**Bacillus subtilis** 5′-nucleotidases with various functions and substrate specificities

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**Abstract**

**Background:** In *Escherichia coli*, *nagD*, *yrfG*, *yyjG*, *yieH*, *yigL*, *surE*, and *yfbR* encode 5′-nucleotidases that hydrolyze the phosphate group of 5′-nucleotides. In *Bacillus subtilis*, genes encoding 5′-nucleotidase have remained to be identified.

**Results:** We found that *B. subtilis* *ycsE*, *araL*, *yutF*, *ysA*, and *yqeG* show suggestive similarities to *nagD*. Here, we expressed them in *E. coli* to purify the respective His₆-tagged proteins. YcsE exhibited significant 5′-nucleotidase activity with a broader specificity, whereas the other four enzymes had rather weak but suggestive activities with various capacities and substrate specificities. In contrast, *B. subtilis* *yktC* shares high similarity with *E. coli* *suhB* encoding an inositol monophosphatase. YktC exhibited inositol monophosphatase activity as well as 5′-nucleotidase activity preferential for GMP and IMP. The *ycsE*, *yktC*, and *yqeG* genes are induced by oxidative stress and were dispensable, although *yqeG* was required to maintain normal growth on solid medium. In the presence of diamide, only mutants lacking *yktC* exhibited enhanced growth defects, whereas the other mutants without *ycsE* or *yqeG* did not.

**Conclusions:** Accordingly, in *B. subtilis*, at least YcsE and YktC acted as major 5′-nucleotidases and the four minor enzymes might function when the intracellular concentrations of substrates are sufficiently high. In addition, YktC is involved in resistance to oxidative stress caused by diamide, while YqeG is necessary for normal colony formation on solid medium.

**Keywords:** *Bacillus subtilis*, Haloacid dehalogenase superfamily, Inositol monophosphatase, Inositol phosphate, Nucleoside/nucleotide metabolism, 5′-nucleotidase, Oxidative stress, Phosphatase, Protein motif

**Background**

The pool sizes of nucleotides and nucleosides are balanced to enable the efficient synthesis of DNA and RNA [1]. Numerous enzymes involved in the biosynthesis and catabolism of nucleic acids are controlled to regulate the appropriate pool size of each compound. For example, 5′-nucleotidases hydrolyze 5′-nucleotides to generate nucleosides and inorganic phosphate, which participate in the regulatory mechanism that opposes the generation of nucleotides with the phosphorylation of nucleosides catalyzed by kinases.

5′-Nucleotidases are ubiquitous among species and reside in different subcellular locations. Extracellular 5′-nucleotidases are produced by certain bacteria. For example, *Vibrio parahaemolyticus* NutA is a 5′-nucleotidase bound to the membrane by a lipid anchor [2]. *Escherichia coli* UshA is a periplasmic enzyme with 5′-nucleotidase and UDP-sugar hydrolase activities [3]. These 5′-nucleotidases degrade extracellular nucleotides to satisfy the cell’s nutritional requirements.

In contrast, bacteria produce numerous intracellular 5′-nucleotidases that belong to various enzyme families. In *E. coli*, the substrates of NagD are UMP, GMP, AMP, and CMP [4]. NagD belongs to the haloacid dehalogenase superfamily (HADSF) characterized by a specific protein motif [5]. The HADSF family comprises numerous proteins in organisms ranging from prokaryotes to higher eukaryotes, including humans. The vast majority of enzymes...
of the HADSF family are phosphoryl transferases, although the superfamily was named after 2-haloacid dehalogenase, because it is the first structurally characterized member. When the amino acid sequences of their entire coding regions are compared, similarities among the HADSF members are not usually very high (15–30 % identical), although their central regions involved in catalytic activity are relatively conserved. Further, correlation between structure and catalytic activity is frequently observed and interrelates with similarities among the structures of substrates [6–8]. E. coli produces other HADSF-family enzymes such as YrfG, YjjG, YieH, and YigL, and each exhibits 5′-nucleotide phosphatase activity [9, 10].

In addition, E. coli produces SurE and YfbR, which exhibit 5′-nucleotidase activity and do not belong to the HADSF family [10]. SurE shows broad substrate specificity, by dephosphorylating 5′-nucleotides as well as 3′-nucleotides, with highest affinity for 3′-AMP. SurE hydrolyzes polyphosphate with preference for short-chain substrates. Homologs of E. coli surE are present in numerous eubacteria and archaea, and SurE represents a family of metal-dependent phosphatases [11]. YfbR belongs to the HD domain superfamily of metal-dependent phosphatases and phosphodiesterases, and the HD domain was named after a protein motif with predicted catalytic residues containing the conserved doublet His–Asp [12]. YfbR specifically hydrolyzes 5′-deoxyribonucleotides.

Inositol monophosphatase denotes phosphatases that liberate inorganic phosphate from myo-inositol 1-monophosphate (MIMP). The mammalian and plant enzymes are involved in the metabolism of inositol phospholipids, including the generation and degradation of inositol phosphates and phosphatidylinositols that mediate cellular signal transduction [13]. Inositol monophosphatases are present in diverse organisms such as bacteria and higher eukaryotes, and evolved from a common ancestral gene [14, 15]. Mammalian inositol monophosphatases exhibit relatively broad substrate specificity for phosphate-containing compounds [16, 17]. For example, the enzyme isolated from rat testis hydrolyzes adenosine 2′-monophosphate [16] and the bovine brain enzyme hydrolyzes β-glycerophosphate and adenosine 2′-monophosphate [17]. E. coli suhB encodes a protein homologous to eukaryotic inositol monophosphatases with equivalent activities [18].

However, phosphatidylinositol is not present in E. coli usually [19], and its physiological function in cells is unknown. In contrast, E. coli SuhB exhibits wider substrate specificity and hydrolyzes β-glycerophosphate and adenosine 2′-monophosphate as well [18]. Thus, we hypothesized that bacterial inositol monophosphatases might also function as 5′-nucleotidase. Moreover, various E. coli enzymes likely dephosphorylate 5′-nucleotides with different specificities and functions and may therefore act together to maintain the sizes of the intracellular pools of nucleotides and nucleosides.

In Bacillus subtilis, the partial nucleotide limitation induces 5′-nucleotidase activity [20], and B. subtilis is used for fermentative inosine production, which may involve the dephosphorylation of IMP catalyzed by an unidentified 5′-nucleotidase [21]. However, B. subtilis genes encoding 5′-nucleotidase have remained to be identified.

Here we selected B. subtilis genes potentially encoding 5′-nucleotidases [22] and expressed them in E. coli to characterize their function. We identified two major and four minor genes encoding 5′-nucleotidases, with various functions and substrate specificities. A major 5′-nucleotidase was involved in resistance to oxidative stress, and a minor enzyme was required for normal growth on solid medium.

Results
Selection of B. subtilis genes similar to E. coli genes encoding 5′-nucleotidase
In E. coli, at least nagD, yrfG, yjjG, yieH, yigL, surE, and yfbR encode 5′-nucleotidases, and their gene products are classified in the families as follows: HADSF (NagD, YrfG, YjjG, YieH, and YigL), SurE, and HD domain (YfbR). In the B. subtilis genome, no gene shares significant homology with surE or yfbR. However, gene products of araL, yutF, yqeG, ysaA, ycsE, gapB, ftsA, and hprP share some suggestive similarities with that of E. coli nagD (Fig. 1); AraL (65.8 % similarity/272 aa overlap),
YutF (73.0 %/256 aa), YqeG (15.7 %/172 aa), YsaA (16.9 %/260 aa), YcsE (51.4 %/249 aa), GapB (6.8 %/340 aa), FtsA (9.5 %/440 aa), and HprP (15.7 %/216 aa). Among them, AraL, YutF, YqeG, YsaA, YcsE, and HprP were supposed to belong to HADSF, since their amino acid sequences contain the HADSF motif (http://pfam.xfam.org/family/PF00702.24).

The araL gene resides within the L-arabinose operon, which encodes enzymes required for the degradation of L-arabinose and is under the control of AraR repressor to be induced in the presence of L-arabinose [23]. AraL was predicted to be a phosphatase that hydrolyzes certain sugar-phosphates [23].

The yutF gene encodes a protein that was predicted to be involved in N-acetyl-glucosamine catabolism and similar to 4-nitrophenyl phosphatase [24]. The condition-dependent transcriptome analysis revealed that yutF was almost constitutively expressed at low levels [25].

The yqeG, ysaA, and ycsE genes encode a putative HADSF-family enzyme of unknown function. The yqeG gene is the first gene of the long operon comprising yqeG, yqeH, aroD, yqeL, nadD, yqeK, and yqeM. This operon encodes essential genes [26], which are expressed constitutively at significant levels that increase during germination and under conditions of oxidative stress [25]. The ysaA and ycsE genes are constitutively expressed and are induced, respectively, during the nutritional shift from malate to malate plus glucose and in the presence of ethanol or diamide [25].

The products of gapB, ftsA, and hprP show relatively lower similarities to NagD (Fig. 1). GapB is NADP-dependent glyceraldehyde-3-phosphate dehydrogenase, which is involved in gluconeogenesis [27]. FtsA is an actin-like ATPase involved in cell division [28]. And HprP is a P-Ser-HPr phosphatase involved in carbon catabolite repression [29].

5′-Nucleotidase activity of B. subtilis enzymes sharing some similarities with NagD

Of the eight B. subtilis genes that share some similarities with nagD as described above, gapB, ftsA, and hprP have respective and specific functions other than as 5′-nucleotidases [27–29]. Therefore, these genes were excluded from the present study. We cloned and expressed araL, yutF, yqeG, ysaA, and ycsE in E. coli as His6-tag fusion proteins.

The His6-tag fusion proteins were purified to form respective single bands in sodium dodecyl sulfate polyacylamide gel electrophoresis (SDS-PAGE, data not shown), and were subjected to phosphatase assays using the substrates as follows: AMP, CMP, GMP, IMP, UMP, and glucose 6-phosphate (G6P) (Fig. 2). G6P was included because it is the substrate of certain HADSF enzymes [9]. YcsE appeared to be the most efficient phosphatase acting on all six substrates at a lower concentration (2 mM), indicating its broad substrate specificity with preference for IMP, CMP, and G6P. Its activity increased in the presence of 10 mM substrates, particularly AMP, GMP, and UMP. The other four enzymes were relatively less active even at the higher substrate concentration. All 5′-nucleotides served almost equally as substrates for AraL at lower and higher concentrations, and AraL activity was enhanced particularly against G6P at the higher concentration. YutF exhibited purine 5′-nucleotidase activity with the substrates GMP and IMP but only at the higher concentration, and did not hydrolyze pyrimidine nucleotides and G6P. YqeG hydrolyzed GMP and G6P only at the higher concentration. YsaA activity was weakest with purine-nucleotide substrates at the higher concentration.

The catalytic properties of YcsE are shown in Table 1. The substrate specificity of YcsE was broad with higher $K_M$ values, suggesting that it efficiently dephosphorylated these substrates at higher concentrations. In contrast, the properties of the other four enzymes were not determined because of their lower activities. Nevertheless, the data do not exclude their physiological function as 5′-nucleotidases, because the activities of two of the 5′-nucleotidases YieH and Yigl identified previously in E. coli were too low to determine their enzymatic properties [9].

5′-Nucleotidase activity of SuhB homolog in B. subtilis

As described above, E. coli SuhB is an inositol monophosphatase, which exhibits wider substrate specificity, as it hydrolyzes β-glycerophosphate and adenosine 2′-monophosphate as well [18]. Thus, we hypothesized that bacterial inositol monophosphatases might also function as 5′-nucleotidase.

The B. subtilis yktC gene, which is the homolog of E. coli suhB (31.7 % similarity of 265 overlapping amino acid residues), was cloned and expressed as a His6-tagged protein in E. coli. His6-tagged YktC was purified to form a single band in SDS-PAGE (data not shown) and was subjected to phosphatase assays in the presence of various substrates (Fig. 2, Table 2). The inositol monophosphatase activity of YktC was indeed demonstrated by its efficient ability to hydrolyze MIMP, and it also turned out to exhibit 5′-nucleotidase activity preferentially against IMP and GMP as substrates. To our knowledge, this is the first identification of an inositol monophosphatase with 5′-nucleotidase activity.

Compared with E. coli SuhB, YktC hydrolyzed G6P less efficiently, while β-glycerophosphatase more efficiently [18]. Expression of yktC is mainly constitutive and is markedly enhanced by stressors such as diamide, ethanol, or high salt as well as at higher and lower temperatures [25].
Physiological functions of YqeG, YcsE, and YktC

As mentioned above, yqeG expression is enhanced by oxidative stress, ycsE is induced in the presence of diamide or ethanol, and yktC is markedly enhanced by stressors such as diamide, ethanol, or high salt, as well as at higher and lower temperatures [25]. These findings inspired us to investigate the three genes encoding 5′-nucleotidase for their possible functional association with the response to oxidative stress.

We used conventional marker replacement to inactivate ycsE and yktC. The yqeG gene is the first of the long operon comprising yqeG, yqeH, aroD, yqeL, nadD, yqeK, yqeL, and yqeM [26]. At least yqeH, yqeI, and nadD are essential genes [26]. Therefore, we tried to introduce an in-frame deletion of yqeG to maintain the expression of the downstream essential genes. However, numerous attempts failed. Therefore, once an additional copy of yqeG was introduced into the amyE locus under the

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**Table 1** Kinetic constants of YcsE

| Substrate | $K_M$ (mM)$^a$ | $V_{max}$ (μmol min$^{-1}$ mg protein$^{-1}$)$^a$ | $k_{cat}$ (s$^{-1}$)$^a$ |
|-----------|----------------|---------------------------------|-----------------------|
| IMP       | 3.2 ± 0.036    | 0.13 ± 0.060                     | 3.7 ± 1.4 × 10$^2$    |
| AMP       | 8.6 ± 0.80     | 0.50 ± 0.029                     | 2.3 ± 0.13 × 10$^3$   |
| GMP       | 10.3 ± 0.12    | 0.45 ± 0.12                      | 2.0 ± 0.55 × 10$^3$   |
| CMP       | 4.7 ± 0.23     | 0.53 ± 0.18                      | 3.4 ± 0.011 × 10$^3$  |
| UMP       | 8.5 ± 0.69     | 0.20 ± 0.072                     | 9.6 ± 3.5 × 10$^2$    |
| G6P       | 22.2 ± 0.96    | 2.3 ± 0.48                       | 1.06 ± 0.22 × 10$^4$  |

$^a$All presented data are the mean values of three independent experiments ± SD
control of \textit{spac} promoter, which is conditionally induced by \textit{yqeG}. And then, in the presence of IPTG, the existing \textit{yqeG} gene was deleted using marker replacement so that the genes downstream were constitutively expressed by \textit{yqeG} by IPTG. And then, in the presence of IPTG, the existing \textit{yqeG} might be dispensable in planktonic growth. However, the \textit{spac} promoter is somewhat leaky; a significant basal level of expression still exists in the absence of IPTG [30]. Therefore, in the absence of IPTG, there might be some production of YqeG, and this might be sufficient for planktonic growth. Nevertheless, NON05 formed fewer colonies in the absence of IPTG (Fig. 4), suggesting that \textit{yqeG} was required to maintain normal growth particularly on solid medium. Similarly, less efficient colony-formation was exhibited by the triple mutant NON06 in the absence of diamide (Fig. 3), and the other enzymes may function as two of the main enzymes involved in intracellular nucleoside/nucleotide metabolism. The other enzymes similar to \textit{E. coli} \textit{NagD}, which belong to the HADSF family (\textit{YutF}, \textit{YsaA}, \textit{YqeG}, and \textit{AlaL}), exhibited lower activity, although the data suggest they may function as minor \textit{5′}-nucleotidases. HADSF phosphatases often have broader substrate specificity and hydrolyze various phosphorylated compounds [5]. Thus, \textit{5′}-nucleotidases with lower substrate affinity might contribute to the balance of intracellular nucleotide concentrations as well as that of other phosphorylated metabolites. The \textit{K_M} values of \textit{E. coli} \textit{NagD} for \textit{UMP}, \textit{CMP}, \textit{GMP}, \textit{AMP}, and \textit{G6P} are $0.16 \pm 0.038$, $1.47 \pm 0.044$, $0.40 \pm 0.130$, $0.84 \pm 0.250$, and $5.90 \pm 0.750$ mM, respectively [4], which are smaller than those of \textit{YcsE} and \textit{YktC} (Tables 1 and 2, respectively). Therefore, \textit{B. subtilis} \textit{YktC} and \textit{YcsE} functioned as \textit{5′}-nucleotidases, suggesting that they may represent two of the main enzymes involved in intracellular nucleoside/nucleotide metabolism. The other enzymes similar to \textit{E. coli} \textit{NagD}, which belong to the HADSF family (\textit{YutF}, \textit{YsaA}, \textit{YqeG}, and \textit{AlaL}), exhibited lower activity, although the data suggest they may function as minor \textit{5′}-nucleotidases. HADSF phosphatases often have broader substrate specificity and hydrolyze various phosphorylated compounds [5]. Thus, \textit{5′}-nucleotidases with lower substrate affinity might contribute to the balance of intracellular nucleotide concentrations as well as that of other phosphorylated metabolites. The \textit{K_M} values of \textit{E. coli} \textit{NagD} for \textit{UMP}, \textit{CMP}, \textit{GMP}, \textit{AMP}, and \textit{G6P} are $0.16 \pm 0.038$, $1.47 \pm 0.044$, $0.40 \pm 0.130$, $0.84 \pm 0.250$, and $5.90 \pm 0.750$ mM, respectively [4], which are smaller than those of \textit{YcsE} and \textit{YktC} (Tables 1 and 2, respectively). Therefore, \textit{B. subtilis} \textit{YktC} and \textit{YcsE} might be less efficient as \textit{5′}-nucleotidase than \textit{NagD}, but it is still suggestive that they may be able to function as two of the \textit{5′}-nucleotidases that are the first to respond to the accumulation of nucleotides, and the other enzymes may function when the concentration of nucleotides is in far excess.

Some of the genes encoding \textit{5′}-nucleotidases might be induced upon partial nucleotide limitation as previously reported [20]. Interestingly, the expression of \textit{yqeG} is enhanced under oxidative stress, that of \textit{ycsE} is induced by diamide or ethanol, and that of \textit{yktC} is markedly enhanced in the presence of the various stressors, including

\begin{table}[h]
\centering
\caption{Kinetic constants of \textit{YktC}}
\begin{tabular}{|l|c|c|c|}
\hline
Substrate & \textit{K_M} (mM) & \textit{V_{max}} (\mu mol min^{-1} mg protein^{-1}) & \textit{k_{cat}} (s^{-1}) \\
\hline
\textit{MIMP} & 0.076 ± 0.006 & 0.82 ± 0.16 & 2.9 ± 0.33 × 10^3 \\
\textit{IMP} & 1.1 ± 0.083 & 0.078 ± 0.0084 & 1.8 ± 0.59 × 10^2 \\
\textit{AMP} & 1.8 ± 0.13 & 0.030 ± 0.0022 & 5.3 ± 0.039 × 10^1 \\
\textit{GMP} & 1.9 ± 0.026 & 0.169 ± 0.028 & 3.0 ± 0.34 × 10^2 \\
\textit{CMP} & 1.6 ± 0.017 & 0.049 ± 0.014 & 1.5 ± 0.52 × 10^2 \\
\textit{UMP} & 2.4 ± 0.052 & 0.11 ± 0.0099 & 9.8 ± 4.6 × 10^1 \\
\textit{GTP} & 2.6 ± 0.51 & 0.021 ± 0.010 & 3.7 ± 1.3 × 10^1 \\
\textit{β-glycerophosphate} & 0.49 ± 0.035 & 1.2 ± 0.32 & 5.7 ± 1.2 × 10^1 \\
\hline
\end{tabular}
\\textsuperscript{a}All presented data are the mean values of three independent experiments ± SD.
\end{table}
Fig. 3 Growth curves of *B. subtilis* strains. Strains 168 (a), NON01 (b ΔycsE), NON02 (c ΔyktC), NON03 (d ΔycsE ΔyktC), NON05 (e and f ΔyqeG Pspac-yqeG), and NON06 (g and h, ΔycsE ΔyktC ΔyqeG Pspac-yqeG) were inoculated into liquid medium and their growth was monitored. At the times indicated by the arrowheads, diamide was added to final concentrations of 0 mM (open circle), 1 mM (solid square), and 4 mM (solid triangle). Strains NON05 and NON06 were grown in the presence (e and g) and absence (f and h) of 1 mM IPTG. All experiments were repeated more than three times and similar results were observed.
diamide, ethanol, and high salt as well as at higher and lower temperatures [25]. These findings indicate that the intracellular levels of phosphorylated metabolites and nucleotides including 5′-nucleotides may be altered in the presence of these stressors, in particular oxidative stress. Our results show that inactivation of yktC led to increased sensitivity to diamide (Fig. 4). When the expression of yqeG was artificially modified, the cells grew normally in liquid medium and did not exhibit increased sensitivity to diamide, depending on repression of yqeG in the absence of IPTG (Fig. 4). Further, in the presence of diamide, the triple mutant NON06, lacking ycsE and yktC and with repressed yqeG, exhibited a growth defect similar to that of NON02 or NON03, which lacks yktC (Fig. 4). Therefore, we conclude that only YktC was involved in resistance to diamide-induced oxidative stress. However, further studies are required to elucidate how its enzymatic functions, including its 5′-nucleotidase activity, are related to resistance to the oxidative stress.

*B. subtilis* induces S-cysteinylation to protect protein thiols after exposure to diamide, because this bacterium does not produce thiols such as glutathione [35]. To our knowledge, enzymes with 5′-nucleotidase activity have not been demonstrated to be involved in resistance to oxidative stress. In response to oxidative stress, *Pseudomonas fluorescens* evokes a metabolic adaptation to increase NADPH synthesis and to decrease NADH production, accompanied by the induction and repression, respectively, of NAD⁺ kinase and NADP⁺ phosphatase, which regulate the levels of NAD⁺ and NADP⁺ [36]. Namely, oxidative stress represses the NADP⁺ phosphatase in *P. fluorescens* [36], whereas it induces yktC in *B. subtilis* [25]. If YktC might function as NADP⁺ phosphatase, it could decrease NADPH synthesis under oxidative stress, which is contradictory to the physiological demand. Therefore, YktC is unlikely to act as a NADP⁺ phosphatase, and we did not determine if NADP⁺ was a substrate of the phosphatase activity of YktC. On the other hand, yqeG is the first gene of a long operon comprising essential genes such as nadD that encodes nicotinamide-nucleotide adenyllyltransfärse required for de novo biosynthesis of NAD⁺ and NADP⁺ [37]. The induction of yqeG by oxidative stress, together with the other members of the operon, might be relevant to the enhanced NADPH synthesis as found in *P. fluorescens*. However, our results indicate that yqeG was dispensable, except for normal colony formation on solid medium (Fig. 4). We were therefore unable to determine the physiological function of yqeG.

In mammalian and plant cells, inositol monophosphates play important roles in the metabolism of inositol polyphosphates and phosphatidylinositol [13, 38]. De
**novo** phosphatidylinositol synthesis in mammals and *Saccharomyces cerevisiae* involves a reaction that combines CDP-diacylglycerol with *myo*-inositol. *Myo*-inositol is produced through the isomerization of glucose 6-phosphate to form MIMP [39] and the subsequent dephosphorylation of MIMP by inositol monophosphatase [16]. Further, MIMP is produced by the dephosphorylation of the second messenger inositol polyphosphates [40]. In *E. coli* and *B. subtilis*, however, phosphatidylinositols are not components of phospholipids usually present in the cell membrane [41, 42], and inositol 1-phosphate synthase has not been identified. Therefore, it is unlikely that MIMP occurs naturally in these bacteria and might serve as the physiological substrate of MIMP [43, 44]. The secY gene encodes a membrane component involved in protein secretion [45], and rpoH encodes σ32, a component of the RNA polymerase involved in heat-shock induction [46]. Therefore, defects in secY and rpoH impair secretion and the heat-shock response, respectively, which are involved in posttranslational control of mRNAs [18]. Therefore, SuhB may control the rates of peptide chain elongation and protein folding [42]. In contrast, the SuhB homolog of *P. aeruginosa* plays an important role in pathogenesis to control the genes required for acute and chronic infection [47]. *Burkholderia cepacia* SuhB is required for the secretion of proteins associated with motility and biofilm formation [48], suggesting that the physiological functions of SuhB homologs are complex, which may be true for *B. subtilis* YktC. Nevertheless, it is intriguing that MIMP is the best substrate for these enzymes. Kozloff and the colleagues detected a small amount of phosphatidylinositol from *E. coli* cells, implying the possibility of a weak phosphatidylinositol biosynthesis under some specific conditions [49]. *E. coli* SuhB and *B. subtilis* YktC might be involved in bacterial metabolism of phosphatidylinositol, which has not been characterized yet.

The broad substrate specificity of YktC revealed here implies the association of this enzyme with sugar-phosphate stress. Although sugars serve as energy and carbon sources, sugar phosphates are produced during their metabolism. Excess accumulation of sugar phosphates impairs cell growth [50] and may trigger cell death [51, 52] through an unknown mechanism. Similarly, accumulation of nucleotides is detrimental to the cell [34]. Therefore, by hydrolyzing sugar phosphates as well as nucleotides, YktC may play a role in correcting imbalances of metabolites. For example, we found that the *K_M* value of YktC for G6P was 2.6 ± 0.51 mM, which is greater than the reported intracellular concentrations of glucose 6-phosphate in *E. coli* (0.8–2.0 mM) [53]. However, under certain conditions, the intracellular concentration of G6P in *Lactococcus lactis* reaches 20–50 mM [54], indicating that G6P accumulates to higher levels in bacterial cells. Accordingly, in the presence of excess intracellular concentrations of G6P, YktC may hydrolyze it to intervene in glucose metabolism. This may be true as well for YcsE, because it hydrolyzed G6P in vitro more efficiently than YktC (Fig. 2). Similarly, YktC may hydrolyze β-glycerophosphate, which is a biosynthetic precursor of phospholipids [55], to interfere with the maintenance of the cell membrane. Therefore, their phosphatase activities with broad-specificity may exert pleiotropic effects on metabolism and other cellular functions. It would be worthwhile to perform metabolomic analyses on the mutants lacking yktC and ycsE to determine concentrations of the various metabolites, including 5′-nucleotides and other sugar phosphates, which would reinforce the hypothesis concerning the role of these enzymes.

Together with sodium glutamate, IMP and GMP are generally used as food additives to enhance taste. Industrial production of IMP is achieved using enzymatic conversion of inosine into IMP. *B. subtilis* strains producing large amounts of inosine in the fermentation medium were generated [56–60], and 5′-nucleotidase activity of these strains is remarkably increased, which may convert IMP into inosine [59]. Thus, optimizing the expression of genes encoding 5′-nucleotidases such as those studied here may further improve the fermentation of inosine production.

**Conclusions**

*B. subtilis* ycsE, araL, ytfF, ysaA, and yqeG show suggestive similarities to *E. coli* nagD encoding 5′-nucleotidase. Among the five, only YcsE exhibited significant 5′-nucleotidase activity with a broader specificity, whereas the other four enzymes had rather weak but suggestive activities with various capacities and substrate specificities. Interestingly, YqeG was required to maintain normal growth on solid medium. On the other hand, *B. subtilis* yktC shares high similarity with *E. coli* suhB encoding inositol monophosphatase, the gene product of which had 5′-nucleotidase activity preferential for GMP and IMP, and turned out to be involved in resistance to oxidative stress.

**Methods**

**Bacterial strains, plasmids, oligonucleotide primers, and culture conditions**

Bacterial strains and plasmids are listed in Table 3. Bacterial strains were usually grown aerobically at 37 °C and maintained in Luria-Bertani (LB) medium (Becton Dickinson, NJ, USA) containing 50 mg/l ampicillin and 50 mg/l kanamycin as required. Oligonucleotide primers are listed in Table 4. To assess the resistance to oxidative stress of *B. subtilis* strains, cells were inoculated into LB and allowed to grow. The oxidizing agent diamide was
added to the exponentially growing culture at various concentrations, and growth was continuously monitored.

**Construction of bacterial strains**

*E. coli* strains expressing *B. subtilis* genes were constructed as follows: DNA fragments corresponding to the open reading frames of genes of interest were PCR-amplified using *B. subtilis* strain 168 chromosomal DNA (Table 3) as template and pairs of respective oligonucleotide primers with generation of unique restriction sites at their 5′-termini (araL-F/araL-R for araL, ycsE-F/ycsE-R for ycsE, yktC-F/yktC-R for yktC, yqeG-F/yqeG-R for yqeG, ysaA-F/ysaA-R for ysaA, and yutF-F/yutF-R for yutF) (Table 4). Each fragment was cloned into the pMD20-T plasmid using the Mighty TA-cloning kit (Takara Bio, Otsu, Japan) to yield pMD20-araL, pMD20-ycsE, pMD20-yktC, pMD20-yqeG, pMD20-ysaA, and pMD20-yutF, which were used to transform *E. coli* strain DH5α. After performing nucleotide sequencing to confirm the constructs, each recombinant plasmid was digested with the appropriate restriction enzyme(s) to prepare a DNA fragment corresponding to its respective open reading frame and then ligated to the arms of pET28b (Takara Bio) digested with the same enzyme(s). The plasmids pET28b-araL, pET28b-ycsE, pET28b-yktC, pET28b-yqeG, pET28b-ysaA, and pET28b-yutF were constructed to express each protein as a C-terminal His6-fusion, and expression was controlled by the T7 promoter in *E. coli* strain BL21(DE3) (Takara Bio).

Mutant strains of *B. subtilis* lacking ycsE and yktC were constructed using conventional marker replacement as follows (Additional file 1: Figure S1): To delete

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**Table 3** Bacterial strains and plasmids

| Strain and plasmid | Description | Source or reference |
|--------------------|-------------|---------------------|
| B. subtilis        |             |                     |
| 168                | trpC2       | Laboratory stock    |
| NON01              | trpC2 ycsE:spc | This study       |
| NON02              | trpC2 yktC:cat | This study       |
| NON03              | trpC2 ycsE:spc yktC:cat | This study       |
| NON04              | trpC2 amyE:(Pspac-yqeG erm) | This study       |
| NON05              | trpC2 amyE:(Pspac-yqeG erm) ΔyqeG:kan | This study       |
| NON06              | trpC2 ycsE:spc yktC:cat amyE:(Pspac-yqeG lacI erm) ΔyqeG:kan | This study       |
| TMO310             | trpC2 aptE:(spec lacI Pspac-mazF) | [62]             |
| TMO311             | trpC2 aptE:(kan lacI Pspac-mazF) | [62]             |
| E. coli            |             |                     |
| DH5α               | F− φ80lacZAM15 Δ(lacZYA-argF) U169 recA1 endA1 hsdR17(k−, mK') phoA supE44 λ− thi-1 gyrA96 relA1 | Laboratory stock |
| BL21(DE3)          | F− ompT gal dcm lon hsdS30(rK2, mC2) λ(DE3 [lacI lacUV5-T7p07 ind1 sam7 ninS]) [malB8][k12]λ3 | Laboratory stock |
| **Plasmids**       |             |                     |
| pCRE-test          | cat         | [63]                |
| pET28b             | kan         | Takara Bio          |
| pET28b-araL        | kan, araL   | This work           |
| pET28b-yutF        | kan, yutF   | This work           |
| pET28b-yqeG        | kan, yqeG   | This work           |
| pET28b-ysaA        | kan, ysaA   | This work           |
| pET28b-ycsE        | kan, ycsE   | This work           |
| pET28b-yktC        | kan, yktC   | This work           |
| pMD20-araL         | amp         | This work           |
| pMD20-yutF         | amp         | This work           |
| pMD20-yqeG         | amp         | This work           |
| pMD20-ysaA         | amp         | This work           |
| pMD20-ycsE         | amp         | This work           |
| pMD20-yktC         | amp         | This work           |
| pMutin2            | erm lacI    | [64]                |
**Table 4 Oligonucleotide primers**

| Name | Sequence (5’→3’)* | Restriction site |
|------|-------------------|------------------|
| araL-F | gggatccatatgccgtaaatggccgtagctgat | NdeI |
| araL-R | ccgctctagcatcaatggcagctgat | XhoI |
| yufF-F | cgccgatcatgaaacatataaggtgatta | BamHI |
| yufF-R | ccgcaaccatgaatgagatcagccgtgctta | HindIII |
| ysaA-F | cgccgatccatgaaacagctgtaatggggtattta | BamHI |
| ysaA-R | ccgctctagctgctatagtggaaccttcca | XhoI |
| YqEG-F | gggatccatatgtatatcagccggaagccggttaatggctta | NdeI |
| YqEG-R | ccgaaaccctcctcctgctgtaatggggtattta | HindIII |
| yscE-F | gggatccatatgtctggcagctgcttta | BamHI |
| yscE-R | ccgctctagctgctatagtggaaccttcca | XhoI |
| yksE-F | ggtgatccatatgtatatcagccggaagccggttaatggctta | NdeI |
| yksE-R | ccgaaaccctcctcctgctgtaatggggtattta | HindIII |
| yksE-3 | aacagtcttttcctcgctgctgtaatggggtattta | BamHI |
| yksE-4 | ggtgatccatatgtatatcagccggaagccggttaatggctta | NdeI |
| yksE-5 | ccgctctagctgctatagtggaaccttcca | XhoI |

*Restriction enzyme cleavage sites are underlined

*ycsE*, two PCR fragments (each approximately 700-bp) corresponding to the upstream and downstream regions of *ycsE* were amplified from the DNA of *B. subtilis* strain 168 using the primer pairs *ycsE*-1/*ycsE*-2 and *ycsE*-5/*ycsE*-6, respectively (Table 4). A PCR fragment containing the spectinomycin-resistance gene cassette was amplified...
from the DNA of strain TMO310 (Table 3) using the primer pair ycsE-3/yscE-4 (Table 4). The three PCR fragments were ligated using recombinant PCR with the primer pair ycsE-1/ycsE-6 to generate a single PCR fragment comprising the spectinomycin-resistance gene cassette flanked by the upstream and downstream regions of ycsE. B. subtilis strain 168 was transformed with the recombinant PCR fragment to confer resistance to spectinomycin, and a transformant with correct resistance to both chloramphenicol and spectinomycin inactivating yktC and ycsE, respectively (strain NON06).

Chromosomal DNAs of mutant strains of B. subtilis were subjected to diagnostic PCR experiments with the respective primer pairs used for recombinant PCR, and their correct construction was confirmed by appearance of the respective PCR fragments with altered length (Additional file 1: Figure S1).

Purification of gene products produced in E. coli

E. coli strain BL21(DE3) bearing the pET28b derivatives were inoculated into 600 ml of LB medium containing kanamycin and were cultured at 37 °C with shaking at 180 rpm. When the optical density of the cultures reached 0.4–0.6, IPTG was added to a final concentration of 1 mM, the cells were cultivated further for 5 h, harvested using centrifugation, and resuspended into 10.8 ml of NP buffer (50 mM NaH₂PO₄, pH 8.0, and 300 mM NaCl). The suspension was incubated in an ice bath, mixed with 1.2 ml of 10 mg/ml lysozyme in NP buffer, incubated for 30 min, and sonicated to disrupt the cells. After centrifugation, the supernatant was subjected to His₆-tag affinity purification using TALON Metal Affinity Resins (Takara Bio). The His₆-tagged proteins were purified in a reaction buffer containing 1 mM NH₄Cl, 250 mM KCl, 10 mM MgCl₂, and 50 mM Tris–HCl (pH 7.8), and subjected to SDS-PAGE followed by Coomassie brilliant blue staining.

Enzyme assay

The phosphatase assay was performed as previously described [61]. Each purified enzyme (0.25 mg/ml) was incubated with substrate in 50 µl of reaction buffer containing 1 mM NH₄Cl, 250 mM KCl, 10 mM MgCl₂, and 50 mM Tris–HCl (pH 7.8) at 37 °C, and the reaction was terminated using 50 µl of 10 % trichloroacetic acid. After centrifugation, the supernatant was mixed with 20 µl of solution A (10 % ascorbic acid in 2.25 M H₂SO₄) and 20 µl of solution B (2.4 % ammonium molybdate and 1 mg/ml potassium antimonyl tartrate in 2.25 M H₂SO₄). The mixture was incubated at room temperature for 10 min, and absorbance at 820 nm was measured to calculate the concentration of the released phosphate ion with reference to a standard curve made with various concentrations of inorganic phosphate. Trace of phosphate released in the reaction mixtures in the absence of purified enzyme was subtracted as background.
Additional file

Additional file 1: Figure S1. Genetic organization of the altered chromosomal loci; ΔycsE::spc(A), ΔyktC::can(B), ΔyqeG::kan(C), and amyE::(Pspac-yeg G lacI erm)(D). Genes and primers are indicated schematically by thick arrows and arrowheads, respectively. (PDF 144 kb)

Abbreviations
G6P: Glucose 6-phosphate; HADSF: Haloalkane acid dehalogenase superfamily; IPTG: Isopropyl β-D-thiogalactopyranoside; MIMP: Myo-inositol 1-monophosphate; SDS-PAGE: Sodium dodecyl sulfate polyacrylamide gel electrophoresis

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Availability of data and materials
The datasets during and/or analyzed during the current study are available from the corresponding author on reasonable request. The phylogenetic tree, sequence data, and alignments used to produce the results displayed in Fig. 1 are deposited in TreeBASE (http://purl.org/phylo/treebase/phylows/ study/TB2S1Z9999).

Authors’ contributions
AT conducted most of the experiments and analyzed the results. The enzymatic experiments were performed under the supervision of KT and ST. KT, AN, AO, SN, and JK conducted experiments with the mutant strains of B. subtilis. KT conceived the idea for the project and wrote the final manuscript with ST. All authors read and approved the final manuscript.

Competing interests
The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

Consent for publication
Not applicable.

Ethics approval and consent to participate
Not applicable.

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References
1. Rampazzo C, Ferraro P, Pontarin G, Fabris S, Reichard P, Bianchi V. Mitochondrial deoxyribonucleotidyltransferases, pool sizes, synthesis, and regulation. J Biol Chem. 2004;279:17019–26.
2. Tamaya Y, Ogata Y, Nakamura T, Yamada I, Hasegawa T, Tanaka H, Tsuda M. Tschuha T. Sequence analysis of nuSI gene encoding membrane-bound Cl(−)-dependent S'-nucleotidase of Vibrio paraahemiobacter. J Biochem. 1988;109:292–4.
3. Burns DM, Beacham IR. Nucleotide sequence and transcriptional analysis of the E. coli ushA gene, encoding periplasmic UDP-sugar hydrolase (S'-nucleotidase); regulation of the ushA gene, and the signal sequence of its encoded protein product. Nucleic Acids Res. 1986;14:3743–42.
4. Tremblay LW, Dunaway-Mariano D, Allen KN. Structure and activity analyses of Escherichia coli K-12 NagD provide insight into the evolution of biochemical function in the haloalkanic acid dehalogenase superfamily. Biochemistry. 2006;45:1183–93.
5. Koonin EV, Tatusov RL. Computer analysis of bacterial haloacid dehalogenases defines a large superfamily of hydrolases with diverse specificity: Application of an iterative approach to database search. J Mol Biol. 1994;244:125–32.
6. Allen KN, Dunaway-Mariano D. Phosphoryl group transfer: evolution of a catalytic scaffold. Trends Biochem Sci. 2004;29:495–503.
7. Lu Z, Dunaway-Mariano D, Allen KN. HAD superfamily phosphotransferase substrate diversification: structure and function analysis of HAD subcellular SB sugar phosphate βT431. Biochemistry. 2005;44:8684–96.
8. Shin DH, Roberts A, Janczak J, Yokota H, Kim R, Wemmer DE, Kim SH. Crystal structure of a phosphatase with a unique substrate binding domain from Thermotoga maritima. Protein Sci. 2003;12:1464–72.
9. Kuznetsova E, Proudfoot M, Gonzalez CF, Brown G, Omlenchko MV, Boroian I, Carmel L, Wolf Y, Mori H, Savchenko AV, Arrowmith SM, Koonin EV, Edwards AM, Yakunin AF. Genome-wide analysis of substrate specificities of the Escherichia coli haloacid dehalogenase-like phosphatase family. J Biol Chem. 2006;281:36149–61.
10. Proudfoot M, Kuznetsova E, Brown G, Rao NN, Kitagawa M, Mori H, Savchenko A, Yakunin AF. General enzymatic screens identify three new nucleotidases in Escherichia coli. Biochemical characterization of SufE, YBR, and YDR687C. J Biol Chem. 2006;281:36149–61.
