Myxococcus xanthus, a Gram-negative bacterium, contains a transmembrane protein serine/threonine kinase that blocks the secretion of β-lactamase by phosphorylation

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A gene, pkn2, encoding a Myxococcus xanthus protein with significant similarities to eukaryotic protein serine/threonine kinases, was cloned using the polymerase chain reaction. The open reading frame for the protein, beginning with a GUG initiation codon, consists of 830 amino acids. The amino-terminal 279 residues show 37% identity to catalytic domain of Pkn1, another protein serine/threonine kinase expressed during the development at the onset of sporulation. The catalytic domain of Pkn2 contains 27% and 25% identity to rat Ca2+/calmodulin-dependent protein kinase and Bos taurus rhodopsin kinase, respectively. In the middle of the carboxy-terminal regulatory domain, there is a typical transmembrane domain consisting of 18 hydrophobic residues. The gene product, Pkn2, produced in Escherichia coli under a T7 promoter was phosphorylated at both serine and threonine residues. TEM-1 β-lactamase produced in E. coli was found to serve as an effective substrate for Pkn2, phosphorylated only at threonine residues, shifting its apparent molecular mass from 29 to 44 kD. The phosphorylated β-lactamase was unable to be secreted into the periplasmic space and localized in the cytoplasmic and membrane fractions. Analysis of phoA fusions with pkn2 demonstrated that Pkn2 is a transmembrane protein with the kinase domain in the cytoplasm and the 207-residue carboxy-terminal domain outside the cytoplasmic membrane. Disruption of pkn2 showed no effect on vegetative growth but reduced the yield of myxospores by 30%–50%. On the basis of the present results, we propose that Pkn2 is a transmembrane protein serine/threonine kinase that regulates the activity of endogenous β-lactamase or related enzymes in response to an external signal yet to be identified.

[Key Words: Protein serine/threonine kinase; myxobacteria; transmembrane protein kinase; β-lactamase]

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Protein kinase cascades play major roles in intracellular signal transduction regulating various cellular functions in both prokaryotes and eukaryotes. In prokaryotes, protein histidine kinases are known to be the key enzymes for the cascades, and it has been believed for a long time that protein serine/threonine kinases do not exist in prokaryotes. These kinases, together with protein tyrosine kinases, are essential in signal transduction in eukaryotes. However, recently, the existence of such eukaryotic-like protein serine/threonine kinases in Myxococcus xanthus, a Gram-negative bacterium, has been demonstrated [Munoz-Dorado et al. 1991; Zhang et al. 1992]. M. xanthus living in the soil shows rather spectacular morphogenesis, including multicellular fruiting body formation upon nutrient starvation (for review, see Shimkets 1990). Within the fruiting body, a significant population of rod-shaped vegetative cells differentiate into spherical heat-resistant spores. It is believed that M. xanthus responds to a large number of signals in its natural habitat to regulate its cell–cell communication, gliding motility, aggregation, fruiting body formation, and differentiation to myxospores. The first prokaryotic protein serine/threonine kinase, Pkn1, has been characterized extensively and shown to be required for normal development of M. xanthus [Munoz-Dorado et al. 1991].

In this study we describe another protein serine/threonine kinase from M. xanthus, designated Pkn2. In contrast to Pkn1, Pkn2 was found to be a transmembrane protein kinase, which was able to effectively phosphorylate threonine residues of TEM-β-lactamase to block its secretion across the membrane. We propose that Pkn2 senses an external signal yet to be identified to regulate the activity of β-lactamase and/or related enzymes by phosphorylation. This is the first demonstration that a

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prokaryotic organism contains a transmembrane protein serine/threonine kinase. The significance of the role of Pkn2 in *M. xanthus* is discussed.

**Results**

**Identification and cloning of the pkn2 gene**

From the polymerase chain reaction (PCR) performed on the *M. xanthus* chromosomal DNA, using degenerate oligonucleotides designed according to the consensus sequences of catalytic subdomains VI and VIII of eukaryotic protein kinases as primers, two different DNA sequences were obtained, PCRPK1 and PCRPK2 (Munoz-Dorado et al. 1991). The sequence of PCRPK2 is shown in Figure 1A. PCRPK2 has a length of 173 bp and contains the consensus sequence DFG characteristic of subdomain VII in the same reading frame as subdomains VI and VIII of eukaryotic protein kinases (Hanks et al. 1988). This indicates that *M. xanthus* contains another eukaryotic-like protein kinase in addition to Pkn1 (Munoz-Dorado et al. 1991). PCRPK2 seems to be quite different from PCRPK1 (Munoz-Dorado et al. 1991). It has one extra codon over PCRPK1, and more significantly, the identity of the deduced amino acid sequences between the two PCR products is only 26%, excluding the sequences of the primers used in the reaction.

Using PCRPK2 as a probe, we attempted to isolate the entire gene corresponding to the PCRPK2 sequence. Southern blot hybridization analysis of *M. xanthus* chromosomal DNA digestions revealed that only one DNA fragment for each restriction enzyme digestion hybridized with the PCRPK2 probe (Fig. 1B). Positive clones were obtained from an *M. xanthus* genomic DNA library constructed in phage λ with use of the same probe. Of 1632 phages screened, 2 were detected as positive and more significantly, the identity of the deduced amino acid sequences between the two PCR products is only 26%, excluding the sequences of the primers used in the reaction.

When the deduced amino acid sequence of Pkn2 was compared with those of other eukaryotic protein kinases, it was found to contain the consensus catalytic domain of eukaryotic protein serine/threonine kinases from *M. xanthus*. The entire sequence of the 2.5-kb *XhoI* fragment of pJMPK2X was determined by using the strategy described in Materials and methods. The DNA sequence obtained, starting from the *XhoI* site (Fig. 2) is shown in Figure 3. The relationship between the two fragments and the position of the pkn2 gene are shown in Figure 2. The ORF was in the same reading frame as that of PCRPK2 and contained the consensus sequences of the 11 subdomains (Fig. 3) defined for the catalytic domain of eukaryotic protein serine/threonine kinases (Hanks et al. 1988). There are no ATG codons upstream of the consensus sequence of subdomain I starting from base 421. Instead, there are two GTG codons (positions 341–343 and 392–394). However, the codons used between the two GTG codons show a low G + C usage at the third position (10 of 16; 62.5%) in contrast to a high G + C usage of the codons used after the second GTG (92.5%), a typical codon usage in *M. xanthus* (Inouye et al. 1989). Thus, the ORF for pkn2 is tentatively assigned from the second GTG (base 392–394) to the TGA termination codon (base 2882–2884). A putative ribosome-binding site, AAGATG, can be identified 8 bases upstream of the initiation codon. The ORF should encode a protein (Pkn2) of 830 amino acid residues, with a calculated molecular weight of 87,594.

When the deduced amino acid sequence of Pkn2 was examined for sequence similarities with other proteins, it was found to contain the consensus catalytic domain of eukaryotic protein serine/threonine kinases from

**Figure 1. Identification of the pkn2 gene.** (A) The sequence of PCRPK2. The sequences of the primers used for PCR are marked by long arrows, and the highly conserved sequence in the subdomain VII of eukaryotic protein kinases is underlined. (B) Southern blot hybridization of the *M. xanthus* chromosomal DNA with the PCRPK2 probe. The digestions used were *BamHI* (lane 1), *EcoRI* (lane 2), *BamHI* and *HindIII* (lane 3), *BamHI* and *PstI* (lane 4), *SalI* (lane 5), *SalI* and *XhoI* (lane 6), and *XhoI* (lane 7). The molecular mass standards are expressed in kilobases.

**Figure 2.** Transmembrane Ser/Thr kinase from *M. xanthus*.

cloned from the same phage λ DNA. A 2.5-kb *XhoI* fragment was cloned into the pUX vector (Apelian and Inouye 1990) at the *XhoI* site, resulting in plasmid pJMPK2X. Using both pJMPK2S and pJMPK2X, a new construct denoted pJMPK2HB was constructed, which contains the 7.3-kb *XhoI*–*SalI* fragment encompassing the entire ORF. The physical map of the region containing the pkn2 gene is shown in Figure 2.

**pkn2 encodes a putative protein serine/threonine kinase**

The entire sequence of the 2.5-kb *XhoI* fragment of pJMPK2X and most of the sequence of the 4.8-kb *SalI* fragment of pJMPK2S was determined by using the strategy described in Material and methods. The DNA sequence obtained, starting from the *XhoI* site (Fig. 2) is shown in Figure 3. The relationship between the two fragments and the position of the pkn2 gene are shown in Figure 2. The ORF was in the same reading frame as that of PCRPK2 and contained the consensus sequences of the 11 subdomains (Fig. 3) defined for the catalytic domain of eukaryotic protein serine/threonine kinases (Hanks et al. 1988). There are no ATG codons upstream of the consensus sequence of subdomain I starting from base 421. Instead, there are two GTG codons (positions 341–343 and 392–394). However, the codons used between the two GTG codons show a low G + C usage at the third position (10 of 16; 62.5%) in contrast to a high G + C usage of the codons used after the second GTG (92.5%), a typical codon usage in *M. xanthus* (Inouye et al. 1989). Thus, the ORF for pkn2 is tentatively assigned from the second GTG (base 392–394) to the TGA termination codon (base 2882–2884). A putative ribosome-binding site, AAGATG, can be identified 8 bases upstream of the initiation codon. The ORF should encode a protein (Pkn2) of 830 amino acid residues, with a calculated molecular weight of 87,594.

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amino acid residue 20–274 (see Fig. 3) and the highest identity to the catalytic domain of Pkn1 (37%), a protein serine/threonine kinase of *M. xanthus* [Munoz-Dorado et al. 1991]. It also shows 27% identity with rat Ca<sup>2+</sup>/calmodulin-dependent protein kinase (Lin et al. 1987) and 25% identity with *Bos taurus* rhodopsin kinase [Lorenz et al. 1991]. After the catalytic domain, there is an unusual sequence of 99 amino acid residues (from residue 275 to 373) containing 20 Ser, 10 Thr, 23 Ala, 17 Gly, 10 Pro, and 0 charged residues (double-underlined in Fig. 3). These amino acid residues account for 81% of the sequence. This Ala/Ser/Thr/Gly/Pro-rich region is followed by a sequence of 467 amino acid residues (from residue 374 to 830), in the middle of which there is a hydrophobic stretch of 18 amino acid residues (from residue 606 to 623; boxed in Fig. 3). This hydrophobic stretch was added to the fusion protein, the phoA activity of the resulting protein, Pkn2–PhoA-2 increased dramatically to 5067 units, which was similar to that of a *phoA<sup>+</sup>* control strain (4014 units; see Fig. 4). Because PhoA is known to be active only when it is translocated across the membrane [Manoil and Beckwith 1985], the present results demonstrate that the hydrophobic domain corresponding to residues 606 to 623 functions as an internal signal sequence to translocate the PhoA domain fused downstream of the hydrophobic domain. Because the internal signal sequence appears to have no cleavage site at the carboxy-terminal end, the hydrophobic domain is likely to serve as a transmembrane domain. Therefore, the present results indicate that Pkn2 is a transmembrane protein consisting of the 207-residue carboxy-terminal domain in the periplasmic space and the 605-residue amino-terminal domain in the cytoplasm, where the kinase domain exists.

Expression of pkn2 in *E. coli*

To characterize Pkn2, the *pkn2* gene was expressed in *E. coli* using a T7 RNA polymerase system. The *pkn2* gene was cloned under a T7 promoter in plasmid pET11a as described in Material and methods. In this construction, the GTG initiation codon for *pkn2* was substituted with an ATG codon. The plasmid thus obtained was designated pET11/*pkn2*. After transformation of *E. coli* BL21(DE3) with this plasmid, the expression of the *pkn2* gene was induced with IPTG, and the proteins were labeled with Tran<sup>35</sup>S-label in the presence of rifampicin. As shown in Figure 5A, four new bands were induced in the presence of IPTG. As described later, band a was identified as Pkn2. Its apparent molecular mass (110 kD) was higher than its expected molecular mass (88 kD). This abnormal migration in a SDS–polyacrylamide gel is considered to be the result of phosphorylation of the product as observed for Pkn1 [Munoz-Dorado et al. 1991]. The apparent molecular masses of bands b, c, and d were 44, 31, and 29 kD, respectively, and all of them were found to be derived from the *bla* gene in the pET11a vector used as described later.

Next, we investigated phosphorylation of Pkn2 using the method applied for Pkn1 [Munoz-Dorado et al. 1991]. *E. coli* cells harboring pET11/*pkn2* were labeled with inorganic <sup>32</sup>P after induction with IPTG as described in
bodies were isolated and analyzed by SDS-PAGE. As bands a and b were labeled with $^{32}$P (lane 2). To identify protein was mainly phosphorylated at threonine residues phosphorylated in these bands, they shown in Figure 5B, of four bands (a, b, c, and d) only derived from the pkn2 gene, we examined the precursor-

Characterization of bands b, c, and d

In addition to band a, the induction of bands b, c, and d was puzzling. To determine whether these bands were derived from the pkn2 gene, we examined the precursor-

Materials and methods. After a 3-hr induction, inclusion bodies were isolated and analyzed by SDS-PAGE. As shown in Figure 5B, of four bands [a, b, c, and d] only bands a and b were pulse-labeled with $^{32}$P (lane 2). To identify amino acid residues phosphorylated in these bands, they were extracted from the gel and hydrolyzed in 6 N HCl. The hydrolysates were separated by two-dimensional thin layer chromatography. The results are shown in Figure 5, C and D. Pkn2 (band a) was phosphorylated at both serine and threonine residues (Fig. 5C), whereas the band b protein was primarily phosphorylated at threonine residues (Fig. 5D).

Figure 3. DNA sequence of 3753 bases encompassing pkn2 [GenBank/EMBL accession no. M94857] and the deduced amino acid sequence of Pkn2. Roman numerals represent conserved subdomains in the catalytic domain of kinases. The Ala/ Ser/Thr/Gly/Pro-rich region is doubly underlined. The sequence homologous to mitochondrial energy transfer protein is underlined. The hydrophobic stretch is boxed.

product relationship between these bands. For this purpose, the cells harboring pET11/pkn2 were pulse-labeled for 5 min with Tran$^{35}$S-label after induction with IPTG for 1 hr and chased for 7.5, 15, 30, and 60 min. As shown in Figure 6, Pkn2 (indicated with arrow a) was detected with a molecular mass of 106 kD during the 5-min pulse (lane 1). When chased, the position of Pkn2 gradually shifted to higher molecular mass, which was probably caused by continuous phosphorylation of Pkn2. In addition to band a, bands b, c, and d of 44, 31, and 29 kD, respectively were pulse-labeled with Tran$^{35}$S-label. The intensity of band c decreased with a concomitant increase of the intensity of band b during the chase period. These results suggest that band c at 31 kD was chased into band b at 44 kD, probably because of the post-translational modification of band c. Band b is unlikely to be derived from Pkn2 (band a), because the intensity of band a remained constant throughout the chase period tested (Fig. 6).
Deletion analysis of pkn2 in E. coli

To confirm further that band a is the pkn2 product, various deletion mutations of the pkn2 gene were constructed, as described in Materials and methods, using the unique restriction enzyme sites in the gene [SacII (1221–1226), KpnI (1415–1420), MluI (1548–1553), NcoI (1899–1904), and XhoI (2462–2467; see Fig. 3)] as shown in Figure 7A. The deletion constructs were expressed under the control of the T7 promoter in pET11a. The mutation designated ΔKn was constructed by introducing an NdeI site at Met-398 (see Fig. 3) followed by cloning the resulting 3.5-kb Ndel–BamHI fragment into pET11a at its Ndel and BamHI sites. Therefore, the resulting plasmid, designated pΔKn, encodes a carboxy-terminal fragment of Pkn2 consisting of 433 amino acid residues, which lacks the entire kinase domain.

The cells harboring pET11/pkn2 and the deletion constructs were labeled with Tran35S-label for 30 min with or without IPTG induction in the presence of rifampicin. The products were then analyzed by SDS-PAGE as shown in Figure 7B. In the case of pET11/pkn2, Pkn2

| Plasmid | PhoA fusion protein | Phosphatase activity (Unit) |
|---------|---------------------|----------------------------|
| None    |                     | 16                         |
| pCH2    | Bla Signal Peptide  | 4014                       |
| pPkn2-PhoA-1 |                 | 58                         |
| pPkn2-PhoA-2 |                 | 5067                       |

Figure 5. Expression of Pkn2 and identification of phosphorylated amino acid residues. (A) Expression of pkn2 under the T7 promoter in E. coli BL21 (DE3). Cells harboring pET/pkn2 were incubated in the presence (lane 1) or the absence (lane 2) of 1 mM IPTG, and then labeled with Tran35S-label in the presence of rifampicin [150 μg/ml] as described in Materials and methods. Total cellular proteins were subjected to SDS-PAGE. Proteins were blotted on an immobilon-P membrane, which was then exposed to X-ray film. Letters with an arrow indicate the positions of Pkn2 (band a) and other products from pET11/pkn2 (bands b, c, and d) [see text]. Bars with numbers at left indicate the positions of molecular mass markers (kD). (B) Phosphorylation of the products from pET11/pkn2. Cells harboring pET11/pkn2 were labeled with either Tran35S-label (lane 1) or ortho-32P [lane 2] after 2-hr induction with IPTG in the presence of rifampicin as described in Materials and methods. [lane 1] SDS-gel analysis of total cellular proteins labeled with Tran35S-label; [lane 2] SDS-gel analysis of inclusion bodies labeled with ortho-32P. Inclusion bodies were isolated by a low centrifugation after sonification as described in Materials and methods. SDS-PAGE was carried out as in A. [C,D] Phosphoamino acid analysis of Pkn2 and band b in lane 2 (in B) [described in Materials and methods]. The positions of the nonradioactive phosphoamino acid standards [|P-Ser| phosphoserine, |P-Thr| phosphothreonine, |P-Tyr| phosphotyrosine] identified by ninhydrin and the position of inorganic phosphate [Pi] are shown by broken lines. The first and second dimensions used for the separation of the phosphoamino acids are indicated by numbers 1 and 2, respectively. The origin is indicated by a dot at the intersection of arrows 1 and 2.
came almost undetectable (Fig. 7B, lanes 8, 10, and 12). In the case of ΔM-B, a faint band appeared slightly below the band b position, suggesting that this faint band may have something to do with band b. Note that the band at 41 kDa in ΔK-B (lane 10) is the truncated product of Pkn2 but not the band b product as discussed later. Interestingly, band c became a major product, and no band b was produced in ΔKn (lane 14).

It should be noted that band b was still produced in ΔN-B (lane 6) and that the plasmid for ΔN-B contained the 1.5-kb fragment from the M. xanthus chromosomal DNA that encompasses only the amino-terminal fragment of Pkn2 from residue 1 to 502. Therefore, band b was not likely to be derived from the M. xanthus DNA fragment. We thus speculated that the bla gene for ampicillin resistance encoding TEM-β-lactamase in the pET11a vector is responsible for the production of band b.

As shown in Figure 8A, the bla gene locates downstream of the pkn2 gene in the same orientation in a circular map of pET11/pkn2. To determine that band b is really a product of the bla gene, the bla gene in pET11/pkn2 was replaced with the kanR gene for kanamycin resistance from the Tn5 transposon as described in Materials and methods, and the resultant plasmid was designated pET11km/pkn2. The cells harboring pET11/pkn2 and pET11km/pkn2 were labeled with Tran35S-label for 30 min in the presence of rifampicin with or without the addition of IPTG. The labeled products were then analyzed by SDS-PAGE. As shown in Figure 8B, band a was still clearly detected in the presence of IPTG in cells harboring pET11km/pkn2 (lane 4) as well as in cells harboring pET11/pkn2 (lane 2). However, band b, together with bands c and d, completely disappeared in the cells harboring pET11km/pkn2 in which the β-lactamase gene was replaced with the kanR resistant gene (compare lane 4 with lane 2).

To confirm further that band b was derived from the bla gene, the same samples used in Figure 7B were analyzed by Western blot using anti-β-lactamase antibody. As shown in Figure 7C, strong bands corresponding to band b were detected not only in the cells harboring pET11/pkn2 (lane 2) but also in the cells harboring pΔX-B (lane 4), and pΔN-B (lane 6). A faint band slightly below the band b position was also detected in cells harboring pΔM-B (lane 8), which also coincides well with the faintly labeled band at the same position in Figure 7B, lane 8.

In contrast to band b, band d was detected in all lanes in Figure 7C, indicating that it is β-lactamase, the product of the bla gene. The apparent molecular mass of band d (29 kDa) also agrees well with the molecular mass of β-lactamase. It is not known at present why band d appears as a doublet form. The intensities of band d were also very similar in all lanes regardless of the absence or the presence of IPTG, indicating that β-lactamase at band d was synthesized from its own promoter. On the other hand, the production of bands b and c was observed only in the presence of IPTG and, thus, likely to be under the control of the T7 promoter of pET11a. On the basis of

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**Band b is the product of the bla gene**

In the deletion analyses of pkn2 shown in Figure 7B, one can observe that in contrast to band a, band b did not disappear in ΔX-B and ΔN-B mutations (lanes 4 and 6, respectively). When the 3′-end region of pkn2 was deleted beyond the Ncol site, the production of band b be-
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Figure 7. Analysis of the band b product using various pkn2 deletion mutants. (A) Schematic diagram of pkn2 and its deletion mutants. The method for constructing the deletion mutations is described in the text. (B) Expression of pkn2 and its deletion mutants. E. coli BL21 [DE3] harboring a plasmid containing individual pkn2 mutants was labeled with Tran35S-label for 2 hr in the presence of rifampicin. (- and +) Without and with IPTG, respectively. (Lanes 1,2) Total cell extract from cells harboring pET11/pkn2; (lanes 3,4) from pAX-B; (lanes 5,6) from pAN-B; (lanes 7,8) from pAM-B; (lanes 9,10) from pAK-B; (lanes 11,12) from pAS-B; (lanes 13,14) from pAKn. The products corresponding to individual deletion mutations are marked with dots. (C) Western blot analysis using anti-β-lactamase antiserum. The same sets of samples as used in B were transferred on PVDF membrane to detect cross-reactive materials. Lanes 1 to 14 were the same as shown in B. Letters with an arrow and bars with a number are the same as in Fig. 6.

the apparent molecular mass of band c (31 kD), this band is considered to be the secretory precursor of β-lactamase, which still contains the uncleaved signal peptide of 23 amino acid residues.

The present results indicate that TEM-β-lactamase serves as a substrate for Pkn2, which can be phosphorylated very effectively at multiple threonine residues (Fig. 5D), resulting in the shift of the apparent molecular mass from 31 [band c] to 44 kD [band b]. It is interesting to note that the Ala/Ser/Thr/Gly/Pro-rich domain of Pkn2 (see Fig. 7A) is required for kinase activity, because AK-B and AS-B were unable to phosphorylate β-lactamase (Fig. 7C, lanes 10 and 12, respectively).

Phosphorylation of β-lactamase and the Pkn2 regulatory domain

Next, we examined whether Pkn2 phosphorylates β-lactamase produced by a separate plasmid. For this purpose, E. coli cells were transformed with both pET11km/pkn2 and pAKn. The cells transformed with either plasmid or both were labeled with Tran35S-label in the presence of rifampicin. As shown in Figure 9A, Pkn2 became the major product in the cells harboring pET11km/pkn2 when IPTG was added [band a in lane 2). In the cells harboring pAKn, three major bands appeared in the presence of IPTG [bands c, d, and f in lane 4]—band d for β-lactamase, band c for pro-β-lactamase, and band f for the truncated product of Pkn2. The same pattern was observed in Figure 7B, lane 14.

When cells harboring both plasmids were analyzed, two extra broad bands [bands b and e in lane 6] were detected in addition to bands a, c, d, and f. When the same sample used in Figure 9A was analyzed by Western blot using anti-β-lactamase antibody, not only bands c and d, but also band b, were detected as shown in Figure 9B, lane 6, indicating that Pkn2 produced from a separate plasmid could phosphorylate β-lactamase precursor.
Transmembrane Ser/Thr kinase from M. xanthus

Most importantly, Pkn2 was also able to phosphorylate the carboxy-terminal fragment of Pkn2 resulting in the formation of band e when both ΔKn and pkn2 were co-expressed.

Association of Pkn2 with phosphorylated β-lactamase

Because eukaryotic protein kinases are known to be associated with their substrates, we then examined whether Pkn2 forms a complex with β-lactamase. BL21(DE3) cells harboring pET11/pkn2 were labeled with Tran35S-label with or without rifampicin, as described in Materials and methods. Cell lysates were immunoprecipitated with anti-Pkn2 or anti-β-lactamase serum. As shown in Figure 10, lanes 4 and 10, phosphorylated β-lactamase [band b] was coprecipitated with Pkn2 [band a] when anti-Pkn2 serum was used. Similarly, when the cell lysate was treated with anti-β-lactamase serum, Pkn2 [band a] was coprecipitated [lanes 6,12]. Note that bands c and d in lanes 6 and 12 are pro-β-lactamase and β-lactamase, respectively. It should be noted that bands c and d could not be detected with anti-Pkn2 serum [lanes 4,10]. These results demonstrate that phosphorylated β-lactamase forms a complex with Pkn2.

Cellular localization of phosphorylated β-lactamase

To examine the cellular localization of phosphorylated β-lactamase, cells harboring pET11/pkn2 were incubated with IPTG for 60 min, and cellular fractionation was carried out by the method of Neu and Heppel (1965). As shown in Figure 11, phosphorylated β-lactamase, together with the precursor form of β-lactamase [band c], was detected in both the membrane and the cytoplasmic fractions [lanes 4 and 5, respectively], but not in the periplasmic fraction [lane 3]. In contrast, the mature form of β-lactamase was found primarily in the periplasmic fraction [lane 3]. The periplasmic localization of mature β-lactamase is particularly evident in the control experiment with p2AKn [lane 8]. Note that no band b was produced in the control experiment [lanes 6–10]. These results indicate that the phosphorylation of β-lactamase blocks its secretion across the membrane resulting in the accumulation of phosphorylated β-lactamase in the cytoplasm as well as in the membrane fraction.

Expression of pkn2 in M. xanthus

M. xanthus strain pkn2/Z was constructed, which contained intact pkn2 as well as pkn2-1acZ fusion. The pkn2/Z strain was plated on CYE agar plate for vegetative growth as well as on CF agar plate for fruiting body formation. X-gal [5-bromo-4-chloro-3-indolyl-β-D-galac-
The assignments are the same as in Fig. 6. Periplasmic, membrane, and cytoplasmic fractions were applied to the gel. The results indicate that Pkn2 was expressed only during the vegetative growth but not during development.

A more quantitative analysis of the expression of pkn2 was obtained by assaying β-galactosidase activity in the pkn2ΔKn strain during both vegetative growth and the developmental cycle. The specific β-galactosidase activities were between 3 and 5 U/mg of protein/min during the exponential growth and sharply dropped to <1 during the stationary phase. β-Galactosidase activity on CF agar plates decreases rapidly and stays at a very low level of <2 U/mg of protein/min. These results are consistent with the plating experiment described above, indicating that pkn2 is mainly expressed during the late exponential growth phase.

**Effects of a disruption mutation of pkn2**

As described in Materials and methods, the pkn2 gene on the chromosome was disrupted by inserting the kan' gene from Tn5 into the unique Sall site that lies within the coding region for subdomain IV of the Pkn2 catalytic domain forming strain Δpkn2. Strain Δpkn2 was found to grow normally as the wild-type strain in CYE medium, indicating that pkn2 is not essential for vegetative growth. Strain Δpkn2 was also able to form well-defined fruiting bodies on CF agar plates. The fruiting body morphology of strain Δpkn2 was somewhat different from that of the wild-type strain, strain Δpkn2 produced continuous mounds that appeared loosely packed, in contrast to the well-separated round mounds of the wild-type strain (not shown).

We also quantitated the number of myxospores produced during development. The Δpkn2 spore production was ~30%–50% of the level of the wild-type strain. The Δpkn2 myxospores were viable at the same level as the wild-type spores (not shown).

**Discussion**

In this study we demonstrated that *M. xanthus* contains a transmembrane protein serine/threonine kinase, designated Pkn2. We have described previously that *M. xanthus* contains a cytoplasmic protein serine/threonine kinase, Pkn1 (Munoz-Dorado et al. 1991). Pkn2, consisting of 830 amino acid residues, contains the 11 subdomains characteristic of the eukaryotic protein kinases at the amino-terminal region. Immediately after the last conserved subdomain XI, there is a 99-residue region (from residue 275 to 373) enriched with serine (20 residues), threonine (10), alanine (23), glycine (17), and proline (10). These residues account for 81% of this region. This domain may be the site for phosphorylation, having a total 35 serine plus threonine residues in this region, and the phosphorylation of this domain may be involved in the regulation of Pkn2 function.

From the PhoA fusion analysis, the 207-residue carboxy-terminal domain immediately after the 18-residue hydrophobic domain was found to be translocated across the membrane, which is likely to serve as a receptor to sense an external signal yet to be identified. Most significantly, it was found that Pkn2 is a protein kinase that is able to phosphorylate threonine residues of TEM-β-lactamase specifically. Our data indicate that only nascent β-lactamase was phosphorylated, which prevented the translocation of β-lactamase across the membrane. TEM-β-lactamase contains a total of 20 threonine residues, of which 5 residues are found in known phosphorylation site motifs (R/KXXT and R/KKT), Hunter 1991). Consistent with this fact, five radioactive tryptic peptides were obtained when phosphorylated β-lactamase was digested with trypsin. These results indicate that Pkn2 phosphorylates β-lactamase at multiple sites to block its secretion. At present, it is not known whether phosphorylated β-lactamase still contains the signal peptide.

*M. xanthus* is known to be highly resistant to ampicillin, and a β-lactamase, which is cross-reactive with anti-TEM-β-lactamase serum has been identified in this
organism (K. O'Connor and D. Zusman, pers. comm.). It is tempting to speculate that Pkn2 may regulate the activity of β-lactamase and/or other enzymes related to β-lactamase such as penicillin-binding proteins by phosphorylating them under certain conditions. In this fashion, cell viability, cell shapes, or morphology may be regulated under certain circumstances.

In eukaryotes, most protein serine/threonine kinases are soluble localizing in the cytoplasm. However, a few protein serine/threonine kinases, such as transforming growth factor (TGF)-β receptor (Massagué et al. 1994) and IRE1 (Cox et al. 1993), are known to be transmembrane protein kinases. Pkn2 is the first transmembrane protein serine/threonine kinase ever found in prokaryotes. In prokaryotes, a large number of transmembrane protein histidine kinases are known, which function as sensors for various external signals (Stock et al. 1989). It remains to be elucidated how M. xanthus differentially utilizes two types of transmembrane kinases in terms of their functions. It is also an intriguing question how M. xanthus acquired the serine/threonine kinase gene during the course of evolution.

Materials and methods

Materials

[α-32P]dCTP, [γ-32P]ATP, [α-35S]dATP, and 32P were purchased from Amersham, and Tran3SS-label containing [3SS]methionine and [3S]selenomethionine from ICN. DNA sequencing was performed with a Sequenase kit obtained from U.S. Biochemical. Taq polymerase for PCR was purchased from Perkin-Elmer Cetus, restriction enzymes from New England Biolabs, and T4 DNA ligase from Bethesda Research Laboratories.

Bacterial strains and growth conditions

M. xanthus DZF1 was grown at 30°C in CYE medium (Campos and Geisselsoder 1978), and 40 μg/ml of kanamycin was added when necessary.

M. xanthus fruiting bodies were obtained on CF agar plates (Hagen et al. 1978), and the samples were harvested and treated as reported previously [Munoz-Dorado et al. 1991].

E. coli JM83 (Vieira and Messing 1982) and E. coli CL83 (Lerner and Inouye 1990) were used as recipient strains for transformation and P1 infection. Cells were grown in LB medium (Miller 1972) supplemented with 50 μg/ml of ampicillin, 25 μg/ml of kanamycin, and/or 10 μg/ml of chloramphenicol when necessary. E. coli BL21(DE3) was used for the T7 RNA polymerase expression system (Studier et al. 1990). This strain was grown in M9 medium containing 19 amino acids, excluding methionine, when cells were labeled with [35S]methionine and/or [3S]selenomethionine.

E. coli BL21(DE3) was used for the T7 RNA polymerase expression system [Studier et al. 1990]. This strain was a derivative of pJMPK2HB and contained the lacZ gene without a promoter from E. coli (Masui et al. 1983). pUC7Kan5 and pUC9Kn[Pst I] were derivatives of pUC7 and pUC9, respectively, constructed by S. Inouye (unpubl.), they contained the kanamycin resistance gene from Tn5. Plage Pich100Cm (Rosner 1972) was used to transduce cloned DNA from E. coli to M. xanthus.

DNA manipulation and sequencing

M. xanthus chromosomal DNA was prepared according to the method of Avery and Kaiser (1983) and phage λ DNA according to the method of Maniatis et al. (1982). Southern blot analysis was performed by the method described by Southern (Southern 1975). Plage plaques were blotted to nitrocellulose filters by contact, and M. xanthus and E. coli colonies were grown on Whatman 3MM filters; the filters were treated as described previously [Inouye and Inouye 1987]. The DNA probes were labeled by nick translation, and hybridization was carried out at 42°C in 50% formamide as reported [Maniatis et al. 1982]. DNA sequencing was determined by the dideoxynucleotide chain termination method [Sanger et al. 1977] using double-stranded plasmid DNA and the universal primers and the synthetic oligonucleotides designed from newly obtained DNA sequences as primers.

Construction of pkn2 and its deletion mutants under a T7 promoter

To clone the pkn2 gene under the control of the T7 promoter in pET11α, an NdeI site was first created at the initiation codon of the pkn2 gene by PCR, using the following two primers: 5′-AAGATAAGCTTCATATGCTGGCCCCTGACTCCCTT-3′; which annealed at position 393–412 (Fig. 3) and contained NdeI and HindIII sites, and 5′-GCAGGCGTCGTCGGCGATT-3′; which annealed at position 633–652, immediately downstream of the TGF-β receptor (Massagué et al. 1994) and 40 μg/ml of kanamycin was added when necessary.

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Expression of pkn2 and its deletion mutants in E. coli

Using E. coli BL21(DE3) harboring pET11/pkn2 or its deletion constructs, the expression of pkn2 or its mutated genes was induced by addition of 1 mM IPTG for 1 hr. Rifampicin [150 μg/ml] was then added, and the culture was incubated further for 15 min. After treatment the cells were labeled with Tran35S-
label for a period specified in each experiment. Pulse–chase experiments were performed by labeling for 5 min without adding rifampicin and chasing in the presence of 200 μg/ml of nonradioactive methionine at various times.

Western blot analysis
After SDS-PAGE, the samples were transferred onto a PVDF membrane using a Sartoblot semidry transfer apparatus as recommended by the manufacturer (Sartorius, Gottingen, Germany). β-Lactamase was detected by anti-β-lactamase serum subjecting anti-rabbit IgG conjugated to alkaline phosphatase (Bio-Rad) with chromogenic substrates.

Immunoprecipitation of Pkn2 and its mutants, and β-lactamase
Ten-milliliter cultures of E. coli BL21(DE3) harboring various plasmids were labeled with Tran35S-label as described previously. The cells were harvested, washed with 50 mM Tris-HCl (pH 8.0), and then suspended in 50 μl sonication buffer [50 mM Tris-HCl (pH 8.0), 1 mM PMSF, 1 mM EDTA, and 2 mM β-mercaptoethanol]. After sonication, cell debris was removed by low centrifugation and 500 μl of TENN [50 mM Tris-HCl (pH 8.0), and then suspended in 50 μl sonication buffer 140 mM NaCl, 5 mM EDTA, 0.5% NP-40] was added. After the addition of 5 μl of anti-β-lactamase serum, the mixture was incubated on ice for 1 hr, and then 5 μl of Staphylococcus aureus ghosts purchased from Sigma was added. After 15 min at room temperature, precipitates were recovered by centrifugation (10,000 g for 1 min), washed three times with TENN, suspended in SDS-PAGE sample buffer [80 mM Tris-HCl (pH 6.8), 2% SDS, 0.2 mM EDTA, 10% glycerol], and analyzed on 15% SDS-PAGE after boiling. The gel was dried and autoradiographed.

Cellular fractionation
Isolation of periplasmic, membrane, and cytoplasmic fractions was carried out according to the method by Neu and Heppel (1965). Briefly, E. coli cells grown at Klett 100 in 10 ml of M9 medium with or without induction of IPTG were harvested by centrifugation at 5000 g for 10 min at 4°C, and the cell pellet was washed with 10 mM Tris-HCl (pH 8.0). The washed cells were resuspended into 1 ml of 20% sucrose in 30 mM Tris-HCl (pH 8.0) containing 1 mM EDTA, and the suspension was incubated for 10 min at room temperature. The cell suspension was then spun at 13,000 g for 10 min at 4°C, and after the addition of 1 ml of ice-cold water to the cell pellet, the cell suspension was incubated for 10 min at 4°C. The cell suspension was centrifuged at 5000 g for 10 min at 4°C, and the supernatant was used as the periplasmic fraction. The cell pellet was then disrupted by sonication in 0.5 ml of 50 mM Tris-HCl (pH 8.0), 1 mM EDTA, 1 mM phenylmethane sulfonyl fluoride (PMSF), and 2 mM β-mercaptoethanol. The cell debris and unbroken cells were removed by centrifugation at 5000 g for 10 min at 4°C, and the supernatant was fractionated into the membrane and cytoplasmic fractions by ultracentrifugation at 100,000 g for 30 min at 4°C.

Construction of pkn2–phoA fusion genes and PhoA assay
The DNA fragments, with or without the region encompassing a putative transmembrane domain, were amplified by PCR using pMPK2EB as a template and the following oligonucleotides as primers. P1 (5'-CTAACGCTTGCCGAACGCCGTCGCG-3') corresponds to the region from amino acid 576 to 580 in Figure 3. This primer contains a HindIII site at the 5' end that enables in-frame fusion of the pkn2 fragments with the 3' fragment of lacZ in pUC9. P2 (5'-CTCTGACCGCTGCGGCTCCGCTCG-3') and P3 (5'-CTCTGACCGGCGGCTCCGCTCG-3') correspond to the region from residue 601 to 605 and from residue 648 to 652 in Figure 3, respectively. They have a PstI site at the 3' end to enable the in-frame fusion of the pkn2 fragments with the phoA gene in pCH2 (Hoffman and Wright 1985). PCR-amplified fragments were digested with HindIII and PstI and cloned in the HindIII and PstI sites of pUC9. The resulting plasmids were digested with PstI, and the 3-kb PstI fragment containing the phoA gene from pCH2 was ligated into the PstI site, followed by determination of the correct orientation of the phoA gene. Using the resulting plasmids, pkn2–phoA-1 and -2, the expression of the phoA gene was under the control of the lac promoter–operator. Before PhoA assay, cells were incubated in M9 medium in the presence of 1 mM IPTG for 2 hr. PhoA activity was measured as described previously (Brickman and Beckwith 1975).

Phosphoamino acid analysis
E. coli BL21 (DE3) harboring pET11/pkn2 was labeled in vivo with ortho-32P as described previously (Munoz-Dorado et al. 1991). Phosphoamino acid analysis was carried out as described previously (Kamps and Selton 1989).

Construction of the pkn2/Z strain
To fuse the pkn2 gene to lacZ, a BamHI site was created in the pkn2 gene in-frame with the BamHI site of lacZ in pKM005 by PCR. The reaction was performed with the oligonucleotides 5'-GGCCGCGGATCCAGGGGCAGCGG-3', which anneals at position 1090–1113 (Fig. 3) and contains a BamHI site of the phoA gene. Using the resulting plasmids, pkn2–phoA-1 to -2, the expression of the phoA gene was under the control of the lac promoter–operator. Before PhoA assay, cells were incubated in M9 medium in the presence of 1 mM IPTG for 2 hr. PhoA activity was measured as described previously (Brickman and Beckwith 1975).

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