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REVIEW

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Regulation of bacterial phosphorelay systems

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In terms of biomass, bacteria are the most successful organisms on earth. This is partly attributed to their tremendous adaptive capabilities, which allows them to sense and rapidly organise responses to changing environmental stimuli. Using complex signalling mechanisms, bacteria can relay cellular information to fine-tune their metabolism, maintain homeostasis, and trigger virulence processes during infection. Across all life, protein phosphorylation represents the most abundant signalling mechanism, which is controlled by a versatile class of enzymes called protein kinases and their cognate phosphatases. For many years, histidine kinase (HK)-containing two-component systems (TCSs) were considered the canonical instruments of bacterial sensing. However, advances in metagenomics has since proven that bacterial phosphorelay is in fact orchestrated by a functionally diverse array of integrated protein kinase types, including Ser, Thr, Tyr and Arg-targeting enzymes. In this review, we provide an up-to-date appraisal of bacterial kinase signalling, with an emphasis on how these sensing pathways are regulated to modulate kinase output. Finally, we explore how selective kinase inhibitors may be exploited to control infections and combat the looming health emergency of multidrug resistant bacteria.

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Life has evolved an abundance of protein-based molecular machines to detect and coordinate responses to environmental stimuli. These biological processes are collectively referred to as signal transduction events and are controlled, in part, by a series of functionally divergent enzymes that posttranslationally introduce covalent chemical modifications onto target proteins in a tightly regulated manner. One such class of enzymes are the protein kinases, which catalyse the reversible addition of a phosphate group from ATP to conserved amino acid residues within a protein substrate, in a process called phosphorylation. Since the discovery of protein kinase activity in 1954, and the first protein kinase (phosphorylase kinase) a few years later, 2-4 it is now accepted that protein phosphorylation controls virtually every aspect of life. Indeed, protein phosphorylation is the most extensively studied posttranslational modification in bacteria, as evidenced by the wealth of literature written on the topic.⁵⁻¹³ Bacterial kinases can be classified into 5 groups: His kinases (HKs), non-Hanks type Tyr kinases, Arg kinases, Hanks-type Ser/Thr kinases (the so-called 'eukaryotic-like Ser/Thr kinases' [eSTKs]), and atypical Ser/Thr kinases. 14 HK-containing two-component systems (TCS) constitute the dominant signal transduction circuitry across all prokaryote lineages and function as conduits between

Department of Biochemistry, Cell and Systems Biology, Institute of Systems, Molecular and Integrative Biology, University of Liverpool, Liverpool L69 7ZB, UK. E-mail: bs0u4193@liverpool.ac.uk extracellular sensing and intracellular responses. Individual bacterial species can encode multiple TCSs, which enables sensing of a diverse and ever-expanding repertoire of known environmental signals. The ubiquitous TCSs are thus considered master regulators of prokaryotic metabolism, some environmental changes, virulence, surface, antimicrobial resistance, smoothity and chemotaxis. Although TCSs are not components of mammalian cellular signalling networks, similar systems have been identified in some eukaryotic fungi, and higher plants. These eukaryotic TCSs are structurally more divergent, often supplemented with additional regulatory modules, and can tune the outputs of downstream signalling proteins, such as other kinases. These

In addition to HKs, phosphorylation of protein Ser/Thr and Tyr residues (*O*-phosphorylation), by enzymes that share striking sequence or structural similarities with canonical eukaryotic Hanks-type kinases (Fig. 1), has also emerged as a pervasive phosphosignalling mechanism in bacteria and archaea. ^{42,43} Since the discovery of the first eSTK, Pkn1 in *Myxococcus xanthus*, ⁴⁴ it is now established that the genomes of bacteria can contain multiple eSTKs, with the prevalence of these kinases approximately correlating with the complexity of the bacteria's replicative niche. ⁴⁵ For example, *Sorangium cellulosum* encodes 317 eSTKs, which may imply a multi-directional signalling network reminiscent of eukaryotic systems. ⁴⁶ In fact, >80% of the *Mycobacterium tuberculosis* proteome (which encodes 11 eSTKs, including PknG (Fig. 1B)), and 63% of the theoretical proteome

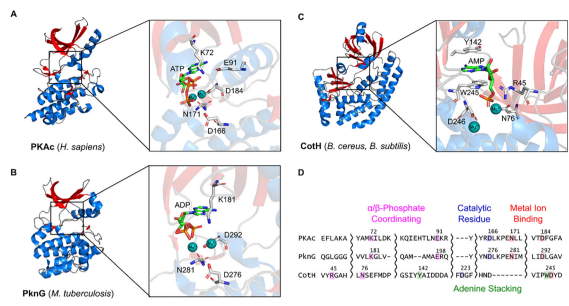


Fig. 1 Structural similarities between eukaryotic and prokaryotic Ser/Thr kinases. Crystal structures of (A) human PKA in complex with ATP (PDB: 4WB5), (B) M. tuberculosis PknG in complex with ADP (4Y0X) and (C) Bacillus atypical protein kinase, CotH, in complex with AMP. Insets show ATP/AMP coordinating amino acid residues (white sticks). α -Helices are shaded blue, β -strands are shaded red, loops are shaded grey, Mg²⁺ ions are represented as turquoise spheres and ATP/AMP is represented by green sticks. (D) Sequence alignment of PKA with PknG and CotH (using MUSCLE). Canonical amino acid residues (or functional equivalents) involved in ATP binding or phosphate transfer are colour coordinated. Note that the atypical kinase, CotH, aligns poorly due to its highly diverged sequence.

of Clostridium difficile is predicted to be O-phosphorylated.^{7,12} Several bacteria also encode atypical Ser/Thr protein kinases which diverge from the traditional Hanks-type kinase fold. For example, the HPr kinase/phosphorylase (HprK/P) of Bacillus subtilis is unrelated to Hanks-type kinases (instead binding nucleotides via a Walker A motif located in the P-loop) and regulates catabolite repression by phosphorylating (and dephosphorylating) the central HPr protein of the phosphoenolpyruvate: sugar phosphotransferase system. 47-49 YihE is another example of an atypical Ser/Thr kinase of E. coli that does not resemble canonical eSTKs in terms of structure or sequence homology, but plays critical role in regulating stress-related programmed cell death in bacteria. 50,51 Interestingly, CotH, an atypical protein kinase of B. subtilis and B. cereus (Fig. 1C) shares sequence similarity with the human Golgi casein kinase, Fam20C.52 Although CotH is not known to serve sensing or signalling functions per se, it is required to maintain the integrity of the spore-coat structure.⁵² CotH retains some features of canonical Ser/Thr protein kinases (including conserved active site residues corresponding to D166, N171, and D184 of the prototypical protein kinase, PKA) but diverges from the typical kinase fold in several crucial ways (Fig. 1C and D); it lacks a putative metal binding DFG motif and utilises an unusual mode of ATP binding involving two aromatic residues, Tyr142 and Trp245, that 'sandwich' the adenine moiety of ATP. 52 In contrast to eSTKs, bacterial tyrosine kinases (BY tyrosine kinases) display limited homology with their mammalian equivalents, and are characterised by Walker A and B ATP/GTP binding motifs.⁵³ Finally, while "classical" sensing and signalling roles have yet to be identified, arginine phosphorylation is known to regulate

several cellular processes, particularly in Gram-positive bacteria, such as B. subtillus and Staphylococcus aureus, where it functions as a protein degradation signal.^{54,55} For example, phosphorylation of proteins by the arginine kinase, McsB of B. subtilis, targets them for recycling by the ClpCP protease system.⁵⁴ In contrast to the other phosphotransfer systems, N-phosphorylation of arginine is a relatively understudied PTM, owing to the intrinsic lability of the phosphoamidate bond impeding conventional proteomics investigations.⁵⁶ As a result, our broader understanding of arginine phosphorylation and its roles within bacterial signaling networks remains limited. Collectively, the varied phosphorelay systems encoded by bacterial species provide a finely tuned and tightly regulated series of signal cascades that govern multiple aspects of bacterial life. This review provides an up-to-date account of the mechanisms that regulate bacterial phosphotransfer events.

Regulation of TCS phosphorelay signals

A prototypical bacterial histidine kinase (HK) is a multidomain His to Asp phosphorelay enzyme (Fig. 2A), consisting of a variable N-terminal "sensing region", transmembrane (TM) domain, and a conserved catalytic transmitter domain (containing catalytic ATPase [CA domain] and homodimeric phosphoacceptor/phosphotransferase [DHp] subdomains⁵⁷). HKs also commonly contain one or more intracellular signalling domains; including HAMP (found in histidine kinases, adenylyl cyclases, methyl binding proteins, and phosphatases), PAS (Per-

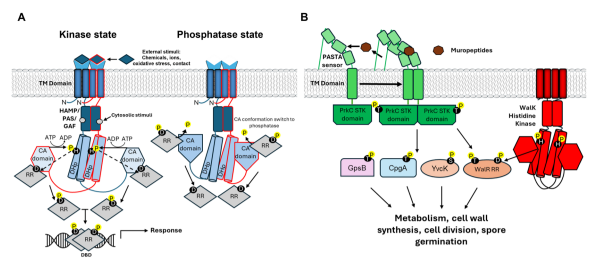


Fig. 2 Overview of bacterial kinase signal transduction mechanisms. (A) Asymmetrical TCS kinase state: classical two component signalling systems (TCSs) respond to extracellular (through periplasmic receptor) or cytosolic (through HAMP/PAS/GAF) sensor domains, leading to dimerization. In a multistep phosphorelay, the catalytic CA domains phosphorylate, in trans, a conserved His in the DHp domain. The phosphate is then shuttled (dashed arrows) to a conserved Asp residue in cognate response regulators (RRs). RRs dimerize and bind to a specific DNA binding domain (DBD) to elicit transcription of target genes. Symmetrical TCS phosphatase state: in the absence of stimuli, the CA domain undergoes a conformation switch to a phosphatase-active state, dephosphorylating RRs in cis, releasing phosphate into the cytosol. (B) Simplified schematic of a B. subtilis eSTK, PrkC, responding to extracytosolic stimuli via PASTA sensor domains. During stationary phase, binding of muropeptides to PASTA repeats induces dimerization of PrkC TM and eSTK domains, leading to in trans autophosphorylation and subsequent phosphorylation of a range of target cytosolic proteins. Substrates include, but are not limited to, GpsB (a cell division protein⁷⁴), YkwC (an oxidoreductase⁷⁵) and CpgA (a GTPase⁷⁶). PrkC also regulates the WalRK TCS by directly phosphorylating the WalR RR, influencing expression of genes relating to cell wall biosynthesis.⁷⁷

ARNT-Sim), STAC (SLC and TCST-Associated Component) or GAF (found in cGMP-specific phosphodiesterase, adenylyl cyclases, and FhlA) domains, 57,58 that augment HK regulation by serving as sensing conductors of extracellular stimuli. Upon stimulation, the signal is transduced to the TCS catalytic core, resulting in ATP binding by the CA domain and autophosphorylation of the conserved DHp acceptor His residue. The phosphate moiety is then relayed to the receiver (REC) domain of a response regulator (RR) module, which catalyses its own aspartic acid phosphorylation.³⁸ Following phosphorylation, most RRs undergo a conformational switch (typically leading to homodimerisation) which facilitates binding to regulatory genomic motifs and activation or repression of gene expression.⁵⁹⁻⁶² Fascinatingly, there is also growing evidence to suggest that some RRs also possess phosphorylation-independent regulatory functions. 63-65 For example, the phosphorylated form of the Salmonella typhimurium RR, RcsB, can homodimerize or form a heterodimer with the ancillary regulator, RcsA, to modulate gene transcription.66 However, recent structural investigations have revealed that an active RcsB dimer can be stabilized just by binding to DNA even in the absence of phosphorylation⁶⁷ and the unphosphorylated form of RcsB can also interact with other transcription factors to regulate gene expression⁶⁸⁻⁷¹ adding an extra layer of regulatory complexity to the traditional TCS model. Some HKs are also 'hybrid kinases' that are supplemented with carboxy-terminal REC and/or histidine phosphotransfer (HPt) domains, which are frequently involved in multi-step phosphorelay systems with contiguous signalling elements, such as DHp and RR proteins. 72 Although TCS-HKs are canonically recognized to adopt a dimeric architecture, monomeric species have also

been reported, including EL346 of Erythrobacter literalis. 73 In the inactive EL346 state, the typical DHp dimer interface of the kinase domain is blocked by direct interaction with a blue-light sensitive light-oxygen-voltage (LOV) domain, which competes with the CA domain for access to the phosphorylatable histidine.

Extra/intracellular or periplasmic signals are typically sensed by N-terminal located sensory PAS domains and propagated along the HK through a series of transient structural rearrangements and inter-domain interactions that regulate catalytic activity and phosphotransfer.58,78 Although the amino acid composition between different PAS domains of HKs is extremely variable (which drives substrate specificity), they adopt a highly conserved multi- β -sheet core structure with adjoining α -helices that forms a ligand-accepting cleft region. 57,78,79 Sensing can also be achieved by PAS-like domains, all α-helix type structures, and Venus flytrap (VFT) domains. 57 Through these mechanisms, TCSs can perceive and respond to a range of chemical and physical stimuli. In the absence of an activation signal, many HKs function as putative phosphatases of RRs, allowing a molecular reset of TCS signalling.80-83 Shifts in the equilibrium of kinase and phosphatase-active states are therefore necessary to balance the net physiological outcomes of TCS signalling. 84,85 The molecular mechanisms of HK signal sensing, as it pertains to the structural dynamics of these mutually exclusive kinase and phosphatase competent states, have been extensively reviewed elsewhere. 8,58,86 In brief, the activity of HKs is contingent on signal-dependent domain reorganisations, whereby the conserved kinase core adopts an asymmetrical homodimer configuration, containing both active and inactive catalytic subunits.8 The CA domain of the inactive conformation can

bind to ATP, whilst the DHp domain remains accessible for phosphotransfer to the REC domain of an RR. In the active conformation, both the DHp and CA domains form a tight complex to catalyse in-cis or in-trans transfer of the γ -ATP phosphate onto the eponymous His site of the DHp domain, which is inaccessible to the RR. ^{8,86,87} Bacteria also encode a range of non-TCS kinase signalling relay systems. A related group of non-TCS HKs are associated with the core signalling complexes of chemoreceptors in motile bacteria and archaea, ⁸⁸ which incorporate soluble cytoplasmic HKs (such as CheA in *E. coli*) as part of a supramolecular sensing array. Of note, several other HKs that are soluble enzymes, lacking transmembrane segments, have been identified. ^{34,89,90}

Regulation of Ser/Thr protein kinases

Membrane-associated (and cytoplasmic) Hanks-type eSTKs serve regulatory functions in numerous cell processes and have also been implicated in the virulence of pathogens. 10,91-96 eSTK-based signalling systems typically rely on the opposing activities of a protein kinase and an associated phosphatase, that (in contrast to HKs) can concurrently (de)phosphorylate Ser/Thr residues of a broad range of protein classes. 10 However, mechanistic details of how branching eSTK signalling cascades are organised are relatively scarce. In addition to a surprisingly well-conserved Hanks-type fold in the kinase catalytic core, 45,94 many eSTKs are buttressed with additional modular regulatory domains that serve as sensor regions, mediate protein-protein interactions and are sites of ligand binding. These include penicillin-binding and Ser/Thr kinase-associated repeats (PASTA) and forkhead-associated (FHA) domains that recognize phosphothreonines. 95,97,98 In particular, extracellular sensory PASTA domains of transmembrane eSTKs (the "PASTA kinase" family), which can interact with ligands (usually peptidoglycan or peptidoglycan precursors) and induce activated kinase domain homodimers, are reasonably well characterised. 45,95,99 PrkC, a membrane-associated eSTK of B. subtilis, contains three PASTA domains and plays crucial roles in spore germination, vegetative growth, modulation of cell wall synthesis and antibiotic resistance. 5,100 PrkC activity is differentially regulated by bacterial growth phases. During the stationary phase, binding of peptidoglycan fragments to the extracellular PASTA domain results in PrkC dimerisation and activation of the cytosolic kinase domain (Fig. 2B),101 whilst during exponential growth, activation is attained by binding to the cell division proteins GpsB, DivIVA, and EzrA. 102 Peptidoglycan binding by PASTA repeats of S. aureus STK similarly induces dimerization and activation of the kinase, 103 which goes on to phosphorylate and inhibit several cell wall synthesis enzymes, such as FemX. 104 PASTA domains also enable ligand-induced modulation of kinase activity and proper localisation of PknB from M. tuberculosis 105,106 with dimerisation of the PknB kinase domain Nlobe resulting in allosteric activation of catalytic function. 107

A common feature of eukaryotic protein kinases and several eSTKs is a requirement for phosphorylation of conserved sites

within their putative activation loop to switch on catalysis. For example, the PASTA-kinase of Enterococcus faecalis, IreK, autophosphorylates at three positions on the activation loop (Thr^{163, 166, and 168}) and an additional site on the C-lobe in response to cephalosporin treatment, resulting in enhanced activity. 108 Autophosphorylation of activation loop amino acid residues appears to be a conserved regulatory mechanism for many eSTKs, including PrkA of Listeria monocytogenes, 109 PrkC from B. subtilis, 110 YegI of E. coli 111 and PknB from M. tuberculosis. 112 The molecular basis of mycobacterial cytoplasmic PknG activation is also predicted to involve N-terminal domain autophosphorylation, which creates a recruitment site for the substrate GarA, and displaces a substrate-occlusive N-terminal rubredoxin domain.113 Phosphorylation of juxtamembrane regions of PknG is also predicted to initiate interaction with the cell wall recruitment protein, FhaA. 114 Other unique modes of eSTK regulation have also been discovered, including for YabT of B. subtilis, which is essential for a robust DNA-damage response, and harbours a DNA-binding motif that activates the kinase upon ds- or ss-DNA binding. 115

Antagonistic regulation of kinase activity by phosphatases

A responsive signalling network requires rapid and reversible addition of phosphate moieties to target proteins to selectively 'switch on' and 'switch off' signalling cascades. Across life, this involves the concerted opposing actions of protein kinases and protein phosphatases. For this purpose, dual-activity HKs can adopt phosphatase-competent states to 'turn-over' their signalling outputs. Moreover, RR dephosphorylation can be further accelerated by the actions of dedicated phosphatases, including CheZ of E. coli, Rap and Spo0E of B. subtills, and CheX of Borrelia burgdorferi. 58 Bacteria have also evolved additional mechanisms to degrade phosphate-dependent signals in response to environmental cues. For example, bacterial phytochrome photoreceptors usually transmit photosensory input through a HK TCS system,82 including Agp1 bacteriophytochrome from Agrobacterium fabrum which exhibits lightsensitive HK-autophosphorylation activity. 116 In contrast, no kinase activity has been detected for a structurally homologous bacteriophytochrome, DrBphP, from Deinococcus radiodurans which functions exclusively as a light-dependent phosphatase for the RR DrRR. 117 In fact, bespoke bacterial phosphatases are prevalent in bacterial genomes, particularly in the context of eSTK- Ser/Thr phosphatase (STP) pairs 10,14,93,118 and even arginine phosphorylation is counteracted by a cognate phosphatase, YwlE. 119 M. tuberculosis expresses 11 eSTKs (PknA-L, two of which, PknG and PknK, are soluble proteins) which collectively phosphorylate a wide range of protein substrates and play a dominant role in bacterial signalling, virulence and viability. 7,120-122 Full activity of the membrane-anchored eSTK, PknB, depends on phosphorylation of two activation loop residues (Thr¹⁷¹ and Thr¹⁷³), which can be reversed by dephosphorylation by the cognate STP, PstP. 112 PstP is the only M.

RSC Chemical Biology Review

tuberculosis phosphatase discovered to-date and is predicted to dephosphorylate all members of the PknA-L family and their substrates. 45 Interestingly, phosphorylation of PstP by PknA/B also regulates its catalytic activity, suggesting a regulatory feedback loop. 123,124

The conserved S. aureus eSTK-phosphatase pair (STK-STP) is co-transcribed on a regulatory operon, with STP dephosphorylating STK. 125,126 Notably, STK-STP has been implicated in controlling expression of secreted virulence factors, including hemolysins α , β and γ , which are drivers of *S. aureus* infection¹²⁷ and the kinase/ phosphatase pair also have antithetical activities on cell wall biosynthesis. 104 Streptococcus agalactiae phosphatase, Stp1, is also required for appropriate regulation of Stk1 eSTK function. 128 eSTK-STP pairs of Streptococci, including Stk1-Stp1 in S. agalactiae, Stk-Stp in S. pyogenes, StkP-PhpP in S. pneumoniae, and PknB-PppL in S. mutans are similarly co-transcribed, to ensure integrative reversible control of Ser/Thr phosphate-signalling. 118 An antagonistic relationship has also been observed between an eSTK of Pseudomonas aeruginosa, PpkA (which is essential for assembly of a type-VI secretion system [H-T6SS] and secretion of hemolysincoregulated protein 1 [Hcp1]) and PppA, a Ser/Thr phosphatase. 129 The lifecycle of Bacillus anthracis, the aetiological agent of human anthrax, comprises vegetative and sporulating phases. Spore germination is partly controlled by the co-expressed kinase-phosphatase pair PrkC-PrpC, 101 with PrkC kinase activity inhibited by PrpC-dependent dephosphorylation. Curiously, PrpC is a dual specificity phosphatase capable of removing a phosphate moiety from Ser, Thr and Tyr residues of PrkD and PrkG kinases. 130 Likewise, although a putative tyrosine kinase of P. aeruginosa has yet to be discovered, TpbA is also believed to be a novel dual specificity Try-phosphatase that negatively regulates biofilm formation in P. aeruginosa. 131

Regulation of kinase function through protein binding and subcellular distribution

Interactions with accessory proteins can determine the catalytic outputs of bacterial kinases, either through direct modulation of catalysis or organisation of their spatial distribution. For example, the McsAB^{B. subtilis} arginine kinase holoenzyme complex is comprised of the McsB kinase domain and an allosteric activator subunit, McsA. 132 As previously discussed, interaction of the eSTK PrkCB. subtilis with GpsB, DivIVA, and EzrA is responsible for appropriate localisation and activation of the kinase. Interestingly, PrkC is variably regulated by GpsB, which can function as both an activator and suppressor of PrkC, with the phosphorylated form of GpsB (itself a substrate of PrkC) providing a negative feedback loop to PrkC activity. 102 EmbR is a transcriptional regulator, and a substrate protein of the mycobacterial eSTK, PknH. Curiously, a structural homolog, EmbR2 (which is a substrate of PknE and PknF) physically interacts with and inhibits PknH. 133 Accessory proteins, GlnX and GlnH, are also required for PknG activation (a cytoplasmic eSTK of mycobacteria that lacks extracellular or

transmembrane sensing regions) in response to extracellular amino acid concentration.134 HipA adopts an atypical protein kinase fold, 135-137 and has been identified in the genomes of E. coli, and several paralogs are also found in other Gramnegative bacteria. 138 HipA functions as a dormancy-driving toxin by inhibiting protein synthesis and is allosterically suppressed under normal growth conditions via direct complexation with the HipB antitoxin. 136,139 Remarkably, HipA activity is also negatively regulated by intermolecular autophosphorylation of Ser150, which stabilises a P-loop 'out-state' and disrupts the ATP-binding pocket. HipT of Legionella pneumophila, which has homology to the N-terminal region of HipA, similarly adopts an inactivate conformation when in complex with its cognate antitoxin, HipS, which leads to the steric occlusion of ATP from the kinase core. 138

HKs can similarly be regulated by accessory proteins. For example, two B. subtilis HKs, KinA and KinB, are essential for initiation of sporulation, and are respectively inhibited by Sda and KipI, which are predicted to repress autokinase activity by sterically blockading interaction between the CA and DHp subdomains. 141,142 In contrast, binding of the accessory protein, PtsN to the DHp domain of the KdpD HK of E. coli stimulates trans-autophosphorylation between protomers of a KdpD dimer. 143 Interaction with protein subunits can also shift the equilibrium of TCS kinase and phosphatase states. For example, the core SaeRS TCS of S. aureus is composed of a sensor kinase (SaeS), RR (SaeR), and two auxiliary proteins, SaeP and SaeQ, which regulate genes associated with biofilm formation.144 SaeQ and SaeP form a complex with SaeS and induce a phosphatase-competent state in the HK, effectively silencing signalling by the TCS. Finally, HK sensing of environmental stimuli (and subsequent activation) can also be mediated through accessory proteins. For example, the BceS HK of B. subtilis forms an activating intermolecular sensory complex with BceAB. 145 In this system, the BceS sensor kinase is indirectly activated by the peptide-based antibiotic, bacitracin, through a flux-sensing mechanism which detects conformational cycling of the BceAB transporter. 146,147 The PhoQP system can also be indirectly regulated by the acid sensing EvgSA system of E. coli via an intermediary membrane protein, SafA. SafA expression is induced following activation of EvgSA, and directly interacts with the PhoQ periplasmic sensor domain to stimulate autophosphorylation. 148-150 The Rcs envelope stress response phosphorelay system (comprising RcsC HK, RcsD phosphotransfer protein, and RcsB RR) is conserved in enterobacteria (reviewed in ref. 66, 67 and 151) and is regulated by the outer-membrane sensor lipoprotein, RcsF. 152 Under normal growth conditions, another inner membrane protein, IgaA, represses the RcsCDB phosphorelay cascade through an inhibitory interaction with RcsD. However, detection of envelope stress by RcsF induces complex formation with AgaA, relieving repression and resulting in autophosphorylation of RcsC.^{153,154}

Subcellular localisation can also be critical for kinase function. Cell pole organising protein PopZ of Caulobacter crescentus self-assembles into a polymeric superstructure that recruits Review

several signalling protein clients (such as the cell fate HKs CckA and DivJ) into a regulatory hub that coordinates cellular polarity. 155,156 DivJ expression at the cell poles is critical for propagating chromosomal replication and stalk appendage biogenesis, 157 and this is controlled by PopZ-dependent recruitment factor, SpmX. 156 A second C. crescentus scaffold protein, PodJ, assembles at the opposite cell pole to PopZ and sequesters several additional proteins, including the HK PleC via its sensory PAS domain. 158 CckA and PleC of C. crescentus are bifunctional HKs that oscillate between kinase and phosphatase activity. 159 CckA exhibits density-dependent activity coordinated through its tandem PAS domains; 159 functioning as an indirect kinase of the CtrA transcription factor when accumulated at high density at a newly formed cell pole, 160 but transitioning to a phosphatase of CtrA in the presence of high levels of the secondary messenger c-di-GMP (cdG). 160-162 Recruitment to the PopZ microdomain, which defines CckA surface density, is partly regulated through complexation with a catalytically inactive pseudohistidine kinase, DivL. 163-165 Curiously, DivL is capable of transforming information relating to subcellular localisation into conformational switches that can toggle CckA between its kinase or phosphatase modes, suggesting a highly dynamic regulatory profile for this HK complex. 166-168 PleC is also mechanistically density-regulated through PAS domains and is recruited to polar PodJ biomolecular condensates, which simultaneously stimulates PleC phosphatase function and suppresses kinase activity. 158 Spatial organisation may also be an evolutionary strategy to limit cross-talk between TCSs, and has been observed in Rhodobacter sphaeroides and for the CheZYA chemotaxis system of E. coli, as reviewed by Sourjik and Armitage, 2010.169

Cross-talk between bacterial phosphorelay systems

Cognate HK-RR pairs are usually highly monogamous and are generally co-expressed under a single promotor. 170,171 As such. encoded TCS paralogs typically serve distinct cellular functions through linear signal transmission processes. 61,171 Retention of signal fidelity is primarily dictated by specificity-determining amino acids that flank the conserved phosphotransfer/acceptor sites in the DHp and REC domains. 172 However, convergence between non-cognate TCS pairs has also been observed in some bacteria. 173-176 For example, in E. coli, NarQ can target its cognate RR, NarP, and also cross-phosphorylate NarL of the NarX-NarL TCS axis, 177 and CreC can interact with PhoB, the cognate RR of PhoR. 178 Far from representing an evolutionary redundancy, crosstalk between TCS may enhance the complexity of responses to individual input events, and promote bacterial growth, metabolism and survival. 173,176 Relaxed specificity is potentially less detrimental in unorthodox 'hybrid' TCS, where spatial tethering of transmitter and receiver domains effectively increases their local concentrations to one another. 179-181 For example, the GacSA hybrid TCS of P. aeruginosa is part of a multi-kinase communication network with

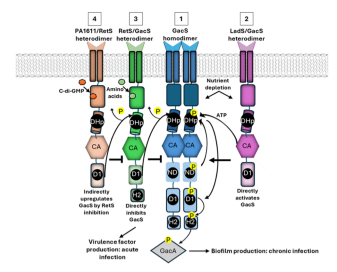


Fig. 3 Cross-talk between GacS/A and other TCSs in P. aeruginosa. (1) Periplasmic or cytosolic sensor domains respond to environmental or cellular signals that cause GacS homodimerization. The ND and CA domains facilitate autophosphorylation of a conserved His in the DHp domain. GacS contains an additional receiver (D1) domain, which becomes phosphorylated by intramolecular phosphotransfer. The phosphate is subsequently shuttled to the H2 domain and finally GacA, the RR. (2) LadS positively regulates GacS function, enhancing kinase activity and supporting chronic infection characteristics. (3) RetS forms a complex with GacS, inhibiting its kinase activity thereby suppressing GacA phosphorylation, leading to enhanced expression of acute virulence factors. (4) PA1611 binds to RetS, preventing its inhibitory interaction with GacS. By sequestering RetS, PA1611 allows GacS to maintain its kinase activity, facilitating GacA phosphorylation and promoting chronic infection pathways.

LadS, RetS and PA1611 hybrid HKs^{182,183} (Fig. 3). Following signal-mediated activation, GacS autophosphorylates at His294 (H1), and the phosphate moiety is then shuttled to Asp717 (D1) and His863 (H2).18 Heterodimersiation of RetS and GacS downregulates GacSA signal transduction by inhibiting GacS autophosphorylation, siphoning phosphate groups from the DHp site, and through direct dephosphorylation of the GacS D1 site by RetS. 184,185 Moreover, RetS-dependent inhibition can subsequently be relieved through formation of a competitive RetS-PA1611 heterodimer. 186 In contrast, LadS forms a multicomponent phosphorelay system with GacS, with the transmitter and REC domains of LadS facilitating trans-phosphorylation of the GacS H2 domain. 182 Through these mechanisms, RetS and LadS can reciprocally regulate virulence factor expression controlled by the GacSA regulon and influence initiation of chronic P. aeruginosa infection. 182 Cross-communication has also been detected between HssSR and HitSR of B. anthracis, 187 NtrBC and NtrYX of Rhodobacter capsulatus, 188 RscS and SypF of Vibrio fisheri¹⁸⁹ HpArsRS and HpNikR of Helicobacter pylori¹⁹⁰ and GraSR and ArlSR of S. aureus.¹⁹¹ Interestingly, RR heterodimers have also been described that can enable differential coactivation of target genes, as reviewed by Agrawal et al., 2016.¹⁷²

HKs and the other major phosphorelay system, eSTKs, may also intertwine to potentiate the output of adaptive signalling responses (Fig. 2B). This is perhaps not surprising given recent

studies revealing that eSTKs exert a broad regulatory influence; phosphorylating 80% of the M. tuberculosis proteome and modulating expression of $\sim 30\%$ of its genes.⁷ Such an expansive regulatory footprint may facilitate overlapping regulons between eSTKs and TCS. Indeed, 170 O-phosphorylation sites were identified on TCSs of M. tuberculosis, suggesting extensive intersection of these two phosphosignalling pathways. 192 Notably, phosphorylation of the *M. tuberculosis* TCS-HK, NarS, at Thr³⁸⁰ by an eSTK, PknL, was sufficient to stimulate its autokinase activity and downstream signalling. 192 The AlgRZ TCS, which is comprised of AlgR, and it's putative HK AlgZ, is an important regulator of P. aeruginosa gene expression and pathogenesis. 193 Strikingly, Ser 143 of the RR, AlgR, is also phosphorylated by the eSTK Stk1, which adjusts its physiological functions. 194 Cross talk between STK of S. aureus and TCSs involved in cell wall metabolism has also been described, through direct phosphorylation of several Thr residues in VraTSR, 195 WalRK, 196 and GraSR. 197 The StkP-PhdP eSTK-STP pair of S. pneumoniae can also reversibly regulate the RRs, RitR, ¹⁹⁸ RR06, ¹⁹⁹ and PknB^{M. tuberculosis} can downregulate DevSR and SenX3/RegX3 signalling by phosphorylating their RRs. 200,201 Fascinatingly, another P. aeruginosa, RR PA3346, contains an N-terminal receiver domain and an active STP domain that dephosphorylates Ser⁵⁶ of another RR, PA3347. The Ser/Thr phosphatase activity of PA3346 is in turn potently stimulated following phosphorylation of the RES domain by a HK, PA2824.202 Direct modulation of HK kinase activity by eSTKs has also been described. For example, the CroS HK of the opportunistic pathogen, Enterococcus faecalis, is phosphorylated at a Thr by a transmembrane PASTA-eSTK, IreK, thereby enhancing HK activity. Of note, this site (Thr346) is situated in the ATP lid of the CroS CA domain, a crucial structure for appropriate nucleotide positioning and phosphate exchange during autophosphorylation.²⁰³ An eSTK of B. subtilis, YbdM, was also reported to directly phosphorylate the sensing domain of an atypical cytoplasmic HK, DegS, and enhance its kinase activity in vitro. 204 Fascinatingly, the S. aureus tyrosine protein kinase CapAB (which positively regulates capsule biosynthesis) can be negatively regulated by PknB-dependent phosphorylation²⁰⁵ suggesting communication between eSTK and tyrosine kinase phosphotransfer networks.

Regulation of kinases by cofactors, secondary messengers, and quorum sensing

Much like their eukaryotic counterparts, bacterial kinases are also regulated by cofactors that can induce conformational changes and interdomain interactions or augment catalytic function. Coordination of haem or flavin adenine-dinucleotide (FAD) cofactors by HK PAS domains enables sensing of cellular redox state. For example, the kinase activity of FixL (found in nitrogen-fixing rhizobia), is potently inhibited by O₂ binding to a PAS-B located haem group.²⁰⁶ Redox sensing cysteines have also been discovered in the CA domain of the

S. aureus HK, SrrB, which form a unique intramolecular disulfide bond that tunes autokinase function.²⁰⁷ Furthermore, oxidation or S-nitrosylation of a Salmonella enterica RR, SsrB, at Cys²⁰³ lowers DNA-binding affinity and thus signalling by the SsrA/SsrB TCS, 208 which may represent a defensive adaptation to promote Salmonella fitness when exposed to nitric oxides (NOs) produced by the host immune response. Similarly, Cys⁶⁷, located in the REC domain of the redox-sensitive WalR RR, can be S-nitrosylated to modulate WalKR signalling and promote S. aureus NO-mediated vancomycin resistance. 209 The cyanobacterium Synechocystis sp. PCC 6803, encodes six eSTKs (SpkA-D, G), one pseudokinase (SpkE), and five atypical kinases (SpkH-L). Of note, SpkB was shown to be inactivated by oxidation of an N-terminal Cys motif, as an adaptation to oxidative stress tolerance.210 As previously discussed, the secondary messenger c-di-GMP binds within the interface between the PAS-B site and CA domain of the HK, CckA, to stimulate phosphatase activity. 159,211 In contrast, the kinase activities of ShKa, a cytosolic hybrid HK of C. crescentus, and RavS, a HK of Xanthomonas campestris, are increased following binding to c-di-GMP. 212,213 The activity of a DNA-damage responsive eSTK of the extremophilic bacteria Deinococcus radiodurans, RqkA, is also enhanced by binding to pyrroloquinoline-quinone (POO).214

S. aureus biofilm formation is partially regulated by intercellular communication and density sensing, commonly known as quorum sensing.215 In this manner, bacteria can detect changes in population numbers based on the local concentration of specific autoinducer signals. Direct binding of an autoinducer peptide (AIP) to the AgrC HK stimulates phosphorylation of the RR AgrA, resulting in positive regulation of toxin genes, autoinduction of AIP biosynthesis, and reduced expression of several surface adhesins, as reviewed by Jenul & Horswill, 2019. 216 The Lux system in Vibrio species is another well-studied TCS that is regulated by quorum sensing.²¹⁷ Low autoinducer concentrations at low cell density induces LuxQ activity and results in phosphorylation of the LuxO RR. In contrast, high autoinducer concentrations (detected by the periplasmic receptor LuxP) activate HK phosphatase activities and reverts the TCS to a pre-activated state. 218

Regulation of secreted effector kinases

In order to establish an infection and subvert host defences, pathogenic bacteria can also deploy an arsenal of signal pathway-hijacking virulence effectors, including eSTKs, into the cytoplasm of target cells. PknB^{M. tuberculosis} (and the phosphatase SapM) was the first discovered effector eSTK, which functions to promote intracellular survival of the bacteria in macrophages by blocking phagosome maturation. 220,221 Since then, secreted bacterial effector kinases have been identified in the genomes of numerous pathogens, including *Yersinia*, *Salmonella*, *Legionella* and *Shigella*²¹⁹ but little is understood about how they are regulated. To avoid premature activation of cytotoxicity, the activities of virulence factors and exotoxins are

extremely tightly controlled, only 'switched on' following secretion or exposure to specific host-derived activation signals. Similar host-dependent activation mechanisms have also been observed for a handful of effector eSTKs. Legionella pneumophila (L.p.) is an opportunistic intracellular pathogen of human alveolar macrophages and the causative agent of Legionnaire's disease. L.p. translocates over 300 survival-promoting effector proteins, including at least 6 experimentally validated eSTKs (LegK1-4, LegK7, and Lem28/Lpg2603) into invaded host cells via the Dot/Icm type IV secretion system (T4SS). 222-231 Fascinatingly, LegK7, which manipulates the host Hippo pathway by functioning as a molecular mimic of Mammalian Ste20-like kinases 1/2 (MST1/2), is allosterically activated by N-terminal binding to a host scaffold protein, MOB kinase activator 1A (MOB1A). 227,228 Similarly, Lem28 is a remote member of the protein kinase superfamily that is allosterically activated by the eukaryoticspecific ligand inositol hexaphosphate (IP₆).²³⁰ Members of the pathogenic Yersinia genus secrete an eSTK domain containing protein, YpkA, which directly phosphorylates the heterotrimeric Gag protein and vasodilator-stimulated phosphoprotein (VASP) to disrupt macrophage cytoskeletal dynamics and phagocytosis. 232,233 YpkA is delivered into host cells in an inactive form via a type III secretion system, where it is allosterically activated following actin binding to the actin binding domain (ABD) and undergoes extensive autophosphorylation. 234,235 The Shigella type-III secretion system effector OspG is an atypical serine/threonine protein kinase that lacks several canonical structural features, including a regulatory activation loop. 236 Despite the extensive degradation of the catalytic core, OspG exhibits intrinsic kinase activity, but only upon binding to components of the host ubiquitin system.²³⁶ In contrast, homologous atypical kinases of E. coli, NleH1, and NleH2, display ubiquitin-independent autophosphorylation which promotes interaction with target proteins.236

TCSs as potential therapeutic targets for the treatment of microbial infections

Multi-drug resistant (MDR) bacterial infections rank globally amongst the leading causes of mortality, and are projected to cause 40 million deaths by 2050.237 Most clinically approved antibiotics are derivatives of small molecules that disrupt a narrow range of essential microbial biosynthetic pathways. Compounds exhibiting alternative modes of action are scarce, ²³⁸ which has exacerbated the spread of MDR bacteria. Accordingly, the many crucial roles that TCSs and kinases serve in bacterial growth, viability and pathogenic processes (including antibiotic resistance mechanisms and biofilm formation) make them attractive alternative therapeutic targets to develop novel antimicrobials. Moreover, the CA and DHp domains of TCS (which are crucially absent in mammalian signalling pathways) are highly conserved across the bacterial kingdom, 84,239 potentially enabling development of compounds with broad-spectrum activity. 240,241 In this final section we discuss some recent advances in kinasetargeting antimicrobial drug strategies.

Several studies have investigated and confirmed TCS inhibition by small-molecule compounds. For example, WalK is an essential HK conserved among Gram-positive bacteria, including S. aureus and B. subtilis. 242 Walkmycins A, B, and C were demonstrated to exhibit strong antibacterial activity against B. subtilis (a model organism for studying pathogenic B. anthracis and B. cereus) and walkmycin B was revealed to specifically inhibit Walk autophosphorylation. 243 Interestingly. walkmycin C was also an effective inhibitor of three other Streptococcus mutans HKs: VicK, CiaH, and LiaS in vitro, 244 and isatin derivatives also exhibit potent antimicrobial activity against S. aureus, which was attributed to direct binding to and inhibition of Walk.²⁴⁵ Other inhibitors of HK autophosphorylation have now been identified for several different bacterium, targeting VicK and AgrC of S. pneumoniae, 246 DosS and DosT of M. tuberculosis, 247 and PhoQ of S. typhimurium, 248 suggesting that TCSs are viable antibacterial therapeutic targets.

Most identified TCS-targeting inhibitors, including walkmycin C, 244 are functional ATP-competitive molecules that 'blockade' ATP binding and thus HK catalysis (HKs such as $HK853^{Thermotoga\ maritima}$, $CheA^{E.\ coli}$ and $VicK^{S.\ pneumoniae}$ can all be inactivated by ATP obstruction²⁴⁹). The ATP-binding pocket of the HK CA domain adopts a distinctive α/β sandwich, consisting of four conserved regions (the N-box, the G1-box, the G2-box, and the G3-box) and a highly variable ATP-lid. 250,251 This structural architecture is known as the Bergerat fold, and is also observed in the ATP-binding domains of the diverse GHKL protein superfamily, which also includes DNA gyrases, HSP90, and the MutL mismatch repair enzyme.²⁵⁰ Guarnieri et al. (2008) demonstrated that the human HSP90 inhibitor, radicicol, binds to the ATPbinding pocket of the Salmonella HK, PhoQ (albeit weakly). 252 Since then, optimised derivatives of ATP-competitive HSP90 inhibitors have been explored as potential HK active agents. 251,253 For example, a fragment of an established HSP90, 3,4-diphenylpyrazole (DPP)-based inhibitor was shown to bind to the CA domain of CheA from Thermotoga maritima, and several DDP analogues are effective inhibitors of EnvZ^{E. coli}, PhoQ^{Salmonella typhimurium}, and CckA^{C. crescentus.} 251,253 Repurposing human protein kinase inhibitors (for which there are ~ 100 FDA approved clinical compounds) is another desirable strategy to identify novel therapeutic agents to target bacterial kinases. Carabajal et al. (2020) screened 686 compounds from the Published Kinase Inhibitor Set (PKIS by GlaxoSmithKline]), and identified quinazoline-based ATP-competitive inhibitors of S. typhimurium PhoQ that repressed autokinase kinase activity.²⁴⁸ However, despite these promising results, potential toxicity associated with 'off-target' inhibition of human kinase signalling pathways may necessitate a more refined, structureguided design approach to develop selective inhibitors which only target bacterial HKs before such agents can be considered therapeutically viable.

Interestingly, several non-ATP competitive bacterial kinase inhibitors have also been discovered. The HK PhoQ (of the conserved PhoP/PhoQ TCS) plays pivotal roles in Salmonella, Shigella, and Pseudomonas pathogenesis, regulating virulence factor production and bacterial survival within host cells.²⁵⁴ Moreover, PhoP/PhoQ contributes to 'last-resort' polymyxin antibiotic resistance by indirectly activating the PmrA/PmrB TCS. 255,256 Fascinatingly, PhoQ (Salmonella typhimurium) HK activity can be repressed by allosteric targeting its second transmembrane region using a hydrazone derivative compound, N^\prime -(thiophen-2-ylmethylene)benzohydrazide (designated $\rm A_{16}B_1$). $^{257-259}$ Watanabe et al., (2012) also identified signermycin B as a potent allosteric inhibitor of WalK that disrupted autokinase activity by sterically blocking the HK dimerization domain. 260 Moreover waldiomycin (and a methyl ester derivative) can effectively inhibit WalK from both S. aureus and B. subtilis (and PhoR and ResE from B. subtilis) by binding to the phosphoacceptor region and blocking autophosphorylation. 261,262

Conclusions

TCSs and bacterial kinases represent a promising frontier for the development of novel therapeutics for microbial infections, offering a versatile platform for addressing both antimicrobial resistance and bacterial virulence. Phosphate-based PTMs, regulated by kinases and phosphatases, are essential for bacterial homeostasis, viability and virulence and clinical compounds with the ability to inhibit conserved domains across multiple protein kinases could provide broad-spectrum antimicrobial activity. Conceptually, TCS inhibitors also show promise as therapeutic adjuvants, potentially synergizing with conventional antibiotics to overcome resistance mechanisms.²⁶³ For example, compounds like PhoQ-targeting A₁₆B₁ can enhance the bactericidal activity of polymyxins.²⁵⁹ However, despite significant interest in this area, no bacterial kinase-targeting small molecule have advanced to clinical approval. To progress this area of research, a combination of structure-guided drug design and innovative screening methodologies will be essential to develop optimised and selective kinase-targeting compounds with robust antimicrobial activity.

Author contributions

Writing – original draft: DPB, DMF, DMC & CW; writing – review and editing: DPB, DMF, DMC & CW; visualisation: DMF, DMC & DPB.

Conflicts of interest

There are no conflicts of interest.

Data availability

No primary research results, software or code have been included and no new data were generated or analysed as part of this review.

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