



Review

Bacterial tyrosine kinases: evolution, biological function and structural insights

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Reversible protein phosphorylation is a major mechanism in the regulation of fundamental signalling events in all living organisms. Bacteria have been shown to possess a versatile repertoire of protein kinases, including histidine and aspartic acid kinases, serine/threonine kinases, and more recently tyrosine and arginine kinases. Tyrosine phosphorylation is today recognized as a key regulatory device of bacterial physiology, linked to exopolysaccharide production, virulence, stress response and DNA metabolism. However, bacteria have evolved tyrosine kinases that share no resemblance with their eukaryotic counterparts and are unique in exploiting the ATP/GTP-binding Walker motif to catalyse autophosphorylation and substrate phosphorylation on tyrosine. These enzymes, named BY-kinases (for Bacterial tYrosine kinases), have been identified in a majority of sequenced bacterial genomes, and to date no orthologues have been found in Eukarya. The aim of this review was to present the most recent knowledge about BY-kinases by focusing primarily on their evolutionary origin, structural and functional aspects, and emerging regulatory potential based on recent bacterial phosphoproteomic studies.

Keywords: tyrosine kinase; bacteria; protein phosphorylation; biological role; structure

1. INTRODUCTION

Since its discovery in the mid-1950s, protein phosphorylation in eukaryotes has been shown to be a versatile and sophisticated mechanism for the regulation of cellular processes such as intercellular communication and coordination of complex functions [1,2]. The presence of this post-translational modification was neglected for a long time in bacteria. Indeed, as early attempts to detect its presence were unsuccessful, it was proposed that protein phosphorylation was a regulatory mechanism that emerged late in evolution to meet the specific needs of organisms composed of multiple and differentiated cells. Protein phosphorylation in bacteria was evidenced in the late 1970s through the pioneering work of several groups that showed the presence of phosphorylated amino acids in bacterial proteins and, in some cases, characterized the phosphorylated proteins [3–5]. Considering that bacteria live in environmental conditions that change frequently and rapidly, they needed to develop devices to monitor their surroundings and adjust their physiology and behaviour accordingly. For that, they possess an arsenal of options, such as moving to more favourable conditions, changing the expression of genes involved for instance in detoxification or nutrient

use, or implementing developmental processes to survive extreme environmental conditions. It therefore came as no surprise that the studies conducted over the next two decades (until the early 1990s) established protein phosphorylation as a major player in bacterial environmental signalling. However, these early studies essentially showed that bacterial signalling and adaptive responses were mainly dependent on histidine kinases [6]. The only exceptions recognized at the time were the three protein kinases sharing no resemblance with eukarvotic ones (the HPr kinase/ phosphorylase, the SpoIIAB from Bacillus subtilis and the isocitrate dehydrogenase kinase/phosphatase from Escherichia coli) catalysing serine phosphorylation [7-9]. During the second half of the 1990s, the availability of complete genome sequences revealed the widespread presence of eukaryote-like serine/threonine protein kinases (STPK) [10]. Like histidine kinases, they have turned out to participate in key regulatory functions of bacterial physiology (for review, see [11]). Surprisingly, eukaryotic-like tyrosine kinases have not been identified in bacterial genomes. Despite the initial detection of phosphotyrosines in bacterial crude extracts and the characterization of two atypical tyrosine kinases possessing some signatures of the eukaryotic kinase domain (WaaP, MasK) [12,13], it has been widely accepted that tyrosine phosphorylation is confined to Eukarya. However, the characterization in 1997 of a peculiar protein autophosphorylated on tyrosine (Ptk from Acinetobacter johnsonii) gave a fresh

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impetus in the field. Ptk shares no sequence homology with eukaryotic tyrosine kinases and possesses the Walker A and B ATP/GTP-binding motifs [14,15]. Ptk homologues have since been biochemically and physiologically characterized in several bacterial species, and the presence of homologous genes has been detected in a majority of sequenced bacterial genomes, but not in Eukarya [16]. Therefore, these enzymes idiosyncratic to bacteria have consequently been unified in the same family named bacterial tyrosine kinases (BY-kinases) [17]. BY-kinases account for the vast majority of tyrosine phosphorylation events detected in bacteria, but there are some other kinases that contribute. These include the protein PtkA of Mycobacterium tuberculosis that does not possess any signature or pattern related to BY-kinases and eukaryotic tyrosine kinases [18], a histidine-kinaselike protein autophosphorylating on tyrosine (DivL from Caulobacter crescentus) and a kinase using a guanidino-tranferase domain that catalyses both arginine and tyrosine phosphorylation (McsB from B. subtilis and Bacillus anthracis) [19-22].

2. CHARACTERIZATION OF FIRST BY-KINASES

The first indication of a protein-tyrosine kinase activity in bacteria was reported in the case of E. coli in which phosphotyrosine was detected in hydrolysates of proteins [23]. This observation was rapidly confirmed in a variety of bacterial species, and in 1996 tyrosine phosphorylation was detected in 22 different bacterial species [24]. However, protein-kinase activities catalysing this reaction were not identified in the initial studies, and proteins phosphorylated on tyrosine were most often characterized only by their molecular mass and isoelectric point. Nevertheless, in some cases, a clear-cut phenotype was associated to tyrosine phosphorylation. For example, it was suggested that tyrosine phosphorylation could be involved, like in eukaryotic cells, in signal transduction, allowing the bacterial cell to adapt to its environment. Indeed, tyrosine phosphorylation was observed to vary during bacterial adaptation to cold and light in Pseudomonas svringae and the cvanobacterium Prochlorothrix hollantica, respectively [25,26]. Phosphotyrosines were detected in pathogenic mycobacteria, but not in non-pathogenic, thus suggesting a possible function in virulence [27]. The identification of certain phosphotyrosine-proteins also suggested that tyrosine phosphorylation could serve other functions involved in metabolism or motility [28,29]. Unfortunately, all these initial studies were not followed up. From the knowledge that we now have about bacterial tyrosine phosphorylation, it would certainly be justified to go back to some of those studies with new methods of the 'omic' era.

The first characterizations of BY-kinases illustrate perfectly that point. In 1993, the group of A. J. Cozzone used massive amounts of radioactive phosphate for *in vivo* labelling and treatment of proteins with bases and acids to detect a tyrosine phosphorylation signal of an 80 kDa protein from *A. johnsonii*. Subsequent purification of a few micrograms of this phospho-protein required litres of cell culture [14,30]. After screening of an *A. johnsonii* genomic library with a degenerate oligonucleotide probe, they successfully identified the gene sequence and named it ptk. The ptk gene encodes a membrane protein capable of autophosphorylating on several tyrosine residues but sharing no resemblance with eukaryotic tyrosine-kinase receptors [15]. Instead of the canonical Hanks-motifs of eukaryotic kinase domains [31], Ptk displays homology with other membrane proteins with Walker A and B ATP/GTP-binding motifs, encoded by genes located in exopolysaccharide synthesis loci. This similarity prompted several groups to test Ptk homologues for their tyrosine-autokinase activity. Thus, proteins Wzc and Etk from E. coli, AmsA from Erwinia amylovora, Orf6 (now renamed Wzc) from Klebsiella pneumoniae, ExoP from Sinorhizobium meliloti and CpsD from Streptococcus pneumoniae were also characterized as possessing tyrosine-autokinase activity [32-35]. However, when analysing the phosphorylation patterns of some of these strains and, for example, E. coli K12, no tyrosine autophosphorylation of Wzc and Etk was detected. etk and wzc genes are located in two distinct capsular polysaccharide loci, expressed either only in pathogenic E. coli strains (for Etk) or dependent upon transcriptional regulators of two-component systems (for Wzc) [33]. Further analyses showed unambiguously that the two-component system RcsABC must be activated to allow Wzc expression and the same effect could be achieved by overproduction of the response regulator RcsA [36]. For Etk, such a clear-cut demonstration has not been made, but its expression requires specific growth conditions, including low pH and low magnesium concentration, that are also required for activation of two other two-component systems, the PmrAB and PhoPQ systems [37].

One now understands better why it was so difficult to characterize the first BY-kinase: it was simply not expressed in standard laboratory growth conditions. Once this expression obstacle had been overcome, BY-kinases-mediated tyrosine phosphorylation was extensively studied *in vivo* over the last decade and shown to be a key regulatory device of bacterial physiology [38]. However, as demonstrated by recent analyses of bacterial phosphoproteomes (discussed below), there are still many uncharacterized substrates of BY-kinases and what we know about their role in bacterial physiology probably corresponds to the tip of an iceberg.

3. EVOLUTION OF BY-KINASES: WHENCE DO THEY HAIL?

BY-kinases clearly share no significant sequence homology with Hanks-type kinases that phosphorylate serines, threonines and tyrosines in Eukarya. However, this does not mean that Hanks kinases evolved in Eukarya after the separation from the bacterial kingdom. Genes encoding Hanks-type kinases are also present in almost all sequenced bacterial genomes, and it is therefore reasonable to assume that they were present before the separation from Eukarya [11] or there was lateral transfer of these genes from a eukaryote. Nevertheless, the bacterial Hanks-type kinases catalyse almost exclusively the phosphorylation of serine and threonine residues, and there are very rare occurrences when they phosphorylate tyrosine [12,13]. One might therefore argue that bacterial Hanks kinases have somehow lost the feature of phosphorylating tyrosines after the separation from Eukarya. But there seems to be no obvious evolutionary benefit gained from such loss of function. A more plausible hypothesis seems to be that ancestral Hankstype kinases, present in the common ancestor before branching off of Bacteria and Eukarya, must have possessed the capacity to phosphorylate serines and threonines, but not tyrosines. After the separation of Eukarya and Bacteria, tyrosine phosphorylation mechanisms appeared in both kingdoms, which effectively means that tyrosine phosphorylation evolved twice, completely independently. In Eukarya, the evolution of tyrosine kinases took the path of modifying the existing Hanks kinases and creating a slightly altered active site that accommodates tyrosines. So what happened in Bacteria? Active sites of contemporary BY-kinases share signatures known as the Walker motifs with a number of ubiquitous ATP/GTPases [17,39]. Walker motifs found in BY-kinases however differ from the canonical sequences found in ATP/ GTP-binding proteins. Indeed, only the GKS/T motif is conserved in the (A/G)xxxxGK(S/T) Walker A motif (phosphate-binding loop or P-loop), whereas the hhhhD canonical Walker B motif is extended to hhhDxxP. A third motif, named A' and located between motifs A and B, is nearly identical to the Walker B motif (hhhhDXDXR). These three motifs have been shown to be essential for the activity of BY-kinases and substitution of either the Walker A lysine, or any aspartic acid of Walker A' and Walker B motifs induced a strong decrease of the autokinase activity of BY-kinases [40,41]. Interestingly, the simultaneous presence of these three Walker motifs is found in certain P-loop kinases of small molecules like the dephosphocoenzyme A, but is absent in the majority of them [42]. Still, they are also found in some ATPases closely related in sequence with BY-kinases and belonging to the SIMIBI class [39]. More precisely, BY-kinases exhibit significant sequence similarity with arsenite ATPases (ArsA) and MinD proteins. ArsA proteins are involved in expulsion of arsenite and antimonite ions [43], whereas MinD is involved in determining the position of the new septum during cell division [44]. The extent of sequence similarity with BY-kinases suggests common evolutionary ancestry and it seems plausible to propose that BYkinases, MinDs and ArsAs evolved by splitting off from an ancestral bacterial ATPase [39]. How soon did this happen, after the separation of Bacteria from Eukarya? To answer this question, we selected a number of bacterial ArsA proteins, MinD proteins and BY-kinases (catalytic domains) and attempted to construct evolutionary trees using different methods. The neighbour joining method produced an evolutionary tree that exhibits three clearly distinct clades corresponding to ArsA, MinD and BY-kinases, and is in more or less perfect accord with the contemporary view on the evolution of bacterial phyla (figure 1a). The BY-kinase and MinD ATPase branches are more closely related, whereas ArsA proteins have clearly

split off from the rest at an earlier point. The maximum-likelihood tree (figure 1b) is in agreement with the neighbour joining tree with respect to positioning of ArsA proteins, but some controversy emerges concerning the separation between MinDs and BYkinases. Most MinD proteins and BY-kinases are grouped in two separate clades. However, some MinD sequences (from Actinobacteria and alpha Proteobacteria) and BY-kinase sequences (from Cyanobacteria, Firmicutes and Actinobacteria) form a third mixed clade (figure 1b, highlighted in orange). It is not uncommon that neighbour joining and maximum-likelihood methods provide evolutionary trees of slightly different topology. Neighbour joining is a simple and robust method, but in a recent benchmark study, maximumlikelihood algorithm was clearly tipped to outperform the competitors with small datasets of under 100 species [46]. Be that as it may, one conclusion that clearly emerges from this analysis is that among the three protein families, ArsA ATPases branched off first, and this node is rooted deeply in the bacterial family tree. By contrast, MinD and BY-kinase families separated more recently and, as suggested by the maximum-likelihood tree, may have still not been fully separated before the branching off of the main bacterial families as we know them today. This observation is in accord with the report by Leipe and co-workers [42] who found that the Walker A' motif (DXD; see \S and 5) is present in MinD ATPases but is not fully conserved in ArsA ATPases. BY-kinases contain the canonical Walker A', which again puts their evolutionary origin closer to MinDs rather than to ArsAs. When considering the evolution of such closely related protein families, one is well advised to keep an open mind concerning the specific protein functions. A onegene-one-function paradigm may well prove to be too simplistic to account for these events. BY-kinases are even today still described as polysaccharide co-polymerases by the researchers interested in polysaccharide biosynthesis [47], and are called BY-kinases by those interested in bacterial signalling and tyrosine phosphorylation of protein substrates [17]. Moreover, BYkinases have phenotypes related to the cell cycle [48] and it would not be utterly unexpected if one day a direct mechanical interplay between BY-kinases and closely related MinD proteins were to emerge in the context of bacterial cell cycle control. This notion is strengthened by the fact that in Firmicutes, where BYkinases have evolved with an option to dissociate from their trans-membrane modulators, BY-kinases were in some growth conditions found to co-localize with cytosolic substrates rather than with the trans-membrane modulators [49].

4. DESCRIPTION OF BY-KINASES

As indicated above, BY-kinases possess Walker A and B motifs [50], which are characteristics of nucleotide-binding proteins (NTPases) called P-loop proteins [39,42,51]. Because of that, they were initially thought to function as ATPases in the extracellular polysaccharide synthesis pathway and it is likely that this idea would have persisted for a long time without the characterization of the autokinase activity of Ptk



Figure 1. Evolutionary relationship among BY-kinase, ArsA and MinD protein families. Protein sequences were obtained from the following species. BY-kinases (catalytic domain): Bacillus subtilis, Bacillus halodurans, Escherichia coli, Klebsiella pneumoniae, Pelobacter propionicus, Geobacter bemidjiensis, Sagittula stellata, Roseovarius sp., Acaryochloris marina, Microcoleus chthonoplastes, Aeromonas hydrophila, Raphidiopsis brookii, Clostridium bolteae, Lactobacillus reuteri, Staphylococcus aureus, Desulfobacterium autotrophicum, Bifidobacterium angulatum, Kytococcus sedentarius and Janibacter sp. ArsA ATPases: Lactococcus lactis, Clostridium botulinum, Listeria monocytogenes, Enterobacter cloacae, Anabaena virabilis, M. chthonoplastes, Acaryochloris marina, Intrasporangium calvum, Rubrobacter xylanophilus, K. pneumonia, E. coli and B. halodurans. MinD proteins: B. subtilis, B. halodurans, E. coli, K. pneumonia, Sagittula stellata, Roseovarius sp., R. xylanophilus, I. calvum, Acaryochloris marina, Microcoleus chtonoplastes, Prochlorococcus marinus, C. butyricum, Yersinia pestis, Rhodopseudomonas palustris, Methylocella silvestris and Bradyrhizobium japonicum. (a) Neighbour joining tree obtained with BioNJ. BY-kinases are highlighted in pink, ArsA proteins in blue and MinD proteins in yellow. (b) Maximum-likelihood tree obtained with PhyML. Same colour-coding was used; a clade with mixed BY-kinases and MinD proteins is in orange. Alignments were prepared with CLUSTALW and trees were visualized using TREEDYN. All cited software is available at http://www.phylogeny.fr/ [45].

from A. johnsonii. An additional motif located at the very C-terminal end of the kinase domain further distinguishes BY-kinases from other P-loop proteins [52]. This motif, named the tyrosine cluster (YC), varies in length (from 10 to 20 amino acids) and contains several tyrosine residues (from three to seven) that constitute the autophosphorylation sites of BYkinases. Albeit designated as autophosphorylation, the event actually necessitates at least two BY-kinase molecules, one phosphorylating the tyrosine cluster of the other, and vice versa. It is therefore also referred to as trans-phosphorylation. The sequence of the tyrosine cluster motifs is not conserved among BY-kinases. It has been shown that each tyrosine could be phosphorylated independently of the others without any preferential order [53]. The presence of these four motifs (Walker A, Walker A', Walker B and YC) was used to constitute a typical signature of BY-kinases [16]. Bioinformatics resources allowed the annotation and collection of more than 2000 genes in sequenced bacterial genomes that are proposed to belong to the family of BY-kinases.

A database (BYKdb) dedicate to BY-kinases is available at http://bykdb.ibcp.fr/BYKdb/ [54].

In Proteobacteria, BY-kinases are essentially found in the form of a membrane protein displaying a large extracellular loop and a cytosolic catalytic domain [17]. By contrast, BY-kinases of Firmicutes possess the cytoplasmic catalytic domain in a separate polypeptide that interacts with a separate membrane protein, homologous to the extracellular domain of proteobacterial BY-kinases, but shorter in size [55]. These two polypeptides are encoded by adjacent genes in Firmicutes. Firmicute-type cytosolic BY-kinases, however, require the membrane protein to undergo autophosphorylation [55]. More precisely, the cytosolic and C-terminal end of the membrane protein (about 30-50 residues) mimics the juxtamembrane region of BY-kinases from Proteobacteria and is sufficient to promote BY-kinase activation [56]. In agreement with this observation, the isolated cytoplasmic domain of a BY-kinase from Proteobacteria is also active in vitro. The cause and functional consequences of this separation remain unclear, although some recent leads suggest that it could allow



Figure 2. BY-kinase architectures among bacterial phyla. The number of BY-kinases predicted in the BYKdb [54] for each bacterial phyla is indicated in black. Those displaying a Firmicute-type organization are numbered in green, whereas those with a Proteobacteria-type organization are numbered in red.

interaction with other proteins to participate in additional regulatory functions. The two structural prototypes, Firmicute- and Proteobacteria-type architectures, are not strictly confined and some representatives of both can be found in other bacterial phyla, as predicted in BYKdb (figure 2).

5. STRUCTURAL ASPECTS OF BY-KINASES

The first structural insights concerning BY-kinases came from the low-resolution electron microscopy structure (14 Å) of the *E. coli* BY-kinase Wzc [57]. Wzc was described as 'molar tooth and its roots' forming a tetrameric ring that oligomerizes *via* the periplasmic domains and with non-interacting cytosolic domains freely protruding in the cytosol. The same organization was observed in the electron microscopy structure of Wzc in complex with the outer-membrane translocon Wza [58]. The Wzc–Wza complex formed a ring-shaped structure spanning the periplasm and connecting the inner and outer membranes of the bacteria. However, this structural organization did not explain the trans-autophosphorylation mechanism demonstrated previously [52].

More recently, the crystal structures of the cytoplasmic domains of Etk, a paralogue of Wzc in E. coli [59], and of the orthologue CapB from the Firmicute Staphylococcus aureus [53] have been solved. Both structures displayed a Rossman fold typical of P-loop proteins [51]. Despite catalysing the same ATP-dependent phosphorylation of tyrosine residues, Etk and CapB did not show any structural similarity with eukaryotic tyrosine kinases [60]. CapB was fused to the C-terminal extremity (29 residues) of the transmembrane activator CapA that was shown to enhance the affinity of CapB for ATP and to stimulate its kinase activity [56]. When produced as a His-tagged recombinant protein in E. coli, Etk was phosphorylated, as well as the CapAB chimaera. Mass spectrometry analysis demonstrated that the four tyrosines of the CapB Y-cluster were heterogeneously phosphorylated [53]. In both structures, the crystallized kinase domain was monomeric and the phosphorylated Y-cluster was either disordered or involved in nonphysiological crystal packing interactions. Again, these two structures did not give any insight into the transautophosphorylation mechanism. They however gave

some interesting clues on the activation mechanism of BY-kinases from Firmicutes and Proteobacteria. As usually observed in P-loop proteins, conserved lysine and threonine residues from the Walker A motif and the aspartates from the Walker A' and B motifs interacted with the phosphate moiety of the bound nucleotide and the associated magnesium ion (figure 3). Interestingly, the juxtamembrane N-terminal region of Etk_{cyto} superimposed with the C-terminal fragment of the trans-membrane activator CapA in the structure of the CapAB chimaera. The side chain of the penultimate residue of CapA, Phe221, was stacked on the base part of the bound ADP molecule, making a strong hydrophobic interaction stabilizing nucleotide binding and thus explaining the activation mode of CapB (figure 3). The C-terminal sequence of CapA orthologues is poorly conserved and it is not clear if this activation mechanism is conserved in all BY-kinases from Firmicutes. In Etk, the adenine base is stacked on the side chain of a phenylalanine residue located in another region of the protein, directly following the Walker A motif. This residue is conserved among BYkinases from Proteobacteria, but not from Firmicutes, suggesting a distinct nucleotide-binding mode. Another difference observed between the two phyla concerns a short stretch of about 20 basic residues, called the RK-cluster, present in Etk but not in CapB [59]. The RK-cluster is 'inserted' in the region corresponding to the junction between CapA and CapB. The RK-cluster is also observed in other BY-kinases from Proteobacteria and it has been shown to be essential for the function of Wzc. Its N-terminal part is involved in the binding of the base part of the nucleotide whereas its disordered C-terminal part has been shown to be essential for the co-polymerase activity of Wzc in the exopolysaccharide synthesis machinery [61]. Its exact role, however, remains to be elucidated. Etk, like its paralogue Wzc, possesses outside its C-terminal Y-cluster an additional tyrosine that had been suggested to be involved in an intramolecular phosphorylation step, preceding transautophosphorylation of the Y-cluster [52]. The analysis of the Etk structure suggested that that this tyrosine points into the active site, and once phosphorylated it could flip out of the active site to allow ATP binding and subsequent phosphorylation of the Y-cluster [59,62]. However, its orientation seems quite incompatible with phosphate transfer, and this residue is not conserved



Figure 3. Close view of the CapAB active site. Bound $ADP-Mg^{2+}$ is shown as sticks in the P-loop (in yellow) with the inactivating K55M mutation of the Walker A motif. The phosphorylatable residue Y225* from the neighbouring Y-cluster is highlighted as well as residues D157 and P160 from the extended Walker B motif and residues D77, D79 and R81 from the Walker A' motif. Residue R81 together with the conserved residues S132 and E133 participate in octamer contact with residues E23*, R26* and R29* from the neighbouring subunit. The adenine base of the bound nucleotide is shown in stacking between residues R212 and the penultimate residue F221 from the CapA activator (in blue).

among all BY-kinases from Proteobacteria. Therefore, this putative additional phosphorylation step remains controversial and needs to be further investigated.

In order to decipher the trans-autophosphorylation mechanism of the Y-cluster, inactive mutants of the cytoplasmic domains of Wzc [61] and of the chimaera CapAB [53] were used. The P-loop lysine was replaced by a methionine, a mutation known to inhibit phosphate transfer without impairing ATP binding in P-loop proteins [63]. The crystal structures of these two nonphosphorylated kinase domains revealed the formation of a conserved ring-shaped octamer. In both cases, the last tyrosine of the Y-cluster of each subunit was bound into the active site of the neighbouring subunit, pointing towards the bound nucleotide, and thus explaining the trans-autophosphorylation mechanism of the BY-kinases (figure 3). The aromatic ring of the tyrosine is in a sandwich interaction between the proline residue of the extended Walker B motif and a lysine residue immediately following the Walker A' motif of the neighbouring monomer. The second aspartate of this hhhhDXDXR motif directly interacts with the phosphorylatable hydroxyl of the tyrosine, most probably acting as an acid catalyst (figure 3). The flexibility of the C-terminal extremity is in agreement with the heterogeneous phosphorylation pattern observed by mass spectrometry but it is not clear why in both crystal structures the last tyrosine of the Y-cluster was preferentially bound into the neighbouring active site. The interaction of the Y-cluster of one subunit with the active site of the neighbouring subunit represents 50 per cent of the surface contact between two monomers. The second half of the interface encompasses the conserved arginine of the Walker A' motifs and played a crucial role in stabilizing the octamer (figure 3) [53,61]. This region of contact involves others residues that are conserved among all bacteria, suggesting that the octamer is a conserved feature of BY-kinases.

Taken together, the crystal structures of the phosphorylated and non-phosphorylated forms of the BY-kinases suggest that the kinase domains dissociate upon phosphorylation, thus allowing endogeneous substrate proteins to interact with the active site of BY-kinases. However, in Proteobacteria, BY-kinases are trans-membrane proteins and the N-terminal periplasmic domains most probably remain associated in an oligomeric ring [64]. The co-polymerase function of the BY-kinases in the exopolysaccharide synthesis machinery has been shown to be dependent on a cyclic process oscillating between the non-phosphorylated and the phosphorylated form of the protein (see later text). This would thus correspond to cycling between a closed octameric ring-shaped conformation and an open conformation with independent cytoplasmic kinase domains linked to a trans-membrane crown, as observed in the low-resolution structure of phosphorylated Wzc [57]. In Firmicutes, the separation of the two domains into distinct polypeptide chains suggests that the phosphorylated kinase domains could dissociate from their cognate transmembrane activator and interact freely with the endogenous substrates and/or cytosolic modulators.

6. COMPARISON WITH BACTERIAL ATP-BINDING PROTEINS

As indicated already, the closest structural homologue of BY-kinases is the bacterial cell division regulator MinD, which was successfully used as initial model for the resolution of the structure of Staphylococcus aureus CapAB. The Y-cluster of the BY-kinases has however no equivalent in the ATPase MinD, which does not autophosphorylate and has never been reported to phosphorylate any other protein. Interestingly, the fold is conserved between the two proteins with an r.m.s. distance of 1.66 Å over 158 aligned Ca atoms, but the C-terminal region of MinD superimposes with the N-terminus of the BY-kinase (figure 4). Such circular permutations can appear in evolution upon duplication of a precursor gene, when one or the other extremity of the resulting tandem protein is lost after introduction of a stop codon [65]. By contrast, BY-kinase structure is much more different from that of an atypical bacterial serine kinase, the bifunctional HPr kinase/phosphorylase (HPrK/P), which belongs also to the P-loop proteins. Indeed, the catalytic domain of HPrK/P displays a Rossman fold with an unusual topology similar to that already observed in the phosphoenolpyruvate carboxykinase [66] and forms a dimer of trimers surrounded by three dimers of N-terminal domains of unknown function [67]. The similarity between the HPrK/P and BYkinases is therefore limited to the conserved P-loop type nucleotide-binding domain (figure 4) and they consequently belong to different classes of P-loop proteins. Disappointingly, the sequence and structure comparison of the BY-kinases with MinD ATPases and the HPrK/Ps does not provide any clear clues for distinguishing between P-loop proteins with and



Figure 4. Structural comparison of CapAB with MinD and HPrK/P. The core of the BY-kinase CapAB conserved with MinD and HPrK/P is shown in green with the P-loop in yellow and the ADP-Mg in orange. (a) CapAB with the Y-cluster in red, the CapA fragment in blue and the N-terminus of CapB in cyan. (b) MinD with the C-terminal extremity superimposing on the CapA fragment and N-terminus of CapAB in red. (c) Catalytic domain of *Lactobacillus casei* HPrK/P with the conserved P-loop core in green and yellow (in the absence of nucleotide).

without protein kinase activity. It would be of utmost importance to find such clues in the future, because P-loop proteins are one of the largest protein families in all kingdoms of life, representing for instance 5 per cent of the human genome [68].

7. BIOLOGICAL ROLE OF BY-KINASE AUTOPHOSPHORYLATION

In the first phase of their characterization, BY-kinases were shown to function essentially as autophosphorylating enzymes and this feature was associated with

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their role in the biosynthesis and export of extracellular polysaccharide. Although the different groups working in the field agreed that the autokinase activity was required in this process, there were initially some apparent discrepancies. Thus, in E. coli K30, a positive regulation of the group I polysaccharide capsule biosynthesis by autophosphorylation of Wzc was observed [32], whereas in E. coli K12 production of the exopolysaccharide colanic acid was decreased when Wzc was phosphorylated [36]. When analysing the Gram-positive bacterium S. pneumoniae, a similar divergence was reported. In the RX1 strain, the BY-kinase CpsD was shown to be dephosphorylated to promote capsule synthesis, whereas an opposite situation was found in the S. pneumoniae D39 strain [34,69]. In Acinetobacter lwoffii, emulsan polysaccharide is produced to a relatively low extent when Wzc is phosphorylated, whereas in E. amylovora the overproduction of the low-molecular-weight-phosphatase AmsI induces a strong reduction in the exopolysaccharide amylovoran synthesis [70,71]. However, the identification of several bacterial phosphotyrosine phosphatases crucial for polysaccharide production shed new light on the regulatory function of BYkinases in polysaccharide production (for example, the low-molecular-weight acidic phosphotyrosine phosphatase Wzb of E. coli and the manganese-dependent phosphotyrosine phosphatase CpsB of S. pneumoniae [72,73]). Hence, it was shown that both the phosphorylated and the non-phosphorylated forms of BY-kinases were alternately required for polysaccharide polymer production, explaining thus the divergent initial observations [74]. On this basis, the current model is that cycling between phosphorylated and non-phosphorylated forms of BY-kinases is required for proper synthesis and export of the polysaccharide polymer. According to their regulatory function in polysaccharide production, BY-kinases were classified as polysaccharide co-polymerases assisting the activity of the polymerase of the polysaccharide assembly membrane machinery [75]. However, the detailed mechanism by which BYkinases interact with this machinery remains obscure. Several hypotheses have been proposed. Structural characterization of BY-kinases CapB and Wzc from Staphylococcus aureus and E. coli, respectively, showed that the non-phosphorylated form of BY-kinases forms a ring-shaped octamer which upon autophosphorylation dissociates to monomers. The monomers would remain in interaction with the trans-membrane domains. Therefore, an interesting possibility would be that BY-kinases might simply influence the amount of polysaccharide by serving as molecular scaffolds for the other proteins of the machinery (figure 5) [47]. Phosphorylation/ dephosphorylation of BY-kinases would thus behave as a conformational switch that would most probably be transmitted to their trans-membrane domain. In turn, the latter would be affected in their interaction with other protein components of the polysaccharide assembly machinery, such as the polysaccharide unit polymerase, the lipid-linked repeat unit flippase or the lipid-sugar transferase [53]. Alternatively, one should also consider that BY-kinases could form a channel across the cytoplasmic membrane, large and hydrophilic enough to allow the polysaccharide polymer to cross the



Figure 5. Role of BY-kinase-mediated phosphorylation in the bacterial cell. The polysaccharide membrane assembly machinery is shown as salmon-pink ovals. BY-kinases (blue oval) form a ring-shaped octamer, which upon autophosphorylation dissociates to monomers. This conformational switch is responsible for controlling polysaccharide biosynthesis and export, but the underlying mechanism remains unclear. Three possible scenarios, are shown with arrows numbered as followed: (1) targeting to the septum; (2) scaffolding of the assembly machinery; (3) forming a channel across the cytoplasmic membrane. Enzymes involved in polysaccharide unit synthesis are shown as a salmon-pink hexagon. Their phosphorylation by BY-kinases regulates the production of the polysaccharide polymer. Multicoloured hexagons represent proteins being phosphorylated by BY-kinases, and/or eukaryotic-like serine/threonine kinases (green oval) and/or histidine kinases of the twocomponent systems (fuchsia oval). Arrow 4 shows that two-component systems could regulate the expression of BY-kinases. Question mark shows the hypothetical stimulus triggering the activity of BY-kinase. The activation of cytoplasmic BY-kinases by proteins (red half-oval) distinct from their transmembrane membrane (dark blue half-oval) is also shown.

membrane [61]. Although cryo-electron-microscopic studies made with Wzc of E. coli K30 suggested that this channel would be filled by a lipid bilayer [57], one should note that in Gram-negative bacteria another protein is required for the polysaccharide to go across the outer membrane. This protein, named Wza in E. coli, also forms an octameric complex and was shown to interact directly with the Wzc oligomer, thus forming a complex that spans both the inner and the outer membranes [58,76]. This finding might suggest that the Wza-Wzc complex could form an octameric channel, directly constituting the polysaccharide secretion system (figure 5). This hypothesis is further supported by the structural similarities observed between the extracellular domain of BY-kinases and another class of polysaccharide co-polymerase involved in lipopolysaccharide biosynthesis [77]. These latter proteins were shown to regulate the length of O-antigens and they self-assembled into ring-shaped oligomers of variable sizes forming a large internal cavity [64]. BY-kinases would therefore play a more central role than assisting the Wzy polymerase in the translocation of capsular polysaccharides. Very recently, another study added a new facet to the role of BY-kinases in extracellular polysaccharide production. The BY-kinase CpsD that was previously found to be required for *S. pneumoniae* capsular polysaccharide production has been shown to localize at the bacterial division septum [78]. The authors further showed that in the absence of CpsD (named Wze in this study), capsular polysaccharides were still produced, but absent from the division septum. On this basis, it could be presumed that BY-kinases might also function as spatial regulators of capsular polysaccharide biosynthesis (figure 5). Considerable efforts are currently directed at understanding the exact role played by BY-kinases in polysaccharide export and the associated molecular mechanisms.

8. BY-KINASE-MEDIATED PHOSPHORYLATION OF BACTERIAL PROTEINS

Beyond their autophosphorylation, BY-kinases were shown to phosphorylate endogenous protein substrates on tyrosine residues and consequently to control some cellular processes. The first identified BYkinase substrates were, quite serendipitously, found to participate in polysaccharide production (figure 5). They are the uridine diphosphate (UDP)-sugar dehydrogenases, and their phosphorylation by BY-kinases has been documented in E. coli, B. subtilis and Staphylococcus aureus [37,55,79,80]. Upon phosphorylation, their dehydrogenase activity is enhanced. In E. coli, Wzc-mediated phosphorylation of Ugd has further been shown to regulate the amount of the exopolysaccharide colanic acid produced, whereas when phosphorylated on the same tyrosine residue by Etk, the synthesis of a modified sugar (UDP-4-amino-4deoxy-L-arabinose) is increased. This sugar is a crucial element in bacterial resistance to antibiotics such as polymyxin and cationic peptides of the innate immune system, and Etk-mediated phosphorylation of Ugd was shown to efficiently induce bacterial resistance [37]. Ugd phosphorylation by the two BY-kinases Wzc and Etk is therefore at the crossroads of two distinct biosynthetic pathways, illustrating thus the regulatory potential of tyrosine phosphorylation in bacterial physiology. The molecular mechanism of activation of UDPsugar dehydrogenases by BY-kinases remains not clearly understood and it could involve precise positioning of either the catalytic amino acid or the phosphorylated tyrosine with respect to the bound substrate in the active site [81]. To illustrate this, the recent characterization of the structure of CapO of Staphylococcus aureus suggests that the phosphorylated tyrosine could participate in the correct positioning of the catalytic cysteine residue [82]. Other proteins involved in polysaccharide production were shown to be phosphorylated by BYkinases. One could cite the Streptococcus thermophilus phospho-glycosyltransferase EpsE and the K. pneumoniae undecaprenolphosphate glycosyltransferase WcaJ that are necessary for capsular polysaccharide production [83,84]. However, in the absence of structural information, the activation molecular mechanism remains unknown. Nevertheless, note that by contrast with UDP-sugar dehydrogenases, which retain basal activity upon Tyr to Phe mutations at the phosphorylation site, mutation of the phosphorylation site of the phospho-glycosyltransferase EpsE seems to induce its inactivation.

BY-kinases are also capable of phosphorylating other endogenous proteins substrates. The BY-kinase Etk was found to phosphorylate the heat-shock sigma factor σ 32 and the antisigma σ E factor RseA and thus to participate in the heat shock response [85]. Similarly, the BY-kinase Wzc was found to phosphorylate the integrase (Int) of certain coliphages, thus suggesting that lysogenization could be regulated by Wzc-mediated phosphorylation of Int [86]. Several BY-kinases, not encoded by genes of polysaccharide loci, were also shown to phosphorylate diverse proteins or to be involved in other cellular processes. The BYkinase PtkA of B. subtilis was shown to phosphorylate several proteins in addition to the UDP-sugar dehydrogenases TuaD and Ugd. Single-stranded DNA binding proteins SsbA and SsbB proteins are phosphorylated on tyrosine and this stimulates their ability to bind DNA [87]. Considering that cells deficient for PtkA expression exhibited defects in the initiation of DNA replication and chromosome distribution, PtkA could be involved in regulating DNA metabolism [48]. The single-stranded DNA exonuclease YorK and the aspartate semialdehyde dehydrogenase Asd are also

activated by PtkA-dependent phosphorylation [49]. A variety of proteins phosphorylated by PtkA are still of unknown function, but, interestingly, a number of them displayed aberrant cellular localization in cells deficient for PtkA expression [49]. This suggested that besides controlling enzymes activity, BY-kinases would have another mode of action by ensuring proper localization of their targets. In the coming years, the use of cell biology approaches should help us to understand the role of BY-kinase mediated tyrosine phosphorylation and decipher the underlying regulatory processes.

9. FUTURE PROSPECTS AND CHALLENGES

In the last 15 years, evidence was provided that tyrosine phosphorylation of proteins is an important post-translational modification involved in the regulation of different biological processes in bacteria [88]. BY-kinases are widespread and account for most tyrosine phosphorylation events in bacteria. Nevertheless, numerous aspects remain obscure and it seems that we are only at the onset of understanding the biological significance of tyrosine phosphorylation.

One big question concerns the regulation of BYkinase activity. The presence of a large extracellular loop suggests, by analogy to eukaryotic tyrosine kinases receptors, that the outside domain of BY-kinase could transmit signals, thus affecting the kinase function. Unfortunately, such signals remain to be identified. Generally speaking, extracellular signals are known for some two-component systems, but are unknown for a vast majority of other bacterial protein kinases [89]. The interaction of the extracellular BY-kinase domain with the polysaccharide translocation complex may prevent it 'sensing' other signals. On the other hand, one can reasonably argue that the very interaction with the translocation complex could indeed act as a signal triggering BY-kinase activity. A recent study showed that distinct mutations in the extracellular domain of the BY-kinase ExoP of Sinorhizobium meliloti either increased or abolished ExoP autophosphorylation [90]. It suggested that tyrosine autophosphorylation of ExoP could be modulated by conformational changes of the N-terminal domain possibly mediated by protein-protein interactions inside the polysaccharide assembly complex. Nevertheless, this possibility is unlikely to be valid for the phosphorylation of the other BY-kinase substrates not involved in polysaccharide production. Structure predictions showed that the extracellular domain of BY-kinases from Proteobacteria would possess more β -structures than the extracellular domain of BY-kinases from Firmicutes [77]. It is tempting to speculate that this feature would confer an additional function to the periplasmic domain of BYkinases of Proteobacteria, namely a sensor activity, to allow phosphorylation of other targets not involved in polysaccharide production. In Firmicutes, the 'split in two' BY-kinase organization would allow the cytoplasmic domain to interact with other activating proteins. This latter possibility is supported by the finding that under certain growth conditions, the BY-kinase PtkA dissociates from the cognate membrane domain, becomes cytosolic and interacts with other cytosolic proteins that are not its substrates, but modulate its

| species | proteins | functions | peptide sequences | |
|--------------------------|------------------|--|---|--|
| Escherichia coli | AsnS | asparaginyl-tRNA synthetase | KFENPV _p YWGVDLSSEHER | |
| | Dps | DNA protection during starvation protein | SpYPLDIHNVQDHLK | |
| | GroS | chaperonin, small subunit of GroESL | VGDIVIFNDGpYGVK | |
| | Ppa | inorganic pyrophosphatase | AQIAHFFEHpYK | |
| | T_{naA} | tryptophanase/L-cysteine desulfhydrase | NIFGpYQYTIPTHQGR | |
| | TufA | elongation factor Tu | GYRPQFpYFR | |
| | YgfB | predicted protein | NIAQLGYDEDEDQEELEMSLEEIIEpYVR | |
| | ${ m RpsD}^{a}$ | 30S ribosomal subunit protein S4 | VVNIAp(SY)QVSPNDVVSIR | |
| | | | pSDLSADINEHLIVELp(YS)K | |
| | RpsE | ribosomal subunit protein S5 | IFpSFpTALTVVGDGNGRVGFGpYGK | |
| | RpsG | ribosomal subunit protein S7 | VGGpSpTpYQVPVEVRPVR | |
| | RpsL | ribosomal subunit protein S12 | VpYTpTTPKKPNpSALRK | |
| | RpsR | ribosomal subunit protein S18 | NpYIpTEpSGKIVPSR | |
| | $RplE^{a}$ | ribosomal subunit protein L5 | LMpTEFNp(YNS)VMQVPR | |
| | : | - | GNp(YS)MGVREQIIFPEIDYDKVDR | |
| | KpIM | ribosomal subunit protein $L13$ | DWpYVVDAIGK | |
| | | | HKAEpYp(TPHVDT)GDYIIVLNADK AFn(YT')PHVDnTGDnYIIVI NADK | |
| | đ | | | |
| | KpIr DalI I | ribosomal subunit protein L13 | VLPYEMDGVFEELAK Mavanforggy | |
| | RpIV | ribosomal subunit protein L22 | VSOALDILDToYoTNK | |
| | | | | |
| Dacutus subturs | Aupr | aikyi iiyuroperoxide reduciase (large subuliit) | | |
| | ASU T | aspartate-semiatoenyde denydrogenase | | |
| | Eno" | enolase | P(YPIIS)IEDGLDENDWEGHK | |
| | InfA | initiation factor IF-I | VTVELSPpYDLTR | |
| | LctE | L-lactate dehydrogenase | NAApYhIIEK | |
| | OppA | oligopeptide ABC transporter (binding lipoprotein) | DGSLHVEPIAGVp(YWY)K | |
| | $RocA^{a}$ | delta-1-pyrroline-5 carboxylate dehydrogenase | p(TVTY)AHEPFTDFTEAK | |
| | YjoA | conserved hypothetical protein | EMGHTELPFpYQQR | |
| | YnfE | conserved hypothetical protein | MDEILKQYMVLpYKK | |
| | YorK | single-strand DNA-specific exonuclease | EFILQLDGpYK | |
| | \mathbf{Y} vyG | putative flagellar protein | pYIQAITQTEDDRIK | |
| | $ m YwqF^{a}$ | UDP-glucose dehydrogenase | LNFETp(SY)EK | |
| Streptococcus pneumoniae | Tuf | translation elongation factor Tu | FKGEVpYILTK | |
| | PgK | phosphopglycerate kinase | VLIGGGMTYTFpYKAQGIEIGNpSLVEEDK | |
| | PfkA | 6-phosphofructokinase | MEDIVApSIKAGpYECGKK | |
| | PfiB | formate acetyltransferase | ISGpYCVNpTKpYLTPEQK | |
| | GlnA | glutamine synthetase, type I | HAPYNYPTAIMNPpTVNSYKR | |
| | PurA Set-D | adenylosuccinate synthetase | pYQGGDNAGHpTIVIDGKK | |
| | | Serine/Inreonine protein kinase | | |

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| species | proteins | functions | peptide sequences |
|--|---|--|--|
| | PrfA | petide chain release factor 1 | pYAEAQGWRFEVMEApSMNGVGGFKEVVAMVSGQSVYpSK |
| | CbpC | choline binding protein C | TYYFGDIGEMVVGWQYLEIPGTGpYR |
| | Spr0084 | conserved hypothetical protein | pYVpSKNELTQAEVIER |
| | InfA | initiation factor IF-I | NpYIRILAGDRVTVEMSPYDLTR |
| | FtsA | cell division protein | LEMRGLLpYpTGPR |
| | LytA | autolysin | pTGWVKpYKDpTWYYLDAK |
| Streptomyces coelicolor A3 | SCO3843 ^a | hypothetical protein | PGGp(YGY)PQPAGGQQPPAAPAPGGR |
| | SCO4973 | putative uncharacterized protein | pYGGTEVK |
| | SCO5128 ^a | putative membrane protein | GGAAp(YPSGT)PPYGTPTASDAGADAGRKTEp(LTT)R |
| | SCO5475 | hypothetical protein | EAHAPAPAQAPAGGp(YGY)PPAGSR |
| | NarH | putative nitrate reductase | YLGVVL _p YDADK |
| Lactococcus lactis | FusA ^a | elongation factor G | LYDGp(SY)HDVDSSETAFK |
| | GinA | glutamine synthase | RLVPGYEAPVpYVAWAGR |
| | GroEL ^a | Chaperonin GroEL | Pp(SY)GSPLJTNDGVTIAK |
| | MalE ^a | ABC-type sugar transport system | TPAp(SQY)DSQLK |
| | PknB | serine/threonine protein kinase | IMPANLINPpYDpTKPLIDKK |
| | Pta ^a | phosphotransacetylase | IAQRLGNFEAIGPILQGLNAPVp(SDLSRGSNEEDVY)K |
| Klebsiella pneumoniae Helicobacter pylori | AhpC atpD ^a CsrA CsrA CsrA CsrA CsrA GalU ManB ManB ManB Rpa Rpa Rpa Rpa Rpa Rpa Rpa Rpa Rpa Rpa | alkyl hydroperoxide reductase ATP synthase subunit beta carbon storage regulator deoxyribose-phosphate aldolase elongation factor P type-1 fimbrial protein, A chain UTP-glucose-1-phosphate uridylyltransferase phosphomannomutase phosphomannomutase phosphoenolpyruvate synthase 50S ribosomal protein L21 DNA-directed RNA polymerase, beta subunit DNA-directed RNA polymerase, beta subunit 30S ribosomal protein S9 putative UDP-glucose lipid carrier transferase putative prophage baseplate assembly protein putative uncharacterized protein gputative uncharacterized protein putative uncharacterized protein | AAQpYVASHPGEVCPAK AAPp(SY)EELSSSQELLETGIK IQAEKSQQSSpY AAIApYGADEVDVVFPYR ATpYYSNDFR YpYATGAATAGIANADATFK GLGHAVLCAHPVVGDEPVAVILPDVILDEpYESDLSR GELGEELNEDIApYR DALpYNGSLLIR pYAALCDVFVMDAFGTAHR LpYTSLGDAAWGR MpYAVFQSGGK MpYAVFQSGGK P(STGSY)SLVTQQPLGGK IApYNVEAAR AENQpYYGTGR AENQpYYGTGR AENQpYYGTGR AFNQpYYGTGR ATLpYHR FGPDPKPFSpY FNALpYEAIHVQGAK SSNNpYLGAAR AAHDFpY GpYLIPK FKDNPPYFPTPLHPLLKDK |
| | VII:D4 | DNA liausief pioteili | (Continued.) |

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Table 1. (Continued.)

| species | proteins | functions | peptide sequences |
|------------------------|---|---|--|
| | SpeA ^a Dxs SpoT UvrD RpoB Rco57lr PepQ Tgt HP0468 HP0112 HP0408 HP0408 HP0586 HP1055 ^a | arginine decarboxylase deoxyxylulose-5-phosphate synthase penta-phosphateguanosine-3'-pyrophosphohydrolase DNA helicase II DNA-directed RNA polymerase, beta subunit Type IIS restriction enzyme R and M protein proline dipeptidase tRNA-guanine transglycosylase conserved hypothetical protein conserved hypothetical protein conserved hypothetical protein conserved hypothetical protein conserved hypothetical protein conserved hypothetical protein | GLN-4p(YGLEAGSKSELIIAMSY) TNPK KSKSAILSPTEApYSNTLLELAK NECNIVGVpSpYLGYK p(TLTS)RLApYLIGVCGVPp(SENTLTLTFTNKAS)K pYpTVREAMER LpYQELKQNPNK pYSLEGKITEGFSDDGEPKGp(SS)GMPMLSVLR YpSKApYLHHLFRAK GVSpYNENIALK HNRLLPRDpYLGpYR pTLpYAFPSLIDSQYCSK DNApYVFRIKDLK p(YFVSYFGTRFY)GDLLLGGGALK |
| Pseudomonas putida | HrpA TxrB PfI01_2957 CcmE PgaA ^a FlgL ^a PfI01_2213 PfI01_2213 | ATP-dependent helicase thioredoxin reductase pyridine nucleotide-disulfide oxidoreductase cytochrome c-type biogenesis protein predicted outer membrane protein flagellar hook-associated protein amino acid adenylation multidrug efflux protein | DLQLSLNKEPADpYPKLHK VIILGSGPAGYSAAVpYAAR MLDANLKAQLKSpYLER TVTTTpYR AGHp(YT)PALSVLR VVLTNGIVGp(T)Aap(TY)NAK FLDDPFNSGRMpYR LPQDAEDPVLSKEAADASALMpYISFFSK |
| Pseudomonas aeruginosa | ThiC PA3965 SucC PvdL SbcB PA0920 PA3238 PA3347 | thiamin biosynthesis protein probable transcriptional regulator Succinyl-CoA synthetase beta chain probable non-ribosomal peptide synthetase exodeoxyribonuclease i hypothetical protein hypothetical protein | GIITPEME _p YIAIRENMK pYDLRILEELQR NLVT _p YQTDANGQPVSK MWFLWQLEPDpSPApYNVGGLARLSGPLDVAR pYRARNFPEpTLNVAER LJIDRITApYR pTFpYEVVLSK NATpYLDSSALGML1LLR |
| Campylobacter Jejuni | Cft Cj0691 Thij | ferritin hypothetical protein 4-Me-5(β-hydroxyethyl)-thiazole phos. synthesis | T _p YEHEQFI _p TK pYNEVCEANEALR KLCGDEI _p YQK |
| Listeria monocytogenes | Atpa1 FusA Imo0208 Imo0310 Imo1871 Imo2324 Imo2373 | ATP synthase subunit alpha 1 elongation factor G UPF0145 protein hypothetical protein hypothetical protein hypothetical protein | IID TTGKML _p YEESEEK VMTDP _p YVGR SQGpYEDELNAR NKELISEN _p YNLDLEIKNK GpYRNVINEALR pYFIEVEKQARK IPVDVIDMAA _p YGTMNGEK |

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Table 1. (Continued.)

kinase activity (Mijakovic *et al.* 2012, unpublished data). One can also not exclude that, when expressed, BY-kinase might be constitutively active, especially in Proteobacteria where the kinase domain is always linked to the extracellular domain.

Another intriguing question about BY-kinases concerns their ability to phosphorylate several substrates with distinct functions. Upon examination of BYmediated protein phosphorylation sites, no clear-cut recognition motifs emerged in the immediate surroundings of the phosphorylated tyrosines. Moreover, if one considers the number of tyrosine-phosphorylation sites detected in recent phosphoproteomic studies (table 1), the conservation of phosphotyrosine sites is nearly non-existent [84,91-99]. The existence of a specific phosphorylation motif is therefore unlikely in proteins phosphorylated by BY-kinases, and it seems likely that they recognize structural motifs distant from the phosphorylated tyrosine. At present, no structure of BY-kinase/substrate complex has been published, and such structures are eagerly anticipated.

Recent bacterial phosphoproteomics studies (table 1) demonstrate that tyrosine phosphorylation has similar abundance in Bacteria (3-10% of phosphoproteome) and Eukarya (3-39% of phosphoproteome) [100]. One has to consider the fact that bacterial phosphoproteomes are incomplete as they represent only snapshots of tyrosine phosphorylation in the bacterial cell in a given environment and growth state. To illustrate this, a recent comprehensive analysis dedicated to E. coli ribosomal proteins surprisingly detected 14 tyrosine phosphorylation sites, whereas none were reported in the first phosphoproteomic analysis of *E. coli* published a year before (table 1) [95]. It will therefore come as no surprise if more phosphotyrosine-proteins are detected in the near future. Quantitative phosphoproteomic of strains with inactivated individual kinase genes, grown in specific conditions are now needed to link tyrosine kinases to their physiological substrates and unravel the cellular function of bacterial tyrosine phosphorylation [101,102].

10. CONCLUSION

Challenging as it may be, there is no doubt that the ongoing effort made in the field of bacterial tyrosine phosphorylation will lead to the characterization of new regulatory processes crucial for the bacterial cell. Moreover, evidence is provided at an increasing rate that connections and cross talk exist between BY-kinases and other phosphorylating systems (i.e. two-component systems and eukaryotic-like serine/threonine kinases) found in the bacterial cell. The field of bacterial posttranslational modifications is rapidly expanding with the characterization of protein glycosylation, acetylation, methylation and pupylation [103]. It is evident that extensive cross talk exists among these various post-translational modifications in Eukarya [104], and we will undoubtedly have to tackle this kind of cross talk in bacterial protein phosphorylation networks, including but not limited to tyrosine phosphorylation.

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08-BLAN-0143 (C.G.) and 10-BLAN-1303-01 (I.M., C.G. and S.N.).

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