

The γ -Aminobutyrate Permease GabP Serves as the Third Proline Transporter of *Bacillus subtilis*

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PutP and OpuE serve as proline transporters when this imino acid is used by *Bacillus subtilis* as a nutrient or as an osmostress protectant, respectively. The simultaneous inactivation of the PutP and OpuE systems still allows the utilization of proline as a nutrient. This growth phenotype pointed to the presence of a third proline transport system in *B. subtilis*. We took advantage of the sensitivity of a *putP opuE* double mutant to the toxic proline analog 3,4-dehydro-DL-proline (DHP) to identify this additional proline uptake system. DHP-resistant mutants were selected and found to be defective in the use of proline as a nutrient. Whole-genome resequencing of one of these strains provided the lead that the inactivation of the γ -aminobutyrate (GABA) transporter GabP was responsible for these phenotypes. DNA sequencing of the *gabP* gene in 14 additionally analyzed DHPresistant strains confirmed this finding. Consistently, each of the DHP-resistant mutants was defective not only in the use of proline as a nutrient but also in the use of GABA as a nitrogen source. The same phenotype resulted from the targeted deletion of the *gabP* gene in a *putP opuE* mutant strain. Hence, the GabP carrier not only serves as an uptake system for GABA but also functions as the third proline transporter of *B. subtilis*. Uptake studies with radiolabeled GABA and proline confirmed this conclusion and provided information on the kinetic parameters of the GabP carrier for both of these substrates.

The soil bacterium *Bacillus subtilis* lives in a taxing habitat where many microorganisms compete with each other for specific ecological microniches and nutritional resources (1, 2). Its genome sequence (3) bears the hallmarks of a bacterium that lives in association with plants and plant detritus (4, 5). It is therefore understandable that *B. subtilis* devotes a considerable portion of its genome coding capacity to direct the synthesis of transporters for the uptake of a wide variety of plant-derived compounds for use as nutrients (6) or as stress protectants (7–9).

L-Proline serves both as a nutrient and as a stress protectant for B. subtilis since the cell can exploit it not only as a sole carbon, nitrogen, and energy source (10-12) but also as an osmostressrelieving compound (13-16). To fuel protein synthesis, B. subtilis produces proline from the precursor glutamate (17, 18), a pathway that is present in many bacterial species (19). The anabolic proline synthesis route (ProB-ProA-ProI [ProG]) is interconnected with the osmostress-relieving production route for proline (ProJ-ProA-ProH) via the y-glutamyl phosphate reductase (ProA) (14). The latter pathway is osmotically controlled and allows B. subtilis to produce very large amounts of the compatible solute proline (20) to fend off the detrimental effects of high salinity on cellular hydration, turgor, and physiology (13–15, 21). Proline-mediated osmoprotection can also be achieved through uptake; the osmotically inducible OpuE transporter of B. subtilis serves this function (15, 16). OpuE also operates as a recapturing device for newly synthesized proline that is released from B. subtilis grown under high-salinity conditions, probably in an effort by the cell to fine-tune turgor (22). Expression of opuE is osmotically inducible (16, 23, 24), but there is no stimulation of opuE expression in response to proline availability.

Externally provided proline not only affords osmostress protection (15, 16), but it also can serve as the sole carbon, energy, and nitrogen source for *B. subtilis* (10–12). Utilization of proline as a nutrient requires its capture from environmental sources, such as root exudates and organic deposits in the rhizosphere (2, 25), and relies on the PutB- and PutC-mediated catabolism to glutamate (10), a central intermediate in the interconnected carbon and nitrogen utilization systems of *B. subtilis* (26, 27). The PutP transporter mediates the uptake of proline for its use as a nutrient, and the induction of the expression of the catabolic *putBCP* operon by an external supply of the substrate proline reflects this role (10, 28, 29).

The two functionally characterized proline import systems of B. subtilis, PutP and OpuE (10, 16), are members of the solute: sodium symporter family (SSS; Transporter Classification Database [TC] accession number 2.A.21) (30) that mediate the import of their substrates tightly coupled to Na^+ symport (31–33). The PutP and OpuE proteins are closely related to each other (61% amino acid sequence identity) and to the biochemically well-studied PutP proline permease from Escherichia coli and Salmonella enterica serovar Typhimurium (31, 33-37). Both the B. subtilis PutP and the OpuE L-proline transporters possess a high affinity for their substrate, with K_m values being in the low μ M range, and they exhibit a substantial transport capacity (10). However, their transport profiles differ significantly. The proline import activity of PutP, as reflected by the Vmax value, is upregulated when proline is present in the environment, and it is gradually inhibited when the salinity of the growth medium is increased (10). Conversely, not only can the proline import activity of OpuE withstand salt stress, but also it is actually increased in response to high osmo-

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TABLE 1 B. subtilis strains used in this study

Strain ^a	Relevant genotype ^b	Source or reference
JH642	trpC2 pheA1	J. Hoch
ACB164	Δ (<i>ywcA</i> :: <i>ery</i>)1	This study
ACB165	Δ (yodF::neo)1	This study
ACB199	$\Delta(putP::spc) 1 \Delta(opuE::tet) 1 \Delta(ywcA::ery) 1 \Delta(yodF::neo) 1$	This study
ACB214	$\Delta(putP::spc) 1 \Delta(opuE::tet) 1 \Delta(ywcA::ery) 1 \Delta(yodF::neo) 1 gabP1$	This study
ACB274	$\Delta(gabP::zeo)$ 1	This study
ACB275	$\Delta(opuE::tet) 1 \Delta(gabP::zeo) 1$	This study
ACB276	$\Delta(opuE::tet) 1 \Delta(gabP::zeo) 1 \Delta(treA::neo) 1 [amyE::\phi(putB'-treA) 1 cat]$	This study
ACB277	$\Delta(putP::spc) \mid \Delta(gabP::zeo) \mid$	This study
ACB278	$\Delta(putP::spc) 1 \Delta(gabP::zeo) 1 \Delta(treA::neo) 1 [amyE::\phi(putB'-treA) 1 cat]$	This study
ACB279	$\Delta(putP::spc) 1 \Delta(opuE::tet) 1 \Delta(gabP::zeo) 1$	This study
ACB288	$\Delta(putP::spc) \Delta(opuE::tet) \Delta(treA::neo) [amyE::\phi(putB'-treA) cat]$	This study
ACB292	$\Delta(putP::spc)1 \Delta(opuE::tet)1 \Delta(gabP::zeo)1 \Delta(treA::neo)1 [amyE::\phi(putB'-treA)1 cat]$	This study
BLOB9	$\Delta(opuE::tet)$ 1	16
JSB8ery	Δ (<i>proHJ</i> :: <i>ery</i>)1	J. Brill
MBB1	$\Delta(treA::neo)$ 1	M. Brosius
SMB10	Δ (<i>treA</i> :: <i>neo</i>)1 [<i>amyE</i> :: ϕ (<i>putB'</i> - <i>treA</i>)1 <i>cat</i>]	10
SMB11	$\Delta(putP::spc)$ 1	10
SMB12	$\Delta(putP::spc) 1 \Delta(opuE::tet) 1$	10

^{*a*} All strains are derivatives of the *B. subtilis* wild-type laboratory strain JH642 (52) and therefore carry the *trpC2* and *pheA1* mutations, in addition to the indicated genetic markers. ^{*b*} The designation *amyE*:: $\phi(putB'-treA)1$ *cat* indicates that the *putB-treA* operon reporter fusion is stably integrated into the chromosomal *amyE* gene as a single copy, thereby rendering the fusion strains defective in the extracellular α -amylase AmyE. The $\phi(putB'-treA)1$ reporter fusion is genetically linked to a chloramphenicol resistance gene (*cat*).

larity (38). Underlying these characteristics of the PutP- and OpuE-mediated proline import are influences that result from the different transcriptional profile of the *putBCP* and *opuE* genes in response to proline availability in the growth medium and its osmolarity (10, 16, 23, 24). However, posttranscriptional effects of high salinity on the transport activity of PutP and OpuE are also noticeable (10, 16).

In connection with the analysis of the utilization of proline as a nutrient by *B. subtilis*, Moses et al. (10) found that a *putP opuE* double mutant strain was still prone to growth inhibition by L-aze-tidine-2-carboxylic acid (AC) and 3,4-dehydro-DL-proline (DHP), toxic proline analogs (39) that can be imported into microbial cells via different types of proline transport systems (36, 40, 41). It was also noted in this study that the expression of the *putBCP* operon remained inducible by an external supply of proline in a *putP opuE* double mutant (10). These two observations therefore hint at the operation of a further proline import system in *B. subtilis*.

Here, we identify this third proline importer using a genetic selection for resistance to DHP and genome resequencing to provide a lead as to the identity of the mutation underlying the loss of the ability to use proline as a nutrient in the absence of functional PutP and OpuE transporters. The third proline import system of *B. subtilis* turned out to be GabP, the permease for γ -aminobutyrate (GABA) (42, 43). The simultaneous inactivation of the PutP, OpuE, and GabP transporters abolished the use of proline as a nutrient, but minor proline uptake activity remained and still allowed induction of *putBCP* expression in response to an external supply of proline.

MATERIALS AND METHODS

Chemicals. The amino acids L-proline and GABA, the toxic proline analogues AC and DHP, the chromogenic substrate *para*-nitrophenyl- α -D-glucopyranoside (PNPG) for the TreA enzyme (44), the ninhydrin reagent for the quantification of proline by a colorimetric assay (45), and the antibiotics chloramphenicol, kanamycin, erythromycin, tetracycline, lin-

comycin, and spectinomycin were all purchased from Sigma-Aldrich (Steinheim, Germany). The antibiotic zeocin was obtained from Invitrogen (Carlsbad, CA). Radiolabeled L-[U-¹⁴C]proline (269 mCi mmol⁻¹) and γ -[2,3-³H]aminobutyrate (35 Ci mmol⁻¹) were purchased from PerkinElmer LAS GmbH (Rodgau, German).

Bacterial strains, media, and growth conditions. The Escherichia coli K-12 strain DH5α (Invitrogen, Carlsbad, CA) was used for maintenance of recombinant plasmids. These strains were propagated either in Luria-Bertani (LB) liquid medium or on LB agar plates (46). For E. coli strains carrying plasmids encoding a beta-lactamase resistance gene, ampicillin was used at final concentrations of 100 µg ml⁻¹. The genetic properties of the B. subtilis wild-type laboratory strain JH642 (kindly provided by J. Hoch, Scripps Research Institute, La Jolla, CA) and its mutant derivatives used throughout this study are summarized in Tables 1 and 3. B. subtilis strains were routinely cultivated in Spizizen's minimal medium (SMM) with 0.5% (wt/vol) glucose as the carbon source, 15 mM (NH_4)₂SO₄ as the nitrogen source, and L-tryptophan (20 mg liter⁻¹) and L-phenylalanine $(18 \text{ mg liter}^{-1})$ to satisfy the auxotrophic requirements of strain JH642 (trpC2 pheA1) and its derivatives (Table 1). A solution of trace elements (47) was added to SMM to improve the growth of B. subtilis strains; the osmolarity of the growth medium was adjusted by adding NaCl to it from a 5 M stock solution. When the potential use of L-proline or GABA as the sole carbon source was tested, glucose was replaced by the addition of 33 mM L-proline or 41 mM GABA. In growth experiments with L-proline or GABA as the sole nitrogen source, the ammonium sulfate was left out from SMM and L-proline or GABA was added to a final concentration of 30 mM. All B. subtilis cultures were inoculated from exponentially growing precultures in prewarmed (37°C) SMM to an optical density at 578 nm (OD₅₇₈) of about 0.1, and the cultures were then subsequently propagated at 37°C in a shaking water bath set to 220 rpm. Growth of the strains was monitored by measuring their OD_{578} . The antibiotics chloramphenicol (5 μ g ml⁻¹), tetracycline (10 μ g ml⁻¹), erythromycin-lincomycin (0.4 μ g ml^{-1} and 15 µg ml⁻¹, respectively), and spectinomycin (100 µg ml⁻¹) were used for the selection of *B. subtilis* strains carrying chromosomal copies of gene disruption mutations with insertions of an antibiotic resistance cassette or of $\phi(putB'-treA)$ 1 cat reporter fusion constructs into the nonessential chromosomal *amyE* locus (Table 1).

The *B. subtilis* wild-type strain JH642 is able to use L-proline both as the sole nitrogen source and as the sole carbon source, but loss of the proline transporter PutP abrogated the use of proline as the sole carbon source when precultures grown with glucose as the carbon source were used to inoculate SMM (10). However, we observed that when strains carrying a *putP* gene disruption but intact *putBC* catabolic genes were pregrown in the presence of both glucose and L-proline, growth could occur with L-proline as the sole carbon source upon reinoculation. Accordingly, for such growth experiments, two precultures were prepared: the strains were initially grown to mid-exponential phase in SMM with glucose (28 mM) as the carbon source, and these precultures were then subsequently used to inoculate SMM that contained both glucose (14 mM) and proline (33 mM) as available carbon sources. When the second preculture had reached mid-exponential growth phase, the cells were carefully washed two times with SMM containing proline (33 mM) but no glucose. These cells were then used to inoculate the main culture that contained only proline (33 mM) as the carbon source. Cells treated in this way were able to grow even when they lacked the PutP proline import system (see Fig. S1 in the supplemental material).

Genetic constructions of bacterial strains. The routine manipulations of plasmid DNA, the amplification of selected regions of the B. subtilis genome via PCR, the isolation of chromosomal DNA from B. subtilis, and the transformation of B. subtilis strains with plasmid or chromosomal DNA were all carried out using standard procedures (47). To construct a deletion of a chosen gene in the genome of *B. subtilis*, a two-step PCR-based method (48) was used, in which the 5' and 3' regions of the gene of interest were first amplified by PCR from chromosomal DNA and connected in a second step by long-flanking-region PCR with a DNA fragment encoding an antibiotic resistance gene. The generated PCR products were then used for the transformation of B. subtilis strain JH642, and their integration into the chromosome via homologous recombination events was selected for by the addition of the appropriate antibiotic to the agar plates. The following strains were constructed in that way, and the primer pairs used for the long-flanking-region PCR are given in parentheses: ACB164 [Δ (*ywcA*::*ery*)1] (5'-CATGTTAAAA AAGATTTGGACAAAGGTTGG-3'/5'-TTTGTCATCAATCTGAAAGAA GCCTAAATC-3'), ACB165 [Δ (yodF::neo)1] (5'-TGTATCCAAAGCTTTC CGTTAACAATTGTA-3'/5'-TGTCTATTCAGAGATGAAAGACAACAAC AT-3'), and ACB274 [Δ (gabP::zeo)1] (5'-CTCTTGTTTAGCAGAAAAAAT AGATTCTCC-3'/5'-TTTTAAGCGTTTTTCCTTTGGGTCATGAAT-3'). The antibiotic resistance genes used for these gene disruption constructs were derived from plasmids pDG642 (ery), pDG783 (neo) (49), and p7Z6 (zeo) (50). The presence of the correct gene disruption constructs in the newly created mutant strains was verified by PCR using primers that flank the deleted genomic region; PCR products derived from genomic DNA of the B. subtilis wild-type strain JH642 were used for comparison. To combine different gene disruption mutations (Table 1), the recipient B. subtilis strain was transformed with chromosomal DNA derived from B. subtilis donor strains carrying a gene disruption linked to an antibiotic resistance marker, and the transformants were then selected on LB agar plates containing the appropriate antibiotic.

Sensitivity of *B. subtilis* strains to toxic proline analogues and selection for DHP-resistant mutants. To test the sensitivity of *B. subtilis* strains to the toxic proline analogues DHP and AC (10), cultures were grown in SMM in the absence and presence of 0.6 M NaCl until they reached exponential growth phase. Three hundred microliters of each culture was then plated on SMM or SMM agar plates with 0.6 M NaCl, and subsequently, a paper filter disk (diameter, 6 mm) soaked with 10 μ l of a 25-mg ml⁻¹ solution of DHP or AC was placed in the middle of each agar plate. The plates were incubated at 37°C for 24 h (SMM agar plates) or 48 h (SMM agar plates with 0.6 M NaCl), and the formation of growth inhibition zones around the filter disk was recorded by photography.

To isolate mutants resistant to DHP, strains with gene disruption mutations in *putP* and *opuE* were grown in SMM to an optical density (OD₅₇₈) of 1.5, and a 300- μ l aliquot was plated on SMM agar plates. A paper filter disk that was impregnated with 10 μ l of a 25-mg ml⁻¹ solution of DHP was placed in the center of each plate. After 2 days of incubation at To determine the frequency with which the DHP-resistant colonies appeared, three cultures of strain SMB12 (*opuE putP*) (Table 1) were grown in SMM with glucose as the carbon source and were then plated on SMM agar plates to determine the total number of viable cells and on SMM agar plates containing 50 μ g ml⁻¹ DHP (51). The plates were incubated for 2 days at 37°C; spontaneous DHP-resistant colonies arose at frequencies of about 6 × 10⁻⁶.

Genome resequencing, sequence assembly, and analysis. To identify the spontaneous mutation(s) conferring resistance to DHP, one of the isolated DHP-resistant strains (ACB214) and its parent strain (ACB199) (Table 1) were subjected to genome resequencing. For this purpose, genomic DNA was isolated using a DNeasy blood and tissue kit from Qiagen. Genome resequencing and mapping of the DNA reads on B. subtilis reference genomes (3, 52) were done by the Göttingen Genomics Laboratory (G2L). Shotgun libraries were generated using a Nextera XT DNA sample preparation kit according to the manufacturer's instructions, and DNA sequences were obtained with a Genome Analyzer IIx instrument (Illumina). The prepared libraries were sequenced in a 112-bp single-read indexed run on 1/4 of one channel, resulting in 8.3 million and 9.2 million reads for strains ACB199 and ACB214, respectively. All shotgun reads were mapped to the genomes of B. subtilis strains JH642 (52) and 168 (3) using the GS Reference Mapper software tool, v2.8 (454 Life Sciences). Sequence differences were considered single nucleotide polymorphisms (SNPs) when the total coverage depth exceeded 25 reads and the variant frequency was >90%.

The *gabP* gene from the chromosomal DNA of 15 DHP-resistant mutants was amplified by PCR in two DNA segments (primer AC182_GabP_Seq1_for [5'-CACGAATTTTCGACAAACTGTATATTT ATG-3'] and primer AC183_GabP_Seq1_rev [5'-GGGCTTTCTAAAAT ATTTGCTGAATTCC-3']; primer AC184_GabP_Seq2_for [5'-CATCTT CTCTTTTATGGGAACTGAAATC-3'] and primer AC185_GabP_Seq2_rev [5'-GTTAATGACAATAAGGACGTGATGGTTAC-3']), and the DNA sequence of the resulting PCR fragment was determined by Eurofins MWG Operon (Ebersberg, Germany). These DNA sequences were compared with the corresponding sequence of the genome region of the JH642 parent strain (52).

Measurements of cellular proline pools. The intracellular proline content of *B. subtilis* cells was determined by a colorimetric assay that detects L-proline as a colored proline-ninhydrin complex that can be quantified by measuring the absorption of the solution at 480 nm (45). For these assays, the *B. subtilis* cells were grown in SMM with 0.4 M NaCl; harvesting and processing of the cells, the details of the assay conditions, and the calculation of the intracellular volume have been described previously (15, 21, 53). Consistent with previous measurements (21), the proline pool of strain JH642 was found to be 95 ± 12 mM in cells that had been grown in SMM containing 0.4 M NaCl; the concentrations of proline pools detected in the 15 tested DHP-resistant mutant strains ranged from 79 ± 4 mM to 98 ± 5 mM. Hence, none of the isolated DHP-resistant mutants overproduced proline.

TreA enzyme activity assays. The expression of a chromosomal *putB-treA* reporter gene fusion was monitored by assaying the TreA [phospho- α -(1, 1)-glucosidase] reporter enzyme activity (44) using the chromogenic substrate PNPG (10). TreA enzyme activity is expressed in units per mg protein; protein concentrations were estimated from the OD₅₇₈ of the *B. subtilis* cell culture (46).

Transport assays with radiolabeled L-proline and γ -aminobutyrate. Uptake of γ -aminobutyric acid in *B. subtilis* was measured using γ -[2,3-³H]aminobutyrate (specific activity, 35 Ci mmol⁻¹) as the substrate. The cells were grown to mid-exponential phase (OD₅₇₈ = 0.3 to 0.4) in SMM without or with 0.6 M NaCl and were used immediately for the transport

TABLE 2 Use of proline and GABA as nutrients by B. subtilis^a

				Growth yield of cultures when the following nutrient was used as:					
	Relevant genotype ^{b}			C source			N source		
Strain	PutP	OpuE	GabP	Glucose	Proline	GABA	$\mathrm{NH_4}^+$	Proline	GABA
JH642	+	+	+	3.61	3.01	0.16	3.48	3.67	3.91
ACB275	+	-	_	3.89	2.83	0.10	4.13	3.04	0.24
ACB277	_	+	_	3.79	0.28	0.12	4.04	0.65	0.35
SMB12	_	-	+	3.71	0.30	0.14	3.80	3.29	3.68
ACB279	-	-	-	3.65	0.10	0.10	3.59	0.63	0.25

^{*a*} Cells of the *B. subtilis* wild-type strain JH642 and its mutant derivatives were grown in SMM with either 28 mM glucose, 33 mM proline, or 41 mM GABA as the carbon source in the presence of 30 mM ammonium as the nitrogen source. When GABA or proline was used as sole nitrogen source, it was present at a concentration of 30 mM. The cells were inoculated to an OD_{578} of 0.2 from precultures, and the growth yields were determined by measuring the OD_{578} of the cultures after 16 h of incubation at 37°C. The values given represent the means of two independent sets of experiments. ^{*b*} +, the presence of the wild-type gene; –, its absence.

assay. The uptake assay was started by the addition of γ -aminobutyrate (final assay concentration, 5 μ M to 1,000 μ M) spiked with γ -[2,3-³H]aminobutyrate (final assay concentration, 2.8 nM) in a total reaction volume of 2 ml. This resulted in a specific activity of γ -[2,3-³H]aminobutyrate of 20 mCi mmol⁻¹ at a final substrate concentration of 5 µM GABA and 0.1 mCi mmol⁻¹ at a final substrate concentration of 1,000 µM GABA in the individual transport assays. Samples (0.3 ml) were taken at various time points and filtered through 0.45-µm-pore-size filters (Schleicher & Schuell GmbH, Dassel, Germany). The cells were washed with 20 ml of isotonic minimal salts solution, and the radioactivity retained on the filters was determined in a scintillation counter (Perkin-Elmer, Rodgau, Germany). The uptake assay conditions for measuring the import of radiolabeled L-[U-14C]proline (269 mCi mmol⁻¹) (10) followed the above-described procedures, except that the following proline concentrations were used: for proline uptake assays involving wild-type strain JH642, 1 µM to 40 µM proline was used; in uptake assays involving strain SMB12 (putP opuE), 40 µM to 2,000 µM proline was used. The proline solution was spiked with L-[U-14C]proline (final concentration, 0.186 μ M) in a 2-ml assay volume. This resulted in a specific activity of L-[U-¹⁴C]proline of 42 mCi mmol⁻¹ at a final substrate concentration of $1~\mu M$ proline, 1.2 mCi mmol^{-1} at a final substrate concentration of 40 μ M proline, and 0.025 mCi mmmol⁻¹ at a final substrate concentration of 2,000 µM proline in the individual transport assays.

Database searches and alignments of amino acid sequences. Functional annotations of proteins of *B. subtilis* (3) were obtained from the SubtiWiki web server (http://subtiwiki.uni-goettingen.de/wiki/index.php /Main_Page) (54). The alignment of amino acid sequences of proteins was performed with ClustalW (55), and the amino acid sequence identities were calculated by using the EMBL-EBI online tool EMBOSS Needle (http://www.ebi.ac.uk/Tools/psa/).

RESULTS

A *putP opuE* double mutant strain can still use proline as a nutrient. The disruption of the PutP proline uptake system greatly impairs the growth of a *B. subtilis* strain with intact PutB and PutC proline catabolic enzymes in a minimal medium (SMM) with proline as the sole carbon and energy source (10). However, use of proline as a nitrogen source is still possible even in a strain with simultaneous defects in the PutP and OpuE proline transporters (10), but use of proline as a carbon source was impaired (Table 2). We then observed, however, that the preculturing of strains lacking either PutP or PutP and OpuE simultaneously in a medium containing both glucose and proline as carbon source (see Fig. S1 in

the supplemental material). The physiological mechanism underlying this difference in the use of proline as a nutrient is currently not firmly understood, but it probably reflects the different amounts of proline required to satisfy the cells' need for carbon and nitrogen under conditions where the main catabolic proline import system (PutP) is not functional. This growth pattern is definitely not the result of the accumulation of suppressor strains (53) allowing a bypass of the defect in the PutP transporter, since single colonies obtained from the proline-grown cultures showed the same growth behavior found for the originally inoculated $putBC^+$ -[$\Delta(putP::spc)$ 1] strain (data not shown). In conclusion, the data documented in Table 2 and in Fig. S1 and S3 in the supplemental material demonstrate that, in addition to the so far functionally characterized PutP and OpuE transporters (10, 16), at least one further proline import system must be active in B. subtilis.

A candidate approach to identify the third proline transporter of B. subtilis. To get a possible hint at the identity of the transport system that mediates proline import in a *putP opuE* double mutant strain, we used the amino acid sequence of PutP as the query sequence in a BLAST search (56) of the B. subtilis genome (3). This database search identified the YwcA and YodF membrane proteins as PutP-related proteins with overall amino acid sequence identities of 24% and 21%, respectively; their amino acid sequence identities to OpuE are 22% and 22%, respectively. A domain search using the SMART database (57) recognized in the YwcA and YodF proteins a solute:sodium symporter (SSS) domain, indicating that these predicted integral membrane proteins are members of the same transporter family to which the PutP and OpuE transporters also belong (10, 16). We constructed gene disruption mutations in the *ywcA* and *yodF* genes and combined the resulting mutant alleles with gene disruption mutations in the *putP* and *opuE* genes. The resulting strains, including the *putP opuE ywcA yodF* quadruple mutant ACB199, all retained sensitivity toward the toxic proline analog DHP (39) (Fig. 1A) and were able to use proline as a nutrient (see Fig. S2 in the supplemental material). We therefore concluded that the YwcA and YodF proteins do not contribute significantly to proline uptake by B. subtilis.

A genetic approach to identify the third proline transporter of *B. subtilis*. A *putP opuE* double mutant strain is still sensitive to the proline analogs 3,4-dehydro-DL-proline (DHP) and L-azetidine-2-carboxylic acid (AC) (Fig. 1B). These compounds can be taken up by bacteria via different types of proline import systems (36, 40, 41), mischarged onto proline-specific tRNAs, and incorporated, as a substitute for proline, into proteins that are then prone to misfolding (39). The resistance of bacteria to AC and DHP can be developed either through the overproduction of proline (58, 59), which dilutes the concentration of these toxic compounds within the overall cellular proline pool, or through the mutational inactivation of proline uptake systems (36, 40, 41), which prevents their uptake into the cell.

Mutant derivatives of the *putP opuE* strain SMB12 that are resistant to DHP were readily detectable in the growth-inhibition zone that developed around a filter disk soaked with this toxic proline analog. DHP-resistant mutants appeared at frequencies of about 6×10^{-6} on SMM agar plates that contained 50 µg ml⁻¹ DHP. We isolated 15 independent spontaneously DHP-resistant mutants from the growth-inhibition zone around a DHP-soaked filter disk (10 µl of a 25-mg ml⁻¹ solution was applied to the disk)

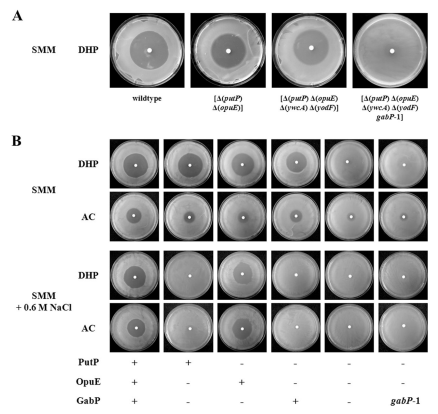


FIG 1 Sensitivity of the *B. subtilis* wild-type strain JH642 and its mutant derivatives to the toxic proline analogues AC and DHP. (A) Cells of the *B. subtilis* wild-type JH642 and its mutants derivatives SMB12 (*putP opuE*), ACB199 (*putP opuE ywcA yodF*), and ACB214 (*putP opuE ywcA yodF gabP1*) were tested for sensitivity to DHP on SMM agar plates. (B) Cells JH642 and its mutants derivatives ACB275 (*opuE gabP*), ACB277 (*putP gabP*), SMB12 (*putP opuE*), ACB279 (*putP opuE gabP*), and ACB214 (*putP opuE ywcA yodF gabP1*) were tested for sensitivity to DHP and AC either on SMM agar plates or on SMM agar plates that contained 0.6 M NaCl. The agar plates were incubated at 37°C for 24 h (cells grown on SMM agar plates) or 48 h (cells grown on SMM agar plates containing 0.6 M NaCl) and then recorded by photography.

with the aim of isolating mutant strains that were defective in proline uptake. Upon retesting, each of these mutants (e.g., ACB214) proved to be resistant to DHP at the concentration originally applied in the filter disk assay (Fig. 1A and B), and none of them overproduced proline (data not shown; see Materials and Methods for details). Hence, each of the 15 picked DHP-resistant strains exhibited the properties expected for a mutant with a defect in a proline uptake system. Indeed, when we tested these strains for the use of proline as either the sole nitrogen or carbon source, each of them was no longer able to use proline as a nutrient (see Fig. S2 and S3 in the supplemental material).

Genome resequencing provides a lead on the identity of the third proline transporter of *B. subtilis.* To get a lead on the type of mutation(s) underlying the DHP resistance phenotype and the defect in proline utilization, we subjected one of the isolated mutants (strain ACB214) and its DHP-sensitive parent (strain ACB199) (Table 1) to genome resequencing using an Illumina sequencing protocol. This resulted in 9.2 million and 8.3 million reads (about 110 bp each) for strains ACB214 and ACB199, respectively, with a minimum genome coverage of 25 reads per nucleotide.

Two mutations were found in the genome sequence of DHPresistant strain ACB214 in comparison to that of parent strain ACB199, which is a derivative of JH642 (52), a member of the domesticated 168 lineage of *B. subtilis* laboratory strains (3): (i) a

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consecutive 2-bp insertion (G-C-G-C) within codon 465 of *cheA*, the structural gene for the CheA signal transduction histidine kinase, a key component of the *B. subtilis* chemotactic system (60), and (ii) a single-base-pair insertion (a A-T base pair) within codon 387 of the *gabP* gene (*gabP1*) encoding the uptake system (GabP) for the nonproteinogenic amino acid γ -aminobutyrate (GABA) that can be used by *B. subtilis* as the sole nitrogen source but not as the sole carbon source (42, 43) (Table 2). Hence, the mutations found by genome resequencing of DHP-resistant strain ACB214 are predicted to cause the production of substantially truncated versions of CheA (467 amino acids) and GabP (423 amino acids), proteins that normally comprise 671 and 469 amino acids, respectively.

The occurrence of the mutation in *cheA* can be understood when one considers that *B. subtilis* displays chemotaxis not only toward L-proline but also toward toxic proline analogs (61). Inactivation of the CheA kinase would thus benefit *B. subtilis* cells by abrogating swimming toward higher concentrations of a poisonous compound (DHP) placed in the middle of the agar plate on a filter disk (Fig. 1A). However, the mutation in *cheA* does not provide a rational explanation for the inability of strain ACB214 to use proline as a nutrient (see Fig. S2 and S3 in the supplemental material). In this respect, the mutation in the *gabP* gene provided a very useful lead since GabP is a functionally characterized permease (42, 43); it is a member of the amino acid-polyamine-or-

 TABLE 3 Mutations in gabP conferring resistance to DHP in a putP

 opuE double mutant strain

Strain ^a	gabP allele	Mutation	Consequence
ACB214	gabP1	A1160	Frameshift in codon 387
ACB215	gabP2	TAT/TAA	Stop at codon 151
ACB254	gabP3	GGT/GTT	G301V
ACB255	gabP4	TTC/TCC	F341S
ACB256	gabP5	TGG/TAG	Stop at codon 100
ACB257	gabP4	TTC/TCC	F341S
ACB262	gabP6	GGT/AGT	G42S
ACB263	gabP7	A1156	Frameshift in codon 386
ACB264	gabP8	TGG/TAG	Stop at codon 118
ACB265	gabP9	GGG/GAG	G338E
ACB266	gabP10	A985	Frameshift in codon 329
ACB267	gabP11	TGG/TGA	Stop at codon 241
ACB268	gabP12	GGC/GAC	G33D
ACB269	gabP13	TAC/TAA	Stop at codon 49
ACB273	gabP14	GGA/AGA	G414R

^{*a*} All strains are derivatives of the *B. subtilis* wild-type laboratory strain JH642 (*trpC2 pheA1*), and each of them carries the $\Delta(putP::spc)1$ and $\Delta(opuE::tet)1$ gene disruption mutations. Strain ACB214 also possess the $\Delta(yodF::neo)1$ and $\Delta(ywcA::ery)1$ alleles, whereas in strain ACB215, only the *ywcA* gene [$\Delta(ywcA::ery)1$] is disrupted.

ganocation (APC) superfamily of transporters (TC accession number 2.A.3) (62). Indeed, the presence of the *gabP1* allele in a *putP opuE* strain background not only conferred DHP resistance (Fig. 1B) and abrogated utilization of proline as the sole nitrogen and carbon source (see Fig. S2 and S3 in the supplemental material) but also caused a strong defect in the use of GABA as the sole nitrogen source by *B. subtilis* (see Fig. S2 in the supplemental material).

If the loss of the GabP-catalyzed transporter activity was indeed causative for the development of the DHP-resistant phenotype and the concomitant inability of such strains to use proline as a nutrient in a *putP opuE* genetic background, then most, if not all, of the originally isolated DHP-resistant strains should carry mutations in gabP. To test this hypothesis, we sequenced the entire gabP gene from all of the remaining 14 independently isolated DHP-resistant mutant strains and found that in each of them the gabP gene had suffered a mutation (Table 3). In Fig. 2, we have projected the positions of these mutations onto a simple topological model for the GabP transporter. This model invokes the presence of 12 transmembrane-spanning regions and suggests that the N and C termini of the GabP protein face the cytoplasm (63), a topological arrangement consistently observed in several crystal structures of members of the APC superfamily (64-66). Two types of lesions were present in the gabP gene of the various DHPresistant strains: (i) stop codons (five isolates) and frameshifts (three isolates) that are predicted to generate truncated versions of the GabP permease (Table 3) and (ii) single amino acid substitutions (seven isolates) (Table 3) that might negatively affect the transporter activity of GabP as well. Indeed, regardless of the type of mutation present in gabP, each of the 15 DHP-resistant strains tested had lost the ability to efficiently use GABA as the sole nitrogen source (see Fig. S2 in the supplemental material).

With respect to the types of mutations recovered in *gabP*, those that lead to single amino acid substitutions in the GabP permease are the most interesting to consider further. In each of these seven GabP variants (the *gabP4* allele [F341S] was isolated twice; Table 3), the mutationally altered residues are predicted to be part of the

membrane-embedded core of the GabP transporter (Fig. 2). Strikingly, five of the six affected amino acids were glycine residues (G33, G42, G301, G338, G414); glycine is a type of residue that has a structural role in membrane-spanning helices, mediates helixhelix interactions in membrane proteins, and contributes to the formation of monomer-monomer interfaces (67, 68).

Targeted disruption of gabP confers DHP resistance, abolishes GABA utilization, and prevents use of proline as a nitrogen source in a *putP opuE* genetic background. The data presented above provided compelling evidence for a link between the loss of the GabP transporter function and the inability of the cells to use proline as a nutrient in a *putP opuE* genetic background (see Fig. S2 and S3 in the supplemental material). However, it seemed remotely possible that these two phenotypes were not causally linked. To avoid any possible bias that might be associated with the originally applied selection scheme for DHP resistance, we constructed a chromosomal gabP gene disruption mutation in which an antibiotic (zeocin) resistance cassette was used to replace the entire coding region of *gabP*. We then introduced the $\Delta(gabP)$: zeo)1 allele in various combinations into B. subtilis strains harboring *putP* and/or *opuE* gene disruption mutations and tested the resulting comprehensive set of mutant strains for their resistance to DHP and AC (Fig. 1B) and their ability to use either proline or GABA as nutrients (Table 2).

Proline utilization occurred as long as one of the PutP, OpuE, and GabP transporters was functional; it was reduced to background levels only when all three transporters were nonoperational (Table 2). This data set thus unambiguously identifies the GabP transporter as the third proline uptake system operating in B. subtilis. Loss of the ability to use proline as a nutrient required the simultaneous inactivation of the PutP, OpuE, and GabP transporters, but use of GABA as the sole nitrogen source was abolished only when the GabP transporter was nonfunctional. The simultaneous loss of the PutP or OpuE transporters did not negatively affect GABA utilization (Table 2). This also firmly excluded the possibility that the cheA frameshift mutation present in the strain (ACB214) whose genome we initially resequenced was required in conferring resistance to toxic proline analogs (Fig. 1A and B) and causing a loss of proline utilization (see Fig. S2 in the supplemental material).

As noted previously (10), the sensitivity of *B. subtilis* strains to

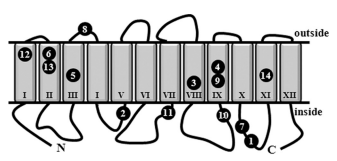


FIG 2 Model for the topological organization of the GabP transporter and position of mutations causing DHP resistance. The predicted 12 transmembrane domains of the GabP transporter with its amino-terminal (N) and carboxy-terminal (C) ends facing the cytoplasm are shown. The indicated numbers refer to the *gabP* alleles whose properties are summarized in Table 3. These *gabP* mutants lead to either truncated variants of the GabP permease (alleles 1, 2, 5, 7, 8, 10, 11, and 13) or single amino acid substitutions in the GabP permease (alleles 3, 4, 6, 9, 12, and 14).

DHP and AC is influenced by the salinity of the growth medium and by which proline transport system is operational. Our isogenic set of strains expressing only one of the PutP, OpuE, and GabP proline transporters or lacking all of them allowed us to assess the individual contribution of these transporters to DHP and AC sensitivity. For these assays, we used SMM agar plates without and with additional NaCl (0.6 M). As shown in Fig. 1B, in a strain possessing only PutP, sensitivity to DHP and AC was abolished when the salinity of the growth medium was increased, whereas in a strain possessing only OpuE, not only was its sensitivity to AC maintained, but its sensitivity was actually increased. The pattern of sensitivity/resistance to DHP and AC of a strain possessing only GabP mimicked that of a strain expressing only PutP (Fig. 1B). There is a good correlation between the pattern of ¹⁴C]proline uptake in cells (cultivated in the absence or presence of increased salinity) and the extent of growth inhibition exerted by DHP and AC on agar plates lacking or containing additional amounts of NaCl (10). Based on the data shown in Fig. 1B, we surmise that the efficiency of GabP-mediated proline uptake is negatively affected in high-salinity growth medium (see below).

Kinetic parameters of proline transport via GabP relative to those of GABA transport. The level of gabP transcription is responsive to the nitrogen source available in the growth medium, and it is subjected to genetic control by the key nitrogen regulatory protein of B. subtilis, TnrA (42, 69), and the globally acting CodY protein (42, 70). As a consequence, GabP-mediated GABA uptake is strongly stimulated when B. subtilis cells are propagated with a growth-limiting nitrogen source such as glutamate (42). We monitored the uptake of radiolabeled GABA under conditions that maximize gabP expression in cells of the wild-type strain JH642 and of a strain (SMB12) that was proficient in the GabP permease but lacked the PutP and OpuE transporters. For these experiments, the cells were grown in either SMM or SMM containing 0.6 M NaCl; both media contained glutamate as the sole nitrogen source. In the uptake assays, we used radiolabeled GABA at a final substrate concentration of 200 µM (Fig. 3A). Both strains showed the same GABA uptake rate [about 132 nmol GABA (min mg $(protein)^{-1}$]. This uptake rate was reduced 2.6- to 3-fold when the strains were cultivated in SMM with increased osmolarity (Fig. 3A), indicating that the activity of the GabP permease is impaired by high salinity. This property nicely explains the reduced sensitivity of a GabP⁺ PutP⁻ OpuE⁻ strain to the toxic proline analogs DHP and AC on high-salinity agar plates (Fig. 1B). The γ -[2,3-³H]aminobutyrate import activity in the GabP-deficient strain ACB279 was reduced to background levels when a substrate concentration of 200 µM was used in the uptake assay, but minor uptake activity was noticeable when the substrate concentration was increased to 700 µM (Fig. 3A). The latter finding is consistent with previous data showing the dominant role of GabP in GABA uptake and the existence of an additional, but very minor, uptake route for this compound in *B. subtilis* (42, 71).

Since the details of the kinetic parameters of the GabP permease in its authentic host strain, *B. subtilis*, have not been reported, we determined them in cells of strain SMB12 (GabP⁺ PutP⁻ OpuE⁻) that were grown in a minimal medium (SMM) with glucose as the carbon source and glutamate as the sole nitrogen source. γ -[2,3-³H]aminobutyrate uptake exhibited Michaelis-Menten kinetics (Fig. 3B) and a K_m of 77 \pm 17 μ M; from the kinetic data, a V_{max} of 310 \pm 21 nmol (min mg protein)⁻¹ was calculated. Hence, GabP is a transport system with a good affinity

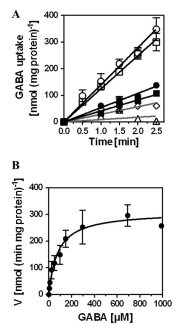


FIG 3 Uptake of radiolabeled GABA by *B. subtilis* and kinetic parameters for its GabP-mediated import. Cells were grown in SMM with glutamate as the sole nitrogen source. After the cultures reached an OD_{578} of 0.4, 2-ml aliquots were withdrawn to determine the uptake of γ -[2,3-³H]aminobutyrate. (A) *B. subtilis* wild-type JH642 (circles) and its derivative, SMB12 (GabP⁺ OpuE⁻ PutP⁻) (squares), were grown in minimal medium either in the absence (open symbols) or in the presence (closed symbols) of 0.6 M NaCl. GABA uptake was measured at a final concentration of 200 μ M. Uptake of radiolabeled GABA in the triple mutant strain ACB279 (GabP⁻ OpuE⁻ PutP⁻) was determined at final substrate concentrations of either 200 μ M (triangles) or 700 μ M (diamonds). Each transport assay was performed in at least two biological replicates. (B) Michaelis-Menten kinetics of GABA uptake in the GabP⁺ strain SMB12. Shown are the means and standard deviations of three biological replicates.

for GABA, and it exhibits a very substantial transport capacity for this nonproteinogenic amino acid. This conclusion is in full agreement with previously reported data on the stimulation of the uptake rates for GABA by *B. subtilis* cells grown in the presence of a poor nitrogen source and transport measurements with recombinant *E. coli* strains expressing the cloned *B. subtilis gabP* gene under the control of the *lac* promoter (42, 43).

To assess the ability of the GabP permease to mediate proline import, we determined under the same growth conditions that we used to measure GABA uptake the kinetic parameters for L-[¹⁴C]proline transport. For these experiments we used a strain (JH642) in which the three proline transporters PutP, OpuE, and GabP were intact and then compared its proline uptake activity with that of a strain in which only the GabP permease was functional (strain SMB12). Proline import into strain JH642 occurred with a K_m of 6.8 \pm 0.7 μ M and a V_{max} of 118 \pm 4 nmol (min mg protein)⁻¹ (Fig. 4A), demonstrating that a *B. subtilis* wild-type strain can readily acquire proline from scarce environmental resources, such as root exudates and plant-mediated deposits in the rhizosphere (2, 25). When proline import was measured at a final substrate concentration of 40 µM, a substrate concentration at which no proline uptake activity was previously detectable in a putP opuE double mutant strain (SMB12) grown in SMM with glucose as the carbon source and ammonium as the nitrogen source (10), L-[¹⁴C]proline import was reduced about 3-fold in

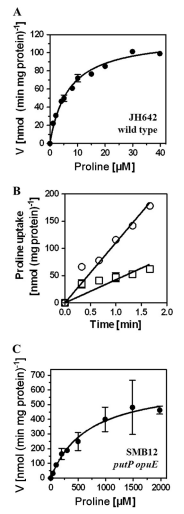


FIG 4 Uptake of radiolabeled proline by *B. subtilis* and kinetic parameters for its GabP-mediated import. Uptake of L-[U-¹⁴C]proline was determined in 2-ml aliquots of cultures that were grown in SMM to the early exponential growth phase (OD_{578} , 0.3). (A) Michaelis-Menten kinetics of proline uptake in cultures of the wild-type strain JH642. Shown are the means and standard deviations of three biological replicates. (B) Proline uptake rates in *B. subtilis* wild-type JH642 (circles) and its derivative SMB12 (GabP⁺ OpuE⁻ PutP⁻) (squares) were measured at a final proline concentration of 40 μ M. Each transport assay was performed in three biological replicates. (C) Michaelis-Menten kinetics of proline uptake in cultures of the GabP⁺ strain SMB12 (*putP opuE*). Shown are the means and standard deviations of three biological replicates.

strain SMB12 in comparison with that in strain JH642 (Fig. 4B). We then measured the GabP-mediated import of proline over a range of substrate concentrations, and from these data (Fig. 4C) a K_m of 700 \pm 330 μ M was estimated. We measured the rates of L-[¹⁴C]proline uptake between 40 μ M and 2 mM, but under these assay conditions, the uptake kinetics did not reach saturation. Hence, it is safe to say that GabP possesses only a rather modest affinity for proline with a K_m value of about 1 mM. In comparison, the K_m values of the PutP and OpuE transporters are 8 \pm 2 μ M and 12 \pm 1 μ M, respectively, in *B. subtilis* cells cultivated with glucose as the carbon source and ammonium as the nitrogen source in the absence of salt stress (10).

Proline can still enter the cells of a mutant strain lacking the PutP, OpuE, and GabP transporters. We observed that a strain that did not possess the PutP, OpuE, and GabP transporters was still not completely resistant to DHP and AC. A diffuse zone of growth impairment was visible around the filter disk impregnated with DHP, and a small but distinct growth inhibition zone was noticeable around the AC-containing filter disk (Fig. 1B). Hence, it seemed possible that there is another proline uptake route present in *B. subtilis* beyond that afforded by the PutP, OpuE, and GabP transport systems. The weak but noticeable growth of such triple mutant strains with proline as a nutrient (Table 2; see Fig. S2 in the supplemental material) fostered this suspicion.

Enhanced transcription of *putBCP* expression can be triggered by very low concentrations (e.g., 25 µM) of proline in a growth medium that contains glucose as the carbon source and ammonium as the nitrogen source (10). A single-copy *putB-treA* operon fusion is thus a very sensitive tool to assess whether proline can enter a given B. subtilis strain. We introduced such a fusion into a set of strains that expressed only one of the PutP, OpuE, and GabP transporters and then monitored the expression of the *putB-treA* reporter in response to the presence of both a low (25 μ M) and a high (5 mM) concentration of proline in the growth medium. We conducted these experiments with cells that were grown either in SMM or in SMM that contained 250 mM NaCl, a salt concentration that already allows good induction of *opuE* expression (23) but that does not strongly impair proline uptake via PutP (10) or GabP (see above). As summarized in Table 4, putB-treA gene expression was inducible by proline, regardless of which of the so far functionally characterized proline transport systems was operational. This was true even in a *putP opuE gabP* triple mutant strain (Table 4), demonstrating that at least one additional proline uptake system must be operational in B. subtilis. However, this trans-

TABLE 4 Induction of <i>p</i>	utB-treA expression	in response to extern	ally provided	proline imported via	different proline transport systems ^a

				TreA activity (U mg protein ⁻¹) in response to the indicated proline concn						
	Relevant	levant genotype ^b		0 M NaCl			0.25 M NaCl			
Strain	PutP	OpuE	GabP	0 M	25 μΜ	5 mM	0 M	25 μΜ	5 mM	
SMB10	+	+	+	19 ± 1	50 ± 2	168 ± 22	20 ± 2	52 ± 4	290 ± 13	
ACB276	+	—	_	15 ± 2	47 ± 5	141 ± 3	19 ± 2	45 ± 3	154 ± 5	
ACB278	_	+	_	19 ± 4	41 ± 2	90 ± 4	19 ± 3	45 ± 5	212 ± 10	
ACB288	—	—	+	20 ± 5	28 ± 6	140 ± 5	19 ± 3	30 ± 4	157 ± 18	
ACB292	—	—	_	20 ± 1	20 ± 3	87 ± 3	22 ± 2	28 ± 1	101 ± 4	

^{*a*} All strains carry the $\phi(putB-treA)$ reporter gene fusion integrated in the *amyE* site on the *B. subtilis* chromosome. Cultures were grown to early exponential growth phase (OD₅₇₈) about 0.5), and proline was then added to the growth medium at a final concentration of either 25 μ M or 5 mM. The cells were then propagated for another 60 min and subsequently harvested for TreA reporter enzyme assay. The values for the TreA activity given represent those for two independently grown cultures, and for each sample analyzed, the TreA activity was determined twice.

^{*b*} +, the presence of the wild-type gene; –, its absence.

port system(s) allowed proline uptake only in quantities that were insufficient to sustain efficient growth (Table 2; see Fig. S2 and S3 in the supplemental material), and its affinity must be low, since the induction of *putB-treA* transcription no longer occurred in response to proline at a concentration of 25 μ M in a *putP opuE* gabP triple mutant strain, but induction occurred at higher substrate concentrations (5 mM) (Table 4).

DISCUSSION

Although the PutP and OpuE proline transporters are closely related to each other with respect to both their amino acid sequences (10, 16) and the mechanism by which they import their common substrate in cotransport with sodium ions (31, 33, 34, 37), they have clearly different cellular functions in *B. subtilis*. Their primary physiological roles can be discerned from the distinct transcriptional profile of their structural genes (10, 16, 23, 28, 29) and the different abilities of the PutP and OpuE proteins to withstand the inhibiting effects of high salinity on their transport activity (10). These features associate PutP with the proline catabolic system of *B. subtilis* (10, 28, 29) and OpuE with the cell's osmostress response systems (16, 22, 72, 73). Nevertheless, it is worth noting that our data now show not only that the OpuE transporter imports proline when it is used as an osmoprotectant (15, 16) but also that it can do so when *B. subtilis* uses it as a nutrient (see Fig. S1 in the supplemental material).

Here we now identify the third proline transporter operating in B. subtilis, the GabP permease. GabP has previously been physiologically and biochemically characterized as an uptake system for GABA (42, 43), a nonproteinogenic amino acid that can be found in the rhizosphere (2) primarily due to the biosynthetic activities of plants (74). The soil bacterium B. subtilis can catabolize GABA to succinate and use it as the sole nitrogen source (42, 71). We have assessed the kinetic parameters of the GabP-mediated uptake of GABA and proline under growth conditions that should maximize gabP expression (42), and these data characterize GabP as a high-affinity uptake system for GABA (K_m , 77 \pm 17 μ M) that possesses a very substantial transport capacity [V_{max} , 310 \pm 21 nmol (min mg protein)⁻¹] for this substrate (Fig. 3B). In contrast, its affinity for proline is rather low, with an estimated K_m of close to 1 mM (Fig. 4C). Despite this limited affinity, enough proline can nevertheless be imported via GabP to serve as the sole nitrogen, energy, and carbon source for B. subtilis (Table 2; see Fig. S2 and S3 in the supplemental material).

The *B. subtilis* GabP transporter not only can import GABA (42) and proline (this study) but also can serve as an effective uptake system for β -alanine. The K_m value exhibited by GabP for β -alanine ($K_m = 9.6 \pm 5.3 \mu$ M) even exceeded that for GABA ($K_m = 37 \pm 13.2 \mu$ M) in a recombinant *E. coli* strain expressing the *B. subtilis gabP* gene under the control of the *lac* promoter (43). Hence, the ligand-binding site of the GabP permease can accommodate both linear molecules (GABA and β -alanine) and also, with considerably reduced efficiency, the cyclic structure of proline. Detailed insights into the molecular determinants for the high-affinity binding of both GABA and proline by the same transporter protein were recently provided through the crystallographic analysis of a periplasmic ligand-binding protein that operates in conjunction with an ABC transport system in the plant-associated bacterium *Agrobacterium tumefaciens* (75).

In contrast to PutP and OpuE, which are both members of the solute:sodium symporter family (SSS; TC accession number

2.A.21) (30), GabP belongs to the amino acid-polyamine-organocation (APC) superfamily (TC accession number 2.A.3) (62). Its amino acid sequence is only distantly related to the amino acid sequences of the PutP and OpuE proteins, with sequence identities of 19% and 17%, respectively. APC-type transporters catalyze the uniport, antiport, and symport of a broad range of substrates across cell membranes (64-66). GabP is a substrate:H⁺-symporter (TC accession number 2.A.3.1.5), and it is categorized as a member of the amino acid transporter subfamily of the APC superfamily (30). Interestingly, this subfamily also comprises ProY, a proline transporter of S. Typhimurium, whose structural gene is normally silent in laboratory strains (76). Notable is also that one of the three GABA transporters operating in Saccharomyces cerevisiae belongs to the same transporter family and functions as a permease (PUT4) that possesses similar affinities for both GABA and proline (77). Furthermore, members of the plant AtProT family of compatible uptake systems can import not only glycine betaine but also both proline and GABA (78). Hence, uptake systems that can import both GABA and proline are not uncommon.

In addition to defining GabP as the third proline transporter of *B. subtilis*, our data also reveal the existence of yet another proline uptake route in this soil bacterium. Efficient use of proline as a nutrient was abrogated in strains with simultaneous defects in the PutP, OpuE, and GabP permeases (Table 2; see Fig. S2 and S3 in the supplemental material), indicating that this fourth proline import route is of limited physiological relevance as far as the use of proline as a nutrient (10) and as an osmoprotectant (15, 16) is concerned. Nevertheless, enough proline can enter the *B. subtilis* cell via this route to induce enhanced expression of a proline-responsive (10, 28, 29) put-treA reporter fusion (Table 4).

The *gabP*, *putP*, and *opuE* proline transporter genes each possess distinct regulatory profiles, but they also share common genetic determinants. Transcription of *opuE* is strongly induced in response to both sudden and sustained increases in the external osmolarity, a process that is mediated by SigA- and SigB-type promoters (16, 23, 24). Data derived from transcriptional profiling experiments suggest that it is also under the positive control of the central carbon metabolism regulatory protein CcpA (79). Transcription of the *putBCP* operon is activated by the proline-responsive PutR regulator (10, 28, 29), whereas that of the *gabP* gene is enhanced through the stimulating activity of TnrA (42, 69), the central regulator of nitrogen metabolism in *B. subtilis* (12, 26, 27). Notably, transcription of *gabP* is not under the control of GabR, a regulatory protein that controls the expression of the *gabTD* GABA catabolic operon (71, 80).

A common determinant for the transcriptional control of the *putP*, *opuE*, and *gabP* genes is their regulation via the globally acting CodY protein, whose DNA-binding activity is sensitively modulated through the cellular pools of branched-chain amino acids and through GTP (70, 81, 82). CodY acts as a repressor for the promoter directing transcription of the *putBCP* operon (28), and it also acts as a negative regulator both for the *gabP* transporter gene (42) and for the GABA catabolic *gabTD* operon (70). CodY also binds to the *opuE* regulatory region (70), but its mode of action for this gene still needs to be studied. One of the cues that CodY avidly responds to is the availability of amino acids in the growth medium (81, 83). Hence, the common CodY-mediated transcriptional control of *putBCP*, *opuE*, and *gabP* allows *B. sub-tilis* to integrate its three major proline importers into its overarching nutrient utilization systems (26), while maintaining situa-

tion-specific control over their expression through regulatory circuits tailored to the individual genes and gene clusters.

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REFERENCES

- Earl AM, Losick R, Kolter R. 2008. Ecology and genomics of *Bacillus subtilis*. Trends Microbiol. 16:269–275. http://dx.doi.org/10.1016/j.tim .2008.03.004.
- Moe LA. 2013. Amino acids in the rhizosphere: from plants to microbes. Am. J. Bot. 100:1692–1705. http://dx.doi.org/10.3732/ajb.1300033.
- Barbe V, Cruveiller S, Kunst F, Lenoble P, Meurice G, Sekowska A, Vallenet D, Wang T, Moszer I, Medigue C, Danchin A. 2009. From a consortium sequence to a unified sequence: the *Bacillus subtilis* 168 reference genome a decade later. Microbiology 155:1758–1775. http://dx.doi .org/10.1099/mic.0.027839-0.
- Belda E, Sekowska A, Le Fevre F, Morgat A, Mornico D, Ouzounis C, Vallenet D, Medigue C, Danchin A. 2013. An updated metabolic view of the *Bacillus subtilis* 168 genome. Microbiology 159:757–770. http://dx.doi .org/10.1099/mic.0.064691-0.
- Beauregard PB, Chai Y, Vlamakis H, Losick R, Kolter R. 2013. Bacillus subtilis biofilm induction by plant polysaccharides. Proc. Natl. Acad. Sci. U. S. A. 110:E1621–E1630. http://dx.doi.org/10.1073/pnas.1218984110.
- Saier MH, Jr, Goldman SR, Maile RR, Moreno D, Weyler MSW, Yang N, Pauslsen IT. 2002. Overall transport capabilities of *Bacillus subtilis*, p 113–128. *In* Sonenshein AL, Hoch JA, Losick R (ed), *Bacillus subtilis* and its closes relatives: from genes to cells. ASM Press, Washington, DC.
- 7. Boch J, Kempf B, Bremer E. 1994. Osmoregulation in *Bacillus subtilis*: synthesis of the osmoprotectant glycine betaine from exogenously provided choline. J. Bacteriol. **176**:5364–5371.
- Hoffmann T, Bremer E. 2011. Protection of *Bacillus subtilis* against cold stress via compatible-solute acquisition. J. Bacteriol. 193:1552–1562. http: //dx.doi.org/10.1128/JB.01319-10.
- 9. Holtmann G, Bremer E. 2004. Thermoprotection of *Bacillus subtilis* by exogenously provided glycine betaine and structurally related compatible solutes: involvement of Opu transporters. J. Bacteriol. **186**:1683–1693. http://dx.doi.org/10.1128/JB.186.6.1683-1693.2004.
- Moses S, Sinner T, Zaprasis A, Stöveken N, Hoffmann T, Belitsky BR, Sonenshein AL, Bremer E. 2012. Proline utilization by *Bacillus subtilis*: uptake and catabolism. J. Bacteriol. 194:745–758. http://dx.doi.org/10 .1128/JB.06380-11.
- Atkinson MR, Wray LV, Jr, Fisher SH. 1990. Regulation of histidine and proline degradation enzymes by amino acid availability in *Bacillus subtilis*. J. Bacteriol. 172:4758–4765.
- 12. Fisher SH. 1993. Utilization of amino acids and other nitrogencontaining compounds, p 221–235. *In* Sonenshein AL, Hoch JA, Losick R (ed), *Bacillus subtilis* and other Gram-positive bacteria. ASM Press, Washington, DC.
- Whatmore AM, Chudek JA, Reed RH. 1990. The effects of osmotic upshock on the intracellular solute pools of *Bacillus subtilis*. J. Gen. Microbiol. 136: 2527–2535. http://dx.doi.org/10.1099/00221287-136-12-2527.

- Brill J, Hoffmann T, Bleisteiner M, Bremer E. 2011. Osmotically controlled synthesis of the compatible solute proline is critical for cellular defense of *Bacillus subtilis* against high osmolarity. J. Bacteriol. 193:5335– 5346. http://dx.doi.org/10.1128/JB.05490-11.
- Zaprasis A, Brill J, Thüring M, Wünsche G, Heun M, Barzantny H, Hoffmann T, Bremer E. 2013. Osmoprotection of *Bacillus subtilis* through import and proteolysis of proline-containing peptides. Appl. Environ. Microbiol. 79:576–587. http://dx.doi.org/10.1128/AEM.01934-12.
- von Blohn C, Kempf B, Kappes RM, Bremer E. 1997. Osmostress response in *Bacillus subtilis*: characterization of a proline uptake system (OpuE) regulated by high osmolarity and the alternative transcription factor sigma B. Mol. Microbiol. 25:175–187. http://dx.doi.org/10.1046/j .1365-2958.1997.4441809.x.
- Belitsky BR, Brill J, Bremer E, Sonenshein AL. 2001. Multiple genes for the last step of proline biosynthesis in *Bacillus subtilis*. J. Bacteriol. 183: 4389–4392. http://dx.doi.org/10.1128/JB.183.14.4389-4392.2001.
- Brill J, Hoffmann T, Putzer H, Bremer E. 2011. T-box-mediated control of the anabolic proline biosynthetic genes of *Bacillus subtilis*. Microbiology 157:977–987. http://dx.doi.org/10.1099/mic.0.047357-0.
- Csonka LN, Leisinger T. 8 August 2007. Chapter 34.6.1.4, Biosynthesis of proline. *In* Böck A, Curtis R, III, Kaper JB, Karp PD, Neidhardt FC, Nyström T, Slauch JM, Squires CL, Ussery D (ed), EcoSal—*Escherichia coli* and *Salmonella*: cellular and molecular biology. ASM Press, Washington, DC. http://dx.doi.org/10.1128/ecosalplus.3.6.1.4.
- Kempf B, Bremer E. 1998. Uptake and synthesis of compatible solutes as microbial stress responses to high osmolality environments. Arch. Microbiol. 170:319–330. http://dx.doi.org/10.1007/s002030050649.
- Hoffmann T, Wensing A, Brosius M, Steil L, Völker U, Bremer E. 2013. Osmotic control of *opuA* expression in *Bacillus subtilis* and its modulation in response to intracellular glycine betaine and proline pools. J. Bacteriol. 195:510–522. http://dx.doi.org/10.1128/JB.01505-12.
- Hoffmann T, von Blohn C, Stanek A, Moses S, Barzantny S, Bremer E. 2012. Synthesis, release, and recapture of the compatible solute proline by osmotically stressed *Bacillus subtilis* cells. Appl. Environ. Microbiol. 78: 5753–5762. http://dx.doi.org/10.1128/AEM.01040-12.
- Spiegelhalter F, Bremer E. 1998. Osmoregulation of the *opuE* proline transport gene from *Bacillus subtilis*: contributions of the sigma A- and sigma B-dependent stress-responsive promoters. Mol. Microbiol. 29:285– 296. http://dx.doi.org/10.1046/j.1365-2958.1998.00929.x.
- Young JW, Locke JC, Elowitz MB. 2013. Rate of environmental change determines stress response specificity. Proc. Natl. Acad. Sci. U. S. A. 110: 4140–4145. http://dx.doi.org/10.1073/pnas.1213060110.
- Vilchez S, Molina L, Ramos C, Ramos JL. 2000. Proline catabolism by *Pseudomonas putida*: cloning, characterization, and expression of the *put* genes in the presence of root exudates. J. Bacteriol. 182:91–99. http://dx .doi.org/10.1128/JB.182.1.91-99.2000.
- Sonenshein AL. 2007. Control of key metabolic intersections in *Bacillus subtilis*. Nat. Rev. Microbiol. 5:917–927. http://dx.doi.org/10.1038/nrmicro1772.
- 27. Gunka K, Commichau FM. 2012. Control of glutamate homeostasis in *Bacillus subtilis*: a complex interplay between ammonium assimilation, glutamate biosynthesis and degradation. Mol. Microbiol. 85:213–224. http://dx.doi.org/10.1111/j.1365-2958.2012.08105.x.
- Belitsky BR. 2011. Indirect repression by *Bacillus subtilis* CodY via displacement of the activator of the proline utilization operon. J. Mol. Biol. 413:321–336. http://dx.doi.org/10.1016/j.jmb.2011.08.003.
- Huang SC, Lin TH, Shaw GC. 2011. PrcR, a PucR-type transcriptional activator, is essential for proline utilization and mediates prolineresponsive expression of the proline utilization operon *putBCP* in *Bacillus subtilis*. Microbiology 157:3370–3377. http://dx.doi.org/10.1099/mic.0 .054197-0.
- Saier MH, Jr, Yen MR, Noto K, Tamang DG, Elkan C. 2009. The transporter classification database: recent advances. Nucleic Acids Res. 37:D274–D278. http://dx.doi.org/10.1093/nar/gkn862.
- Jung H, Hilger D, Raba M. 2012. The Na⁺/L-proline transporter PutP. Front. Biosci. 17:745–759. http://dx.doi.org/10.2741/3955.
- Raba M, Baumgartner T, Hilger D, Klempahn K, Hartel T, Jung K, Jung H. 2008. Function of transmembrane domain IX in the Na+/proline transporter PutP. J. Mol. Biol. 382:884–893. http://dx.doi.org/10.1016/j .jmb.2008.07.070.
- Mazier S, Quick M, Shi L. 2011. Conserved tyrosine in the first transmembrane segment of solute:sodium symporters is involved in Na⁺coupled substrate co-transport. J. Biol. Chem. 286:29347–29355. http://dx .doi.org/10.1074/jbc.M111.263327.

- Olkhova E, Raba M, Bracher S, Hilger D, Jung H. 2011. Homology model of the Na⁺/proline transporter PutP of *Escherichia coli* and its functional implications. J. Mol. Biol. 406:59–74. http://dx.doi.org/10.1016/j .jmb.2010.11.045.
- 35. Wood JM. 1988. Proline porters effect the utilization of proline as nutrient or osmoprotectant for bacteria. J. Membr. Biol. 106:183–202. http://dx .doi.org/10.1007/BF01872157.
- Csonka LN. 1982. A third L-proline permease in Salmonella typhimurium which functions in media of elevated osmotic strength. J. Bacteriol. 151: 1433–1443.
- 37. Cairney J, Higgins CF, Booth IR. 1984. Proline uptake through the major transport system of *Salmonella typhimurium* is coupled to sodium ions. J. Bacteriol. **160**:22–27.
- Hahne H, Mäder U, Otto A, Bonn F, Steil L, Bremer E, Hecker M, Becher D. 2010. A comprehensive proteomics and transcriptomics analysis of *Bacillus subtilis* salt stress adaptation. J. Bacteriol. 192:870–882. http://dx.doi.org/10.1128/JB.01106-09.
- 39. Bach TM, Takagi H. 2013. Properties, metabolisms, and applications of L-proline analogues. Appl. Microbiol. Biotechnol. 97:6623–6634. http://dx.doi.org/10.1007/s00253-013-5022-7.
- Wood JM. 1981. Genetics of L-proline utilization in *Escherichia coli*. J. Bacteriol. 146:895–901.
- May G, Faatz E, Villarejo M, Bremer E. 1986. Binding protein dependent transport of glycine betaine and its osmotic regulation in *Escherichia coli* K12. Mol. Gen. Genet. 205:225–233. http://dx.doi.org/10.1007/BF00430432.
- 42. Ferson AE, Wray LV, Jr, Fisher SH. 1996. Expression of the *Bacillus subtilis gabP* gene is regulated independently in response to nitrogen and amino acid availability. Mol. Microbiol. 22:693–701. http://dx.doi.org/10.1046/j.1365-2958.1996.d01-1720.x.
- Brechtel CE, King SC. 1998. 4-Aminobutyrate (GABA) transporters from the amine-polyamine-choline superfamily: substrate specificity and ligand recognition profile of the 4-aminobutyrate permease from *Bacillus subtilis*. Biochem. J. 333:565–571.
- Gotsche S, Dahl MK. 1995. Purification and characterization of the phospho-alpha-(1,1)-glucosidase (TreA) of *Bacillus subtilis* 168. J. Bacteriol. 177:2721–2726.
- Bates SL, Waldren RP, Teare ID. 1973. Rapid determination of free proline for water-stress studies. Plant Soil 39:205–207. http://dx.doi.org /10.1007/BF00018060.
- Miller JH. 1972. Experiments in molecular genetics. Cold Spring Harbor Laboratory, Cold Spring Harbor, NY.
- Harwood CR, Archibald AR. 1990. Growth, maintenance and general techniques, p 1–26. *In* Harwood CR, Cutting SM (ed), Molecular biological methods for *Bacillus*. John Wiley & Sons, Chichester, United Kingdom.
- 48. Kuwayama H, Obara S, Morio T, Katoh M, Urushihara H, Tanaka Y. 2002. PCR-mediated generation of a gene disruption construct without the use of DNA ligase and plasmid vectors. Nucleic Acids Res. 30:E2. http: //dx.doi.org/10.1093/nar/30.2.e2.
- Guerout-Fleury AM, Shazand K, Frandsen N, Stragier P. 1995. Antibiotic-resistance cassettes for *Bacillus subtilis*. Gene 167:335–336. http://dx .doi.org/10.1016/0378-1119(95)00652-4.
- Yan X, Yu HJ, Hong Q, Li SP. 2008. Cre/lox system and PCR-based genome engineering in *Bacillus subtilis*. Appl. Environ. Microbiol. 74: 5556–5562. http://dx.doi.org/10.1128/AEM.01156-08.
- Stalmach ME, Grothe S, Wood JM. 1983. Two proline porters in *Escherichia coli* K-12. J. Bacteriol. 156:481–486.
- Srivatsan A, Han Y, Peng J, Tehranchi AK, Gibbs R, Wang JD, Chen R. 2008. High-precision, whole-genome sequencing of laboratory strains facilitates genetic studies. PLoS Genet. 4:e1000139. http://dx.doi.org/10 .1371/journal.pgen.1000139.
- 53. Zaprasis A, Hoffmann T, Wunsche G, Florez LA, Stülke J, Bremer E. 19 June 2013. Mutational activation of the RocR activator and of a cryptic *rocDEF* promoter bypass loss of the initial steps of proline biosynthesis in *Bacillus subtilis*. Environ. Microbiol. [Epub ahead of print.]. http://dx.doi .org/10.1111/1462-2920.12193.
- Mäder U, Schmeisky AG, Florez LA, Stülke J. 2012. SubtiWiki—a comprehensive community resource for the model organism *Bacillus subtilis*. Nucleic Acids Res. 40:D1278–D1287. http://dx.doi.org/10.1093/nar /gkr923.
- 55. Thompson JD, Plewniak F, Thierry JC, Poch O. 2000. DbClustal: rapid and reliable global multiple alignments of protein sequences detected by

database searches. Nucleic Acids Res. 28:2919–2926. http://dx.doi.org/10 .1093/nar/28.15.2919.

- Altschul SF, Gish W, Miller W, Myers EW, Lipman DJ. 1990. Basic local alignment search tool. J. Mol. Biol. 215:403–410.
- Letunic I, Doerks T, Bork P. 2012. SMART 7: recent updates to the protein domain annotation resource. Nucleic Acids Res. 40:D302–D305. http://dx.doi.org/10.1093/nar/gkr931.
- Csonka LN. 1981. Proline over-production results in enhanced osmotolerance in *Salmonella typhimurium*. Mol. Gen. Genet. 182:82–86. http://dx .doi.org/10.1007/BF00422771.
- Smith CJ, Deutch AH, Rushlow KE. 1984. Purification and characteristics of a gamma-glutamyl kinase involved in *Escherichia coli* proline biosynthesis. J. Bacteriol. 157:545–551.
- 60. Bischoff DS, Ordal GW. 1992. *Bacillus subtilis* chemotaxis: a deviation from the *Escherichia coli* paradigm. Mol. Microbiol. 6:23–28. http://dx.doi .org/10.1111/j.1365-2958.1992.tb00833.x.
- Ordal GW, Villani DP, Nicholas RA, Hamel FG. 1978. Independence of proline chemotaxis and transport in *Bacillus subtilis*. J. Biol. Chem. 253: 4916–4919.
- Wong FH, Chen JS, Reddy V, Day JL, Shlykov MA, Wakabayashi ST, Saier MH, Jr. 2012. The amino acid-polyamine-organocation superfamily. J. Mol. Microbiol. Biotechnol. 22:105–113. http://dx.doi.org/10.1159 /000338542.
- 63. Hu LA, King SC. 1998. Membrane topology of the *Escherichia coli* gamma-aminobutyrate transporter: implications on the topography and mechanism of prokaryotic and eukaryotic transporters from the APC superfamily. Biochem. J. 336:69–76.
- 64. Shaffer PL, Goehring A, Shankaranarayanan A, Gouaux E. 2009. Structure and mechanism of a Na⁺-independent amino acid transporter. Science 325:1010–1014. http://dx.doi.org/10.1126/science.1176088.
- Gao X, Lu F, Zhou L, Dang S, Sun L, Li X, Wang J, Shi Y. 2009. Structure and mechanism of an amino acid antiporter. Science 324:1565– 1568. http://dx.doi.org/10.1126/science.1173654.
- 66. Ma D, Lu P, Yan C, Fan C, Yin P, Wang J, Shi Y. 2012. Structure and mechanism of a glutamate-GABA antiporter. Nature 483:632–636. http: //dx.doi.org/10.1038/nature10917.
- Javadpour MM, Eilers M, Groesbeek M, Smith SO. 1999. Helix packing in polytopic membrane proteins: role of glycine in transmembrane helix association. Biophys. J. 77:1609–1618. http://dx.doi.org/10.1016/S0006 -3495(99)77009-8.
- Dong H, Sharma M, Zhou HX, Cross TA. 2012. Glycines: role in alpha-helical membrane protein structures and a potential indicator of native conformation. Biochemistry 51:4779–4789. http://dx.doi.org/10 .1021/bi300090x.
- Wray LV, Jr, Zalieckas JM, Ferson AE, Fisher SH. 1998. Mutational analysis of the TnrA-binding sites in the *Bacillus subtilis nrgAB* and *gabP* promoter regions. J. Bacteriol. 180:2943–2949.
- Belitsky BR, Sonenshein AL. 2013. Genome-wide identification of *Bacillus subtilis* CodY-binding sites at single-nucleotide resolution. Proc. Natl. Acad. Sci. U. S. A. 110:7026–7031. http://dx.doi.org/10.1073/pnas.1300428110.
- 71. Belitsky BR, Sonenshein AL. 2002. GabR, a member of a novel protein family, regulates the utilization of gamma-aminobutyrate in *Bacillus sub-tilis*. Mol. Microbiol. 45:569–583. http://dx.doi.org/10.1046/j.1365-2958 .2002.03036.x.
- Bremer E. 2002. Adaptation to changing osmolarity, p 385–391. *In* Sonenshein AL, Hoch JA, Losick R (ed), *Bacillus subtilis* and its closes relatives: from genes to cells. ASM Press, Washington, DC.
- Bremer E, Krämer R. 2000. Coping with osmotic challenges: osmoregulation through accumulation and release of compatible solutes, p 79–97. *In* Storz G, Hengge-Aronis R (ed), Bacterial stress responses. ASM Press, Washington, DC.
- 74. Shelp BJ, Mullen RT, Waller JC. 2012. Compartmentation of GABA metabolism raises intriguing questions. Trends Plant Sci. 17:57–59. http: //dx.doi.org/10.1016/j.tplants.2011.12.006.
- 75. Planamente S, Vigouroux A, Mondy S, Nicaise M, Faure D, Morera S. 2010. A conserved mechanism of GABA binding and antagonism is revealed by structure-function analysis of the periplasmic binding protein Atu2422 in Agrobacterium tumefaciens. J. Biol. Chem. 285:30294–30303. http://dx.doi.org/10.1074/jbc.M110.140715.
- Liao MK, Gort S, Maloy S. 1997. A cryptic proline permease in Salmonella typhimurium. Microbiology 143:2903–2911. http://dx.doi.org/10 .1099/00221287-143-9-2903.
- 77. Jauniaux JC, Vandenbol M, Vissers S, Broman K, Grenson M. 1987.

Nitrogen catabolite regulation of proline permease in *Saccharomyces cerevisiae*. Cloning of the PUT4 gene and study of PUT4 RNA levels in wild-type and mutant strains. Eur. J. Biochem. **164**:601–606.

- Grallath S, Weimar T, Meyer A, Gumy C, Suter-Grotemeyer M, Neuhaus JM, Rentsch D. 2005. The AtProT family. Compatible solute transporters with similar substrate specificity but differential expression patterns. Plant Physiol. 137:117–126. http://dx.doi.org/10.1104/pp.104.055079.
- Lulko AT, Buist G, Kok J, Kuipers OP. 2007. Transcriptome analysis of temporal regulation of carbon metabolism by CcpA in *Bacillus subtilis* reveals additional target genes. J. Mol. Microbiol. Biotechnol. 12:82–95. http://dx.doi.org/10.1159/000096463.
- Belitsky BR. 2004. *Bacillus subtilis* GabR, a protein with DNA-binding and aminotransferase domains, is a PLP-dependent transcriptional regulator. J. Mol. Biol. 340:655–664. http://dx.doi.org/10.1016/j.jmb.2004.05.020.
- Shivers RP, Sonenshein AL. 2004. Activation of the *Bacillus subtilis* global regulator CodY by direct interaction with branched-chain amino acids. Mol. Microbiol. 53:599–611. http://dx.doi.org/10.1111/j.1365-2958.2004 .04135.x.
- Levdikov VM, Blagova E, Colledge VL, Lebedev AA, Williamson DC, Sonenshein AL, Wilkinson AJ. 2009. Structural rearrangement accompanying ligand binding in the GAF domain of CodY from *Bacillus subtilis*. J. Mol. Biol. 390:1007–1018. http://dx.doi.org/10.1016/j.jmb .2009.05.077.
- Molle V, Nakaura Y, Shivers RP, Yamaguchi H, Losick R, Fujita Y, Sonenshein AL. 2003. Additional targets of the *Bacillus subtilis* global regulator CodY identified by chromatin immunoprecipitation and genome-wide transcript analysis. J. Bacteriol. 185:1911–1922. http://dx.doi .org/10.1128/JB.185.6.1911-1922.2003.