been shown that for prototypical TCSs, the interaction between cognate protein partners is highly specific, preventing unwanted cross-talk between TCSs3–5. However, domain-domain interactions within hybrid HKs have relaxed domain-domain specificity and affinity compared to the interprotein interactions of prototypical HKs and RRs.
Relaxed specificity is tolerable in hybrid HKs because the tethering of transmitter and receiver domains within a single protein increases the effective concentration of each domain in the vicinity of the other.  

Although prototypical TCS protein partners encoded by adjacent genes are frequent, isolated HK and RR genes (orphans) are also abundant, as are complex gene loci encoding multiple ‘TCS proteins’ 2. This makes the identification of signaling partnerships non-trivial. The P2CS database (http://www.p2cs.org) provides paired HK and RR phylogenetic trees, which shed light on evolutionary heritage and can additionally provide clues about protein HK–RR partnerships 15. Indeed, HKs and RRs that are known to interact, fall into corresponding subfamilies 10–12. This phylogenetic relationship is mainly attributed to the co-evolution of the interacting domains of TCS partners 13 and the need to prevent unwanted cross-talk between non-cognate TCs.  

Nevertheless, because of their shared evolutionary heritage, non-cognate interactions are possible between phylogenetically-related TCS domains 14–17. Sometimes, observed ‘non-cognate’ interactions turn out not to be unwanted cross-talk, but rather are phenotypically important ‘cross-communication’ between TCS, forming signaling networks 14,18,19. Phylogenetic analysis indicated that there was no common ancestor for hybrid HKs, and that their origin and expansion were achieved by lateral recruitment of a receiver domain into an HK molecule and then duplication as one unit 20.  

The majority of hybrid HK network interactions described in the literature concern the GacS-GacA TCS in the model bacterial species Pseudomonas aeruginosa, an opportunistic human pathogen causing respiratory infections of hospitalized patients. GacS is an unorthodox HK, which phosphorylates the response regulator GacA. GacA and GacS are part of a multikinase network, in which the LadS and RetS hybrid HKs affect the activation of GacA by GacS 21–23. The GacS-GacA TCS network and additional hybrid HKs (PA1611, PA1976 and PA2824) control the expression of small regulatory RNAs (sRNAs), which orchestrate several functions in gammaproteobacteria 21–23. In P. aeruginosa, phosphorylated GacA specifically and exclusively activates the expression of four sRNA genes (rsrX, rsrY and rsmZ), whereas in P. fluorescens CHA0, phosphorylated GacA activates the expression of four sRNA genes, rsmX, rsmY, rsmZ, and mga 24,25. The deletion of rsmY and rsmZ genes in P. aeruginosa 24 and of rsmX, rsmY, and rsmZ in P. fluorescens CHA0 25 result in phenotypes that are similar to those of gacS and gacA mutants.  

Pseudomonas brassicaeae was isolated as the major root-associated bacterial species in the rhizosphere of Arabidopsis thaliana and Brassica napus 26. P. brassicaeae has the ability to suppress plant pathogens by producing antifungal compounds, such as 2,4-diacyltrehaloside and cyaniode. The two-component GacS-GacA system by activating the expression of three small regulatory RNAs (RsmX, RsmY, RsmZ), positively controls the expression of secondary metabolites (e.g. the antifungal compounds DAPG and cyaniode), plant hormones (e.g. auxin), exoenzymes (e.g. lipase and protease) 24, AHLs, the type VI secretion machinery, alginate synthesis, and biofilm-forming ability 24. Mutations in the gacS-gacA system result in drastic pleiotropic changes in P. brassicaeae 25. This bacterial species combines effective plant colonization 27,28 with other beneficial attributes for plant health and growth, including release of plant growth hormones (such as indole-3-acetic acid) 29, modulation of ethylene by the enzyme 1-aminoacyclopropane-1-carboxylate deaminase 30, production of two different siderophores 31 and protection against phytopathogens by producing at least two antifungal compounds DAPG and cyanide 25. The genome of P. brassicaeae contains homologues of the P. aeruginosa hybrid HKs, making it a relevant model for deciphering the GacS-GacA network.  

Here, we report how the evolutionary relationships between hybrid HKs can be used as predictors of potential intermolecular interactions. We used domain-swap gene chimera of P. brassicaeae Gacs to assay interactions between signaling domains of hybrid HKs predicted by their evolutionary relationships. As well as showing the utility of the ‘phylogenetic promiscuity’ approach, we propose that additional hybrid HKs might be involved in the Gacs cascade that controls phytobeneficial traits, and suggest likely new signaling interactions within the GacS multikinase network. This approach also allows us to suggest potential intermolecular hybrid HK interactions in other bacteria using phylogenetic analyses available through the P2CS database.  

Results  

Kinase and receiver domains of ‘GacS cluster’ hybrid HKs have co-evolved. In the latest version of the P2CS database, paired phylogenetic trees were implemented for evolutionary analysis and identification of candidate partners for orphan TCS proteins 16. The genome of P. brassicaeae NFM421 30 was added to P2CS and analyzed to produce trees of kinase and receiver domains from encoded TCS proteins (Supplementary Fig. S1). The kinase domain of P. brassicaeae Psebr_a4261 (Gacs) was found in a cluster (the GacS kinase cluster) alongside seven other hybrid HKs: Psebr_a4938 (LadS), Psebr_a625 (RetS), Psebr_a1633, Psebr_a1408, Psebr_a3082 and Psebr_a4122 (PA1611) (Fig. 1, protein identifiers are abbreviated in the figure and in the rest of the manuscript). These seven proteins also cluster together in the P. brassicaeae receptor domain tree (the Gacs receiver cluster), which also includes the receiver domain of 3692 (Fig. 1), another hybrid HK which is encoded adjacent to a further receiver domain-containing protein (the LuxR-family RR, 3693). Protein 3692 is also included, if we slightly extend domain kinase cluster of GacS (Supplementary Fig. S1). Two hybrid HKs include more than one receiver domain, RetS contains two receivers (625_1 and 625_2) and 3082 contains three. The receiver domains 625_1, 3082_1 and 3082_2 do not belong to the GacS cluster. Homologues of the P. brassicaeae hybrid HKs are found in the P. aeruginosa genome (4261 = Gacs, 4938 = LadS, 625 = RetS, 1633 = PA2824 and 4122 = PA1611) (Fig. 2).  

We further applied Direct Coupling Analysis (DCA), a statistical inference method based on coevolution 13,22,23, to compute the interaction scores (specifically, effective interaction energies 8) between the P. brassicaeae Gacs HK domain and all the receiver domains involved in P. brassicaeae hybrid HKs. Small values of this interaction score have proved successful at predicting the correct interaction partners among prototypical cognate HK–RR pairs, through an iterative pairing algorithm (IPA) which starts from unpaired sequences only 24.
In Table 1, we list the results obtained for all the receiver domains considered in our experimental study. They are sorted from smallest to largest interaction score, and we provide their rank among all the receiver domains involved in \( P. \) brassicacearum hybrid HKs, as well as the interaction score values, and the score gap with the next ranked one. We also indicated the standard deviation on the interaction scores, and the reliability of our ranking, both obtained by bootstrapping. We found that the top predicted partner for the GacS HK domain is the GacS receiver domain, with a gap substantially larger than the standard deviation, and a reliability equal to 1 (which is the maximum possible). The next predicted potential partners of the GacS HK domain are the receiver domains from 4122, 3082 and 1633. Gaps between their interaction scores are smaller than standard deviations, and accordingly, ranking reliabilities are smaller than 1. In contrast, the gap between the last of these three domains (1633) and the next ranked one (625_2, second receiver of RetS) is substantially larger than standard deviations. In summary, this analysis predicts that the three top potential cross-talk partners for the GacS HK domain among all the receiver domains involved in \( P. \) brassicacearum hybrid HKs, are the receiver domains of 4122, 3082 and 1633. While this approach cannot order them reliably, it predicts that they are consistently better candidates than all other ones. These results are consistent with the phylogeny-based analysis.

Sequence analysis of the receiver and kinase domains shows amino acid residues common to all hybrid HKs of the GacS cluster, including phosphorylation sites: aspartic acid and histidine residues. Only one residue (asparagine) is common and exclusive to the group of receivers consisting of GacS and the three phylogenetically closest HKs (Supplementary Fig. S2). Coevolution analysis has shown that hybrid HKs do not exhibit the same extended amino acid coevolution between kinase and receiver domains as canonical kinase–receiver pairs. Collectively, these results suggest the seven hybrid HKs of the GacS kinase cluster share a common evolutionary history (recently duplicated from a common ancestor), and are therefore potentially able to engage in cross-talk/cross-communication with each other. To explore potential phosphotransfer interactions between the different hybrid HKs of the GacS kinase cluster, we focused experimentally on the receiver domains of the hybrid HKs, creating chimeric proteins by replacing the GacS native receiver domain with receiver domains from each of the GacS receiver cluster hybrid HKs.
Phylogenetic relationships mirror the potential for intramolecular phosphotransfer in hybrid HKs. Nine chimeric GacS hybrid kinases were constructed, seven of which had the GacS receiver domain swapped with other receiver domains from the GacS receiver cluster. The other two chimeras were engineered to contain receiver domains from outside the GacS receiver cluster—the receiver of 2650 (a hybrid HK from outside the GacS cluster) and the first receiver domain of RetS (625_1), only the second receiver domain of which (625_2) belongs to the GacS receiver cluster (Fig. 1). In all chimeras the C-terminal GacS HPt domain was retained, potentially allowing phosphotransfer to GacA when the transplanted receiver domain was able to substitute functionally for the original GacS receiver domain. Expressing the chimeras in a \( gacS \) mutant and monitoring GacA-dependent phenotypes (protease activity and colony size) thus provided a read-out of whether the chimeric HK was functional.

Table 2 shows the phenotypes of wild-type \( P. \) brassicacearum, the \( gacS \) mutant and the \( gacS \) mutant strain complemented with \( gacS \). The wild-type and complemented \( gacS \) strains showed protease activity and small non-fluorescent colonies, while the \( gacS \) mutant exhibited no protease activity and large fluorescent colonies due to enhanced pyoverdine production. Three chimeras (containing receiver domains from the three hybrid HKs closest to GacS in the receiver cluster—1633, 3082_3 and 4122) successfully restored wild-type phenotypes, with the exception of the colony size for chimera 1633. In contrast, chimeras harboring receiver domains from

| Receiver domain | Interaction score | Standard deviation | Gap | Reliability |
|-----------------|------------------|--------------------|-----|-------------|
| 4261            | -275.5           | 5                  | 22.9| 1           |
| 4122            | -252.6           | 5.2                | 4.2 | 0.7         |
| 3082_3          | -248.4           | 5.4                | 1.5 | 0.4         |
| 1633            | -246.9           | 5.1                | 18.7| 0.4         |
| 625_2           | -228.2           | 5.4                | 0.2 | 0.2         |
| 2955            | -228             | 5                  | 0.8 | 0.1         |
| 3905            | -227.2           | 4.8                | 12.2| 0.1         |
| 1657            | -215             | 4.9                | 1.2 | 0           |
| 4938            | -213.8           | 5.1                | 0.1 | 0           |
| 3692            | -213.7           | 5                  | 9.1 | 0           |
| 5306            | -204.6           | 5                  | 0.6 | 0           |
| 1408            | -204             | 5                  | 0.7 | 0           |
| 2831            | -203.3           | 4.6                | 3.6 | 0           |
| 3973            | -199.8           | 4.4                | 4.7 | 0           |
| 1484            | -195.1           | 4.9                | 2.4 | 0           |
| 900             | -192.7           | 5.3                | 0.4 | 0           |
| 625_1           | -192.3           | 5.3                | 0.1 | 0           |

Table 1. Direct coupling analysis. The results show the interaction scores (effective interaction energies) with the kinase domain of GacS. All the receiver domains involved in \( P. \) brassicacearum hybrid HKs were considered, and ranked from smallest to largest interaction scores. The gap corresponds to the difference in interaction score between two successive ranks. The reliability score represents the fraction of bootstrapping replicates where the ranking down to this domain remained identical to the one obtained without bootstrapping (see “Materials and Methods”). The receiver domains belonging to the GacS cluster are marked in bold.

| Strains          | Protease activity | Colony size |
|------------------|-------------------|-------------|
| Wild-type        | +                 | Small       |
| \( gacS \)       | -                 | Large       |
| \( gacS-gacS^* \)| +                 | Small       |
| \( gacS-625_1^* \)| -                 | Large       |
| \( gacS-625_2^* \)| -                 | Large       |
| \( gacS-1408^* \)| -                 | Large       |
| \( gacS-1633^* \)| +                 | Large       |
| \( gacS-2650^* \)| -                 | Large       |
| \( gacS-3082_3^* \)| +               | Small       |
| \( gacS-3692^* \)| -                 | Large       |
| \( gacS-4122^* \)| +                 | Small       |
| \( gacS-4938^* \)| -                 | Large       |
RetS-625, 1408, 3692 and LadS-4938 displayed gacS mutant phenotypes (Table 2). The fact that only three out of seven chimeras whose receiver is phylogenetically close to that of GacS lead to restoration of wild-type strain phenotypes, suggests that the observed phenotypes are not only the result of the high local concentration but also specificity, because four chimeras were ineffective despite their local concentrations likely being high (although we cannot exclude the possibility of misfolding or a high level of dephosphorylation for non-functional chimeras). Taken together, these results suggest that the most phylogenetically similar domains retain the capacity for intramolecular phosphotransfer, a principle we call ‘phylogenetic promiscuity’.

GacS chimeric proteins are capable of both intra- and intermolecular signal transduction. As an unorthodox hybrid HK, phosphorylated GacS transfers phosphoryl group intramolecularly to its C-terminal HPt domain, and then intermolecularly to its cognate response regulator GacA. To determine whether this still happens for chimeras that completely (3082, 4122) or partially (1633) restored wild-type phenotypes to the gacS mutant (Table 2), GacA-dependence was tested by expressing each of the three chimeras in a gacA mutant. This resulted in phenotypes comparable to the control gacA mutant (Fig. 3), demonstrating that phosphotransfer to GacA is required for the 1633, 3082 and 4122 chimeras to restore wild-type phenotypes to a gacS mutant.

As the expression of rsmX is completely controlled by the GacS-GacA system23, we examined its expression, by quantitative reverse transcription-PCR (qRT-PCR). The rsmX expression of the mutant gacS harbouring each of the seven chimeras belonging to the GacS cluster was compared to that of the wild-type and the gacS mutant. The expression of rsmX is significantly higher in the wild-type and 3082–4122 chimeras, than in the gacS mutant strain and the four remaining chimeras including 1633, which only partially restored the wild-type phenotypes (Fig. 4, Supplementary Fig. S3).

**Figure 3.** GacA is required for GacS chimeras to restore wild-type phenotypes. Lack of restoration of the wild-type strain phenotypes by expression of GacS chimeras in the gacA mutant. Protease activity (lysis zone) on skim milk medium (A) and colony appearance on PAF medium (B).

**Figure 4.** rsmX expression levels. Relative expression of rsmX for the wild-type strain, gacS mutant and gacS mutant complemented with one of seven chimeras belonging to the GacS cluster. The reported values represent the means of three experiments, with standard errors indicated by error bars. All data are presented as a percentage relative to the wild-type. Welch’s heteroscedastic F test was performed and * indicates significantly different comparisons (p < 0.05) to the wild-type.
Potentially, GacS chimeric proteins might not be fully functional, interacting with, and receiving phosphoryl group from the hybrid HK that had donated the receiver domain to the chimera. This possibility was tested for 3082 by introducing the GacS-3082 chimera into the gacS—3082 double mutant, which restored wild-type phenotypes (Fig. 5).

These observations suggest that functional GacS chimeras are able to perform both the intramolecular phosphotransfer reactions (from kinase to receiver to HPt domains) and also the intermolecular phosphotransfer to GacA, as performed by wild-type GacS.

Transplanted receiver domains in GacS chimeric proteins can be phosphorylated by donor hybrid HKs. Additionally, it is possible that phosphorylation of the transplanted receiver domain in a GacS chimera also occurs via cross-talk from another TCS (as indeed the native GacS receiver domain can). To test this possibility, we generated a 'kinase-dead' GacS-3082 chimera in which the phosphoaccepting histidine residue (H294) in the kinase domain was substituted by alanine, preventing autophosphorylation. Expression of this construct in a gacS mutant resulted in restoration of wild-type phenotypes (Fig. 6), suggesting cross-talk between the GacS chimera and another TCS protein. Indeed, expression of the GacSH294A-3082 chimera in the gacS—3082 double mutant did not restore wild-type phenotypes (Fig. 6), implying that it is 3082, which transfers phosphoryl group to GacS-3082 chimeras.

**Figure 5.** GacS chimeras perform both intra- and intermolecular phosphotransfer. Restoration of the wild-type strain phenotypes by expression of 3082_3+ chimera in the gacS—3082 double mutant, compared to the gacS—3082 double mutant. Protease activity on skim milk medium (A) and colony appearance on PAF medium (B).

**Figure 6.** Transplanted receiver domain in a GacS chimera also occurs via cross-talk. Restoration of the wild-type strain phenotypes by expression of GacSH294A—3082_3+ chimera in the gacS mutant and lack of restoration in the gacS—3082 double mutant. Protease activity on skim milk medium (A) and colony appearance on PAF medium (B). The phenotypes of the gacS mutant and the gacS—3082 double mutant are also shown.
To test whether such cross-talk involves the aspartate residue of the transplanted receiver domain and/or the histidine residue of the GacS HPt domain, a mutant version of the GacS-3082 was engineered in which the aspartate residue in position 718 was substituted with an alanine residue. The GacS-3082D718A construct did not restore wild-type phenotypes to the \( gacS \) mutant (Fig. 7), likely indicating that the GacS-3082_3 chimera receives phosphoryl group onto the transplanted receiver domain from the 3082 HK.

These observations raised the question of whether the 3082 kinase is involved in the GacS–GacA system, under physiological conditions. To examine this possibility, we constructed a 3082 mutant, which displayed the same phenotypes as the wild-type. This is in accordance with the work of Lee et al. on \( Pseudomonas \) \textit{alkyphenolica}, which showed that the expression of \textit{rsmZ}, one of the final targets of the GacS-GacA system, was not significantly altered in a \( bmsA \) (orthologous to the 3082 gene) mutant, and overexpression of \( bmsA \) in a \( gacA \) mutant did not produce the wild-type phenotype. This does not exclude the possibility that the 3082 gene, and by extension \( 1633 \) and \( 4122 \) genes, participates indirectly to the GacS-GacA TCS. Indeed, the genomic context of the 3082 gene, for instance, includes three genes encoding CheR (3081), CheB (3080) and a hybrid histidine kinase (3079). This cluster, as well the GacS-GacA system, have been shown to be involved in common processes such as biofilm formation and motility within the \( Pseudomonas \) \textit{genus}, probably through common mechanisms that have yet to be elucidated.

**Discussion**

**Phylogenetic promiscuity.** Evolutionary analysis suggested that hybrid HKs were not derived from a common ancestor, and their origin and expansion were achieved by lateral recruitment of a receiver domain into an HK molecule and then duplication as one unit. For example, a comparative study of six species of \textit{Xanthomonas} have shown that the individual domains of a hybrid histidine kinase in one species, originated directly from a gene fusion event that combined cognate HK and RR from closely related species. Such event likely occurs through a mutation of the stop codon, resulting in read-through of the open reading frame. In comparison to orthodox HKs, hybrid HKs have special evolutionary characters, manifested by a higher level of DNA polymorphism and faster evolutionary rate, and frequent gene fusion or fission and duplication.

Consequently, the expansion of hybrid HKs by gene duplication raises the question of their potential cross-talk. Although available data indicate that cross-talk between TCSs generally decreases TCS signaling activity, it cannot be ruled out that such systems could tolerate the presence of more than one interaction partner. It has been postulated that phylogenetic proximity and the sequence similarity of hybrid HKs are elements that may indicate the degree to which a set of HKs cross-talk or not. The squid symbiont \textit{Vibrio fischeri} uses a signaling network composed of two interacting hybrid HKs to promote biofilm formation and colonization. The receiver domain of hybrid HK RscS and the HPt domain of hybrid HK SyrP interact to phosphorylate two downstream RRs. A second example shows intra- and interprotein phosphotransfer between two hybrid HKs to control the developmental program in \textit{Myxococcus xanthus}. Other examples of interacting hybrid HKs have been described in \textit{Pseudomonas} spp., including the RcsC/PvrS and GacS multikinase networks. Through GacA activity, the GacS network controls the transcription of several sRNAs, which orchestrate diverse functions in gammaproteobacteria. In all these cases, phylogenetic trees of hybrid HK kinase and receiver domains in
the P2CS database show that interacting hybrid HKs fall within the same phylogenetic clusters (Supplementary Fig. S4).

In *P. brassicacearum*, similar phylogenetic clustering of hybrid HK kinase and receiver domains is observed, suggesting that they may interact with each other in a multikinase network.

**Interconnection of hybrid HKs.** Current knowledge of hybrid HKs stipulates that a spatial tethering of kinases to their substrates relaxes evolutionary constraints on specificity4–8, implying that a covalently linked receiver domain can be functional even in the absence of specific interactions. Nevertheless, using chimeras transplanted into GacS with receiver domains from donor hybrid HKs, we showed that phylogenetic proximity reflects whether a transplant retains biological activity, suggesting interactions were specific (although it is possible non-functional chimeras were misfolded or had high dephosphorylation rates). Interestingly, identification of transmitter–receiver partnerships was more successful for hybrid HKs if domains from hybrid HKs were used as the training set, than if domains from prototypical TCS were used (see “Materials and Methods”). This also suggests that intramolecular interactions within hybrid HKs require specificity, not just tethering of domains to one another.

In addition, the expression of chimeras (with and without mutated kinase domains), in single and double mutant backgrounds, demonstrated that the 3082 third receiver domain can receive signal from both the GacS and 3082 kinase domains, i.e. both intra- and intermolecularly. These observations suggest that 1633, 3082 and 4122 are likely targets to be phosphorylated by GacS in vivo.

The experiments described in this work investigated flow of signal through the GacS-GacA TCS with domains transplanted into GacS from other TCS proteins. The use of chimeras in mutant backgrounds precluded us from directly defining which/whether specific phosphotransfer interactions occur within/between wild-type proteins. However, in our experiments the signaling domains under study were maintained in the same protein context in which they are found in the wild-type proteins, and the use of chimeras allowed assessment of physiological phenotypes which bypassed many of the problems associated with in vitro studies involving the phosphorylation of purified proteins40.

In *P. aeruginosa*, the hybrid HK RetS attenuates GacS signaling via three mechanisms, including phosphotransfer from the GacS kinase to the second receiver (Rec2) domain of RetS19. However, in *P. brassicacearum*, a GacS chimera carrying the Rec2 of RetS (625_2), does not restore wild-type phenotypes to a gacS mutant (Table 2). This can be explained by the difference in phylogenetic distance between the two sets of receiver domains. While in *P. brassicacearum* the receivers of the two proteins are in two distinct clusters (Fig. 1), the Rec2 domain of RetS in *P. aeruginosa* is the closest receiver to that of GacS (Supplementary Fig. S4B).

The avoidance of unwanted cross-talk is a major selective pressure for classical HK-RR two-component pathways, driving the diversification of specificity residues to produce post-duplication insulated pathways41. However, duplicate hybrid HKs such as those exemplified by the GacS multikinase network, converge towards a common signal output and biological response functions. This convergence allows horizontal interconnection of hybrid HKs, presumably producing sophisticated decision-making networks able to assimilate multiple sensory inputs into a single response42. It is also possible that additionally non-hybrid HKs may influence signaling through the network.

It should be noted that the GacS network comprises unusual hybrid sensor kinases that harbor more than one receiver domain. RetS contains two receiver domains, while 3082 contains three. In each hybrid HK, only one of the receiver domains phylogenetically clusters with GacS, or can function within GacS chimeras, implying separate roles and interaction partners for the different receiver domains, as seen for hybrid HKs in similar and diverse organisms4,19,43. It is likely that these proteins have multifunctional roles, potentially monitoring multiple environmental stimuli.

**Multikinase network.** Through in silico prediction and the use of chimeras, we have identified three proteins (1633, 3082 and 4122) whose receiver domains function interchangeably with that of GacS, enabling signal transduction via GacA and resulting in transcriptional activation of *rsmX*. A receiver domain of one of these proteins, 3082_3, is shown to be able to receive signal (phosphoryl group) from its cognate kinase domain both intra- and intermolecularly, suggesting that 1633, 3082 and 4122 might all be involved in the GacS network in *P. brassicacearum*. Further experiments are needed to confirm whether this is the case in vivo, for example by identifying signals which stimulate 1633, 3082 and 4122 signaling and testing for a GacS response to those signals. Our ‘phylogenetic promiscuity’ approach can be used to suggest additional hybrid HKs that may be involved in multikinase networks from publicly available data in the P2CS database. For instance, the *P. aeruginosa* proteins PA_2824 and PA_3462 are likely to be involved in the GacS network44, the VF_A0072 protein of *V. fischeri* is likely to interact with the RscS-SypF network45, and we would also predict that MXAN_2386 and MXAN_0314 of *M. xanthus* are part of the EspB/EspC network44.

The use of evolutionary relationships to predict signaling interactions also implies that signaling relationships can act as indicators of evolutionary heritage. For instance, our results would support a model of GacS network evolution in *P. brassicacearum* by duplication of an ancestral GacS (producing the ancestors of 4122 and 1633) and recruitment of a copy of the GacS receiver domain by another hybrid HK (the ancestor of 3082).

In summary, our results show that the phylogenetic relationships between kinase and receiver domains can be used to guide the rational ‘rewiring’ of communications between and within the HKs forming multikinase networks. This opens possibilities for engineering sophisticated signaling networks by simple domain transplantation, as well as investigating novel functional associations.
Materials and methods
Bacterial strains, plasmids and growth conditions. The bacterial strains and plasmids used in this study are listed in Supplementary Table S1. P. brassicacearum NFM421 and its mutants were grown at 30 °C in tenfold-diluted tryptic soy broth (TSB 1/10) or in 20-fold-diluted tryptic soy broth (TSB 1/20). For detection of extracellular protease activity, bacteria were plated on TSB 1/10 agar plates containing 1% skimmed milk. Pseudomonas agar F (PAF) (Difco) was used to reveal colony morphology. Escherichia coli strains GM2163, TOP10, and DH5α were grown in lysogenic broth (LB) at 37 °C. For growth on plates, media were solidified with 15 g/l agar (Sigma).

Construction of mutants. Mutant of gacA was generated by deletion mutagenesis as previously described23 and gacS mutant was developed according to the same procedure, using primers listed in Supplementary Table S2. The mutants obtained by deletion were tested for their phenotypes and then complemented, confirming the expected phenotypes. To construct the 3082 mutant and the gacS—3082 double mutant, the plasmid pCM18445, unable to replicate in strain NFM421, was used. To generate a vector carrying only kanamycin resistance, pCM184-Km' was created by eliminating ampicillin and tetracycline resistances from pCM184 through Scal/XmnI and EcoRV/PshAl digestion and re-ligation of the blunt cuts. The plasmid pCM184-Km' was then engineered by inserting an internal fragment of the target gene. Integration of the resulting plasmid into the chromosomal copy disrupts the target gene leading to the production of truncated non-functional proteins. A 0.78 kb fragment, between positions 1190 and 1977 of the chromosomal copy disrupts the target gene leading to the production of truncated non-functional proteins.

GacS chimeras synthesis and expression. All chimeras consisted of full-length hybrid histidine kinases including transmembrane regions. The GeneArt service (Invitrogen) was used for construction of chimeric protein expression vectors. Briefly, synthetic gene chimeras were assembled from synthetic oligonucleotides and/or PCR products. Synthetic genes were inserted into pMA-RQ (ampR), and verified by sequencing before inserting the chimeric genes into the expression vector pME603246. Plasmids were transformed into E. coli strain GM2163 to avoid DNA methylation and then introduced into recipient P. brassicacearum strains by tri-partite conjugation (GM2163 with chimeric protein expression plasmids, helper DH5α pRK2013, and the P. brassicacearum strain). Potential mutants were obtained by growth on TSB 1/10 agar plates and screening for tetracycline (and kanamycin). Mutation of the 3082 gene was confirmed by PCR amplification between the plasmid Km' gene and the chromosomal regions flanking the 3082 gene, then sequencing the PCR amplification products.

Sequence analysis. The software MAFFT (http://mafft.cbrc.jp/alignment/software/) was used to align kinase and receiver domain sequences from HK proteins (http://www.p2cs.org). A threshold of 40% identity across domains was used to identify conserved amino acid residues. The multiple sequence alignment is then visualized using Jalview (https://www.jalview.org).

Phylogenetic analysis of kinase and receiver domains and interaction prediction. The generation of phylogenetic trees was carried out as previously described47. Briefly, the kinase and receiver domains of TCS proteins were identified as previously described47 and used to construct pairwise alignments using TULIP 1.548, which uses the Smith–Waterman algorithm, with 1000 sequence shuffles, to estimate pairwise Z-values and infer a distance matrix. In the R environment, a hierarchical cluster analysis was undertaken using this distance matrix and the hclust function with ‘ward’ as a linkage method. An in-house R function then plotted dendrograms to display the hierarchical relationship between domains.
To predict potential interaction partners among paralogs of hybrid HKs in *P. brassicacearum*, we built a direct coupling analysis (DCA) model for hybrid HK kinase and receiver domains using intra-protein pairs between such domains from the P2CS database. Unorthodox proteins were included, but the set of pairs used to construct the model was restricted to those proteins that comprise just one kinase domain and one receiver domain, in order to avoid any ambiguous cases. The resulting training set comprised 12,462 domain pairs. The mean-field approximation was employed to compute DCA couplings between amino acid sites9–11, and we used the same alignment methods and parameters as in12. Note that in principle we could have used a training set that also included prototypical cognate HK-RR pairs, or a training set based only on prototypical cognate HK-RR pairs, as described previously13. However, we found that such a model was less good at predicting intra-protein domain pairs from hybrid HKs than the model constructed using hybrid domains only (the former predicts 29% of all true positives, compared to 96% for the latter), which presumably reflects differences in composition and coevolution between prototypical and hybrid TCSs. The couplings obtained from our DCA model were then employed to compute interaction scores (effective interaction energies) between the *P. brassicacearum* GacS kinase domain and all the receiver domains involved in *P. brassicacearum* hybrid HKs, as previously defined14. All the hybrid HKs included in our study were predicted with a ‘response_reg’ receiver domain, except for HK 2650 which carries a REC domain (data available on www.p2cs.org). This precluded its inclusion in the DCA analysis. To assess the statistical significance of the computed scores, we performed 1000 bootstrapping replicates of the predictions by randomly resampling sequences from the training set (with replacement). This allowed us to compute the variance of each interaction score. For each receiver domain ranked by increasing interaction score with the GacS kinase domain, we further calculated the fraction of bootstrapping replicates where the ranking down to this domain remained identical to the one obtained without bootstrapping. This fraction is indicative of the reproducibility of the ranking by interaction scores, and is called “reliability” in our results.

**Data availability**

Almost all data generated or analysed during this study are included in this published article (and its Supplementary Information files). Upon request raw data would be made available.

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Author contributions
M.B. and P.O. conceived and designed the experiments. M.B., S.F. and D.L. performed the experiments and acquired the data. The expertise on GacS-GacA system, two-component systems and Direct Coupling Analysis was provided respectively by W.A., D.W. and A.F.B. M.B., P.O., A.F.B., D.W., C.S., T.H., and W.A. interpreted the results. M.B. and D.W. wrote the manuscript and all authors read, revised and approved the final manuscript.

Competing interests
The authors declare no competing interests.

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