

Studies of Regulation of Expression of the Propionate (*prpBCDE*) Operon Provide Insights into How *Salmonella typhimurium* LT2 Integrates Its 1,2-Propanediol and Propionate Catabolic Pathways

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Expression of the *prpBCDE* operon of *Salmonella typhimurium* LT2 required (i) the synthesis of propionyl-coenzyme A (CoA) by the PrpE protein or the acetyl-CoA-synthesizing systems of the cell and (ii) the synthesis of 2-methylcitrate from propionyl-CoA and oxaloacetate by the PrpC protein. We propose that either 2-methylcitrate or a derivative of it signals the presence of propionate in the environment. This as yet unidentified signal is thought to serve as a coregulator of the activity of PrpR, the member of the sigma-54 family of transcriptional activators needed for activation of *prpBCDE* transcription. The CobB protein was also required for expression of the *prpBCDE* operon, but its role is less well understood. Expression of the *prpBCDE* operon in *cobB* mutants was restored to wild-type levels upon induction of the propanediol utilization (*pdu*) operon by 1,2-propanediol. This effect did not require catabolism of 1,2-propanediol, suggesting that a Pdu protein, not a catabolite of 1,2-propanediol, was responsible for the observed effect. We explain the existence of these redundant functions in terms of metabolic pathway integration. In an environment with 1,2-propanediol as the sole carbon and energy source, expression of the *prpBCDE* operon is ensured by the Pdu protein that has CobB-like activity. Since synthesis of this Pdu protein depends on the availability of 1,2-propanediol, the cell solves the problem faced in an environment devoid of 1,2-propanediol where propionate is the sole carbon and energy source by having *cobB* located outside of the *pdu* operon and its expression independent of 1,2-propanediol. At present, it is unclear how the CobB and Pdu proteins affect *prpBCDE* expression.

Understanding metabolic pathway integration is a central issue in cell physiology. Learning more about this important aspect of cell function requires that we uncover and dissect the strategies used by the cell to ensure the coordinated and timely synthesis or degradation of metabolites.

In the recent past, the catabolism of propionic acid in *Salmonella typhimurium* LT2 (11, 13, 34) and *Escherichia coli* (9, 32) has been investigated. The cluster of genes required for the catabolism of propionate in these bacteria was first identified in and characterized for *S. typhimurium* LT2 (11, 13), with the closely related gene cluster in *E. coli* being reported as part of the genome project of this bacterium (3). These genes are referred to as the *prp* genes and are located in the 8.5-centisome region of the chromosome. These genes constitute a locus comprised of five genes organized in two transcriptional units (Fig. 1). One of these units contains the *prpR* gene, which encodes a putative member of the sigma-54 (RpoN) family of transcriptional activators (27). PrpR activity is required for the catabolism of propionate in *S. typhimurium* (20). The second transcriptional unit contains the *prpBCDE* gene cluster, which is organized as an operon that encodes propionate-degrading enzymes (13, 14). Work with *E. coli* has identified PrpC as the 2-methylcitrate synthase (9, 32). The 2-methylcitric acid cycle was first discovered in *Yarrowia lipolytica* (2, 29, 30), and in this cycle propionyl-coenzyme A (CoA) is α -oxidized to pyruvate (Fig. 1).

Our laboratory discovered that in addition to the *prpBCDE*

genes, catabolism of propionate in *S. typhimurium* requires the activity encoded by the *cobB* gene (34). The CobB gene product was first identified as an activity required for the expression of a phosphoribosyltransferase enzyme that restored synthesis of adenosylcobalamin in strains defective in the late steps of the biosynthesis of this coenzyme (33).

We have also reported that the inability of *cobB* mutants to catabolize propionate was corrected by the induction of the *pdu* genes required for the catabolism of 1,2-propanediol (1,2-PDL) (34). This observation was of interest to us because propionate, or an activated form of it (not clear at this point), is the end product of the pathway responsible for the catabolism of 1,2-PDL (4, 16, 37). We noted that induction of the *pdu* operon, without catabolism of 1,2-PDL, was sufficient to correct the Prp⁻ phenotype of *cobB* mutants. This finding was interpreted to mean that 1,2-PDL-dependent correction of this phenotype was due to the synthesis of a Pdu protein that compensated for the lack of CobB when propionate was the sole source of carbon and energy (34).

To better understand the implications of these results, it was essential to learn more about the roles of the Pdu and CobB proteins in propionate catabolism. These proteins may be enzymes that catalyze the same reaction, they may be regulatory proteins, or they may be both. Of particular interest to us was the possibility that CobB plays a regulatory role in the catabolism of propionate, since compensating for its absence through the induction of the *pdu* genes would provide evidence for pathway networking.

The regulation of expression of the *prpBCDE* operon is complex. PrpR is required, and at this point we assume that it has a coregulator, although it has not yet been identified. Propionate per se is not the coregulator, since it fails to induce

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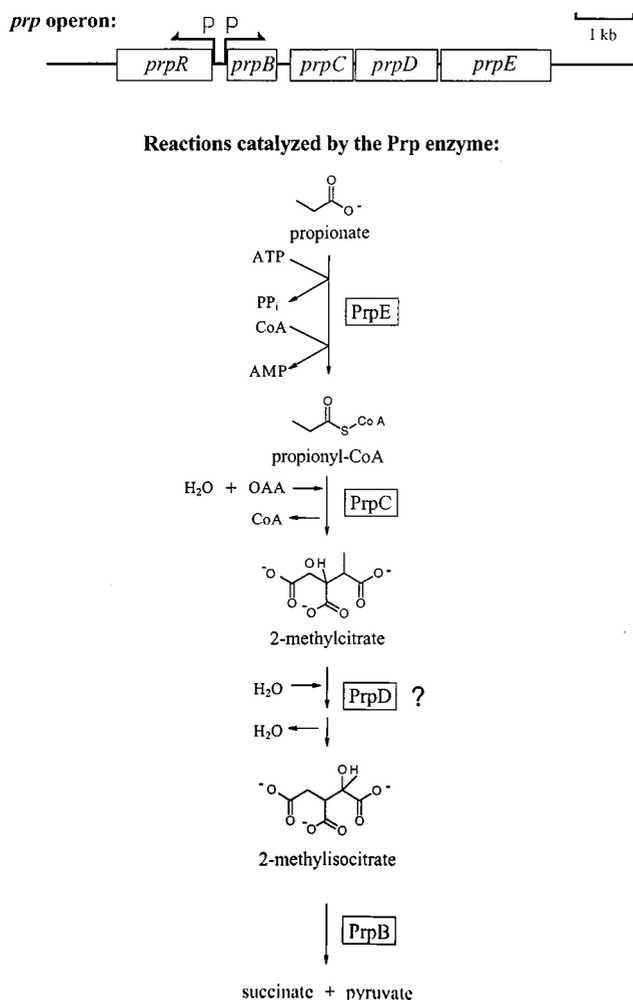


FIG. 1. Organization of the *prpBCDE* gene cluster and reactions proposed to be catalyzed by the *prp* gene products in the 2-methylcitric acid cycle. PrpE, propionyl-CoA synthetase (14); PrpC, 2-methylcitrate synthase (15); PrpD, putative 2-methylisocitrate synthase; PrpB, putative 2-methylisocitrate lyase. P, promoter.

transcription of the operon. This finding was shown by using transcriptional fusions of MudI1734 (*lacZ*⁺) elements (5) under the control of the *prpBCDE* promoter ($P_{prpBCDE}$) (11). Expression of the fusion was observed only in merodiploid strains that carry a wild-type copy of *prpBCDE* and a second copy of the operon, into which the MudI1734 element was inserted. These results were interpreted to mean that a catabolite of propionate was the signal for the presence of propionate in the environment.

In this paper, we show that CobB, PrpC, and PrpE are needed for transcription of the *prpBCDE* operon. We also show that the *pdu* function that allows growth of *cobB* mutants on propionate does so by fully restoring transcription of the *prpBCDE* operon. We suggest that 2-methylcitrate, the proposed product of the reaction catalyzed by PrpC (9, 32) or a derivative of it, is the coregulator needed for PrpR to activate transcription of the operon. Consistent with the requirement for PrpC activity, reduction of the intracellular levels of propionyl-CoA in a *prpC*⁺ strain mimicked the effect of a *prpC* mutation on the expression of the *prpBCDE* operon. The roles of CobB and of the uncharacterized Pdu protein are discussed within the framework of metabolic pathway integration and

physiological strategies used by the cell to ensure expression of target genes in response to multiple environmental stimuli.

MATERIALS AND METHODS

Bacterial strains, medium, and growth conditions. All strains used in this work were derivatives of *S. typhimurium* LT2, and their genotypes and all plasmids used are listed in Table 1. Cells were grown as detailed in Table 1 or the figure legends. The no-carbon E (NCE) medium was used as minimal medium. Nutrients and their concentrations in the medium were as follows: propionate, 30 mM; 1,2-PDL, 12 mM; glycerol, 22 mM; methionine, 0.5 mM; D-(+)-arabinose, 500 μM; and MgSO₄, 1 mM. Antibiotics were used at concentrations previously reported (8).

Genetic techniques. (i) **Transductions.** All genetic crosses were performed with the high-level-transduction mutant phage P22 *HT105 int-201* (25, 26) as described elsewhere (7). Transductants were freed of phage by streaking on indicator plates (6).

(ii) **Complementation analysis of *prp* point mutants.** The *prpB*, *prpC*, *prpD*, and *prpE* mutants used in experiments aimed at determining the involvement of *prp* functions in the expression of the *prpBCDE* operon were isolated after localized chemical mutagenesis with hydroxylamine (7, 12) as previously described (13). It was assumed that a single mutation was responsible for the inactivation of a gene product; however, it was not determined how many mutations were present in each affected gene. It should be noted that each one of these mutants was complemented by plasmids carrying the wild-type allele of the appropriate gene under the control of the arabinose-inducible promoter P_{araBAD} (13). The construction of these plasmids has been reported previously (13, 14). Plasmids were introduced into recipient strains by transformation (31). Inheritance of the plasmid was ensured by selecting for the antibiotic resistance carried by the cloning vector. Transformants were replica printed to NCE minimal medium supplemented with magnesium, propionate, and methionine.

(iii) **Construction of a *cobB* mutant carrying a duplication of the *prpBCDE* genes.** A Tn10-held duplication of the *S. typhimurium* LT2 spanning the 8- to 26-centisome region of the chromosome (DUP1033[*proA-pyrC*]) was moved by transduction into a strain carrying a deletion of *cobB* by selecting for tetracycline resistance. The insertion *prpC114::MudJ* was placed by transduction into one of the copies of the *prpBCDE* operon present within the duplicated region (Fig. 2). As reported elsewhere, the resulting *cobB* mutant was unable to grow on propionate (34). *cobB* mutant strains containing this duplication displayed a Prp⁻ phenotype and were routinely grown in the presence of tetracycline to avoid segregation of the duplicated material (1).

Recombinant DNA techniques. (i) **DNA sequencing.** Plasmid DNA was isolated with a QIAprep Spin Plasmid Miniprep kit of Qiagen Inc. (Chatsworth, Calif.) by following the manufacturer's instructions without modifications. PCR sequencing reaction mixtures were prepared with an ABI PRISM Dye Terminator Cycle Sequencing kit (Perkin-Elmer, Norwalk, Conn.) according to the manufacturer's instructions. Reaction mixtures were purified in AutoSeq G-50 columns (Pharmacia Biotech, Piscataway, N.J.), dried in a SpeedVac concentrator (Savant Instruments, Farmingdale, N.Y.), and sequenced at the Biotechnology Center of the University of Wisconsin—Madison.

(ii) **Plasmid constructions.** A ca.-790-bp *EcoRI* fragment from plasmid pPRP8 (13) containing $P_{prpBCDE}$ was cloned into the *EcoRI* site of plasmid pRS551 (28) to place the promoterless *lacZ*⁺ gene in pRS551 under the control of $P_{prpBCDE}$. The resulting plasmid, pPRP25 (Fig. 3), was used as a reporter of promoter activity. The orientation of this fragment was confirmed by DNA sequencing. Plasmid pPRP35 carries a wild-type allele of *prpC* under the arabinose-inducible promoter P_{araBAD} . The construction of this plasmid has been reported previously (13).

(iii) **Electroporation.** Plasmids were introduced into recipient cells by electroporation under conditions described elsewhere (19). Resistance to the antibiotic encoded by the plasmid was used to assess inheritance. Plasmid DNA was isolated as described above.

β-Galactosidase enzyme activity assay. β-Galactosidase activity assays were performed as described elsewhere (8). A unit of activity was defined as the amount of enzyme required to catalyze the hydrolysis of 1 nmol of *o*-nitrophenyl-α-D-galactopyranoside (ONPG) per min. Specific activity was reported as units per unit of absorbance at 650 nm (A_{650}).

RESULTS

In a previous report, we showed that expression of a *prp-lacZ* transcriptional fusion occurs only in merodiploid strains that carry a wild-type copy of the *prpBCDE* operon. This result suggested that the activity of one or more of the Prp proteins was needed to generate the catabolite of propionate required for activation of transcription of the operon (11).

Identification of *prp* functions encoded by the *prpBCDE* operon needed for expression of *prpBCDE*. To determine which functions encoded by the *prpBCDE* operon were re-

TABLE 1. Strain and plasmid list

Strain ^a or plasmid	Relevant characteristic(s)	Source or reference
Strains		
LT2	Wild type	Lab collection
Derivatives of LT2		
SMS209	<i>pta209::Tn10</i>	R. LaRossa
SMS408	<i>ack408::Tn10</i>	R. LaRossa
JE4265	DUP1033[<i>proC-pyrC</i>] <i>prpC114::MudI1734^b DELI184cobB</i>	
JE4334	DUP1033[<i>proC-pyrC</i>] <i>prpC114::MudJ</i>	
TR6583 (formerly SA2979)	<i>metE205 ara-9</i>	K. Sanderson via J. Roth
Derivatives of TR6583		
JE4044	pPRP25	
JE4388	<i>prpB210/pPRP25</i>	
JE4389	<i>prpB195/pPRP25</i>	
JE4390	<i>prpC173/pPRP25</i>	
JE4391	<i>prpC167/pPRP25</i>	
JE4392	<i>prpD174/pPRP25</i>	
JE4393	<i>prpD169/pPRP25</i>	
JE4394	<i>prpE213::kan/pPRP25</i>	
JE4395	<i>prpE213::kan DELI231 acs/pPRP25</i>	
JE4396	<i>prpE213::kan DELI231 acs ack408::Tn10/pPRP25</i>	
JE4397	<i>prpE213::kan DELI231 acs pta209::Tn10/pPRP25</i>	
JE4199	<i>prpC114::MudJ</i>	
JE4359	DUP1033[<i>proC-pyrC</i>] <i>prpC114::MudJ pdu-8::MudA DELI184cobB</i>	
JE4401	<i>prpC114::MudI1734/pBAD30</i>	
JE4402	JE4265/pCOBB5 <i>cobB⁺ cat⁺</i> (vector pSU19)	
JE4521	<i>prpE213::kan ack408::Tn10</i>	
JE4524	<i>DELI231 acs ack408::Tn10</i>	
JE4334	DUP1033[<i>proA-pyrC</i>] <i>prpC114::MudJ</i>	
JE4265	<i>DELI184cobB DUP1033[proA-pyrC] prpC114::MudJ</i>	
Plasmids		
pRS551	Cloning vector with a promoterless <i>lacZ⁺</i> gene, <i>kan⁺</i>	28
pSU19	Cloning vector, <i>cat⁺</i> (Cm ^r)	17
pBAD30	Expression vector, P _{<i>araBAD</i>} <i>bla⁺</i> (Ap ^r)	10
pPRP21	<i>prpB⁺</i> in pBAD30, <i>bla⁺</i> (Ap ^r)	13
pPRP25	P _{<i>prpBCDE</i>} - <i>lacZ kan⁺</i>	
pPRP35	<i>prpC⁺</i> in pBAD30, <i>bla⁺</i> (Ap ^r)	13
pPRP36	<i>prpD⁺</i> in pBAD30, <i>bla⁺</i> (Ap ^r)	13
pPRP54	<i>prpE⁺</i> in pBAD30, <i>bla⁺</i> (Ap ^r)	14
pCOBB5	<i>cobB⁺</i> cloned into pSU19 <i>cat⁺</i> (Cm ^r)	35

^a All strains used were derivatives of *S. typhimurium* LT2, and unless otherwise stated, they were constructed during the course of this study.

^b Hereafter and throughout the text, MudI1734 is also referred to as MudJ.

quired for its transcription, we investigated the effect that mis-sense mutations in *prpB*, *prpC*, or *prpD* had on the expression of a plasmid-encoded *lacZ* reporter gene placed under the control of P_{*prpBCDE*} (plasmid pPRP25). Allele *prpE213*, a kana-

mycin resistance cassette, was inserted into *prpE* by recombinant means (14). This insertion element ensured inactivation of *prpE* function, and since *prpE* is the most downstream gene in the operon, polarity of this insertion was not a concern.

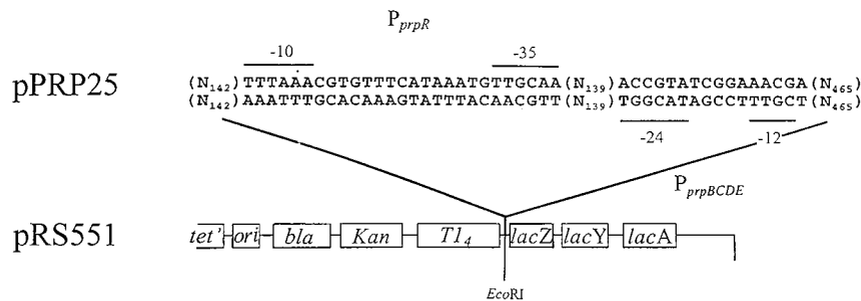


FIG. 2. Structure of reporter plasmid pPRP25. Details of the cloning are described in Materials and Methods. The orientation of the cloned fragment was verified by sequencing.

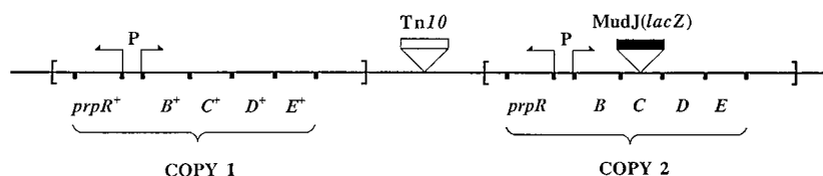


FIG. 3. Tn10-held chromosomal tandem duplication of the region containing the *prpBCDE* loci. The MudJ insertion places the *lacZ* gene under the control of $P_{prpBCDE}$ promoter.

Strains carrying mutations in *prpB*, *prpC*, or *prpD* displayed a severe Prp⁻ phenotype that, in all cases, was corrected by introducing into the strains a plasmid carrying a wild-type allele of the gene affected by the mutation. *prpE* mutants with an otherwise wild-type genetic background did not display a Prp⁻ phenotype due to redundant functions in the background (13, 14). When these functions were inactivated, however, the Prp⁻ phenotype of *prpE* mutants was complemented by a wild-type allele of *prpE*.

Complementation of the Prp⁻ phenotype of *prpB*, *prpC*, *prpD*, and *prpE* mutant strains by a single-gene plasmid showed that the mutations tested did not affect the synthesis of other gene products in the operon. Except for *prpE*, two independently isolated alleles of each gene were used in these studies to help validate the conclusions drawn. All strains tested carried a wild-type allele of *prpR*, the putative activator of *prpBCDE* transcription (13).

Loss of *prpC* function resulted in a severe reduction (ca. 95%) in the level of expression of the operon relative to the level of expression in the wild-type strain (Table 2). Lack of any of the remaining *prp* functions did not have as severe an effect. Noteworthy was the effect observed in strains lacking PrpD. In these mutants the level of *prpBCDE* expression was reduced ca. 66% (Table 2). Lack of PrpE reduced the level of expression by 30% (Table 2), and lack of PrpB resulted in a ca. 10% reduction in the level of expression of the operon relative to that in the wild type (Table 2). The roles of PrpD and PrpE are discussed further below.

These results indicated that PrpC activity was necessary for expression of the *prpBCDE* operon. We hypothesize that PrpC is probably involved in synthesizing the coregulator needed by PrpR to activate transcription of the *prpBCDE* operon.

As shown below, regulation of expression of the *prpBCDE* operon requires additional activities; that is, although PrpC activity was necessary, it was not sufficient to ensure expression of the operon.

PrpC provided in trans restores activity of the *prpBCDE* promoter in a haploid strain. Data presented in Table 3 demonstrate that the lack of PrpC activity was the reason why the *prpC114::MudJ* transcriptional fusion present in the chromosome was not expressed if the strain did not carry an additional, wild-type copy of the *prpBCDE* operon (11). When *prpC*⁺ was provided in trans, a 28-fold increase in expression of the *prpC114::MudJ* fusion was measured.

We note that similar levels of expression were measured when arabinose was not included in the medium, indicating that residual expression of *prpC* in the absence of the inducer resulted in levels of PrpC protein that were sufficient to restore full expression of the fusion. In contrast, the presence of arabinose was required to correct the Prp⁻ phenotype of a *prpC* missense mutant; that is, complementation of function was not observed in the absence of arabinose (data not shown). These results demonstrated the ability of arabinose to substantially increase the level of PrpC in the cell. As expected, the growth phenotype of strain JE4199 (*prpC114::MudJ/pPRP35 P_{araBAD} prpC*⁺) was not corrected in the presence or absence of arabinose (data not shown but see reference 13). These results were consistent with the need for *prpD* and *prpE* gene products to catabolize propionate. Neither PrpD nor PrpE is synthesized in strain JE4199 due to the polar effect of the MudJ element on *prpD* and *prpE*.

The dramatic increase in expression of the fusion in the absence of *prpD* and *prpE* gene products strongly suggested that the propionate catabolite needed for expression of the *prpBCDE* operon was either 2-methylcitrate (Fig. 1) or a derivative of it.

The pathway shown in Fig. 1 predicted that expression of the *prpBCDE* operon in a strain deficient in the synthesis of propionyl-CoA would not be restored by providing *prpC*⁺ in trans. This prediction was tested and proven to be correct. The results of the experiments addressing this point are discussed below.

TABLE 2. Effect of *prp* mutations on the expression of the *prpBCDE* operon

Strain	Relevant genotype ^a	Mean β -galactosidase activity \pm SD (U/A ₆₅₀ unit) ^b	% Expression ^c
JE4043	<i>prpRBCDE</i> ⁺ /pRS551 (vector control)	27 \pm 17	NA
JE4044	<i>prpRBCDE</i> ⁺ /pPRP25 ($P_{prpBCDE}$ - <i>lacZ</i> ⁺)	20,696 \pm 850	100
JE4390	<i>prpC173</i> /pPRP25 ($P_{prpBCDE}$ - <i>lacZ</i> ⁺)	1,320 \pm 130	6
JE4391	<i>prpC167</i> /pPRP25 ($P_{prpBCDE}$ - <i>lacZ</i> ⁺)	974 \pm 160	5
JE4393	<i>prpD169</i> /pPRP25 ($P_{prpBCDE}$ - <i>lacZ</i> ⁺)	6,987 \pm 687	34
JE4392	<i>prpD174</i> /pPRP25 ($P_{prpBCDE}$ - <i>lacZ</i> ⁺)	7,568 \pm 655	36
JE4394	<i>prpE213</i> /pPRP25 ($P_{prpBCDE}$ - <i>lacZ</i> ⁺)	14,039 \pm 1,578	68
JE4389	<i>prpB195</i> /pPRP25 ($P_{prpBCDE}$ - <i>lacZ</i> ⁺)	18,767 \pm 1,102	91
JE4388	<i>prpB210</i> /pPRP25 ($P_{prpBCDE}$ - <i>lacZ</i> ⁺)	18,267 \pm 395	88

^a All strains were derivatives of strain TR6583 (*metE205 ara-9*).

^b Assays were performed with mid-log-phase cultures grown in NCE medium supplemented with propionate, glycerol, and methionine. Assays conditions have been described elsewhere (7). A unit of activity was defined as the amount of enzyme that catalyzed the hydrolysis of 1 nmol of ONPG per min.

^c Obtained by dividing the level of expression of the *lacZ* gene in the mutant by the level of expression of the *lacZ* gene in the wild-type strain and multiplying by 100. Numbers are rounded up to the closest integer. NA, not applicable.

TABLE 3. *prpC* function provided in *trans* restores *prpBCDE* operon expression in a haploid strain

Strain	Relevant genotype ^a	Mean β -galactosidase activity \pm SD (U/A ₆₅₀ unit) ^b	
		- Arabinose	+ Arabinose
JE4401	<i>prpC114::MudJ/pBAD30</i> (vector control)	56 \pm 7	61 \pm 2
JE4199	<i>prpC114::MudJ/pPRP35</i> <i>P_{araBAD} prpC⁺</i>	1,390 \pm 136	1,564 \pm 59

^a All strains were derivatives of strain TR6583 (*metE205 ara-9*).

^b Assays were performed with mid-log-phase cultures grown in NCE medium supplemented with propionate (30 mM), glycerol (22 mM), methionine (0.5 mM), and arabinose (0.5 mM) whenever indicated. Assay conditions have been described elsewhere (8). A unit of activity was defined as the amount of enzyme that catalyzed the hydrolysis of 1 nmol of ONPG per min.

Propionyl-CoA is a precursor of the catabolite that signals the presence of propionate in the environment. Figure 4 shows the effect of the lack of propionyl-CoA on the expression of the *prpBCDE* operon. Work from our laboratory recently demonstrated that the *prpE* gene encodes a specific propionyl-CoA synthetase (14). It was also demonstrated that *prpE* mutants compensate for the lack of this enzyme through the activity of the acetyl-CoA synthetase encoded by the *acs* gene.

As discussed above, expression of the *lacZ* gene in plasmid pPRP25 in *prpE* mutant strain JE4394 was only slightly reduced relative to that in the *prpE*⁺ strain JE4044 (Table 2). Data in Fig. 4 show that expression of the reporter in the *prpE acs* double-mutation strain JE4395 was not significantly reduced relative to the level measured in the *prpE acs*⁺ strain JE4394 (36 and 32% reductions, respectively). A very drastic reduction in the expression of the reporter, however, was observed in triple-mutation strains carrying null alleles of *prpE acs ack* (strain JE4396) or *prpE acs pta* (strain JE4397). The Ack (acetate kinase) and Pta (phosphotransacetylase) enzymes are known to contribute to the synthesis of acetyl-CoA in the cell (18, 36). In strain JE4396 the level of β -galactosidase activity (mean \pm standard deviation, 1,661 \pm 407 U/A₆₅₀ unit) was reduced 92% relative to that in the wild-type strain JE4044 (20,696 \pm 850 U/A₆₅₀ unit). In strain JE4397 the level of

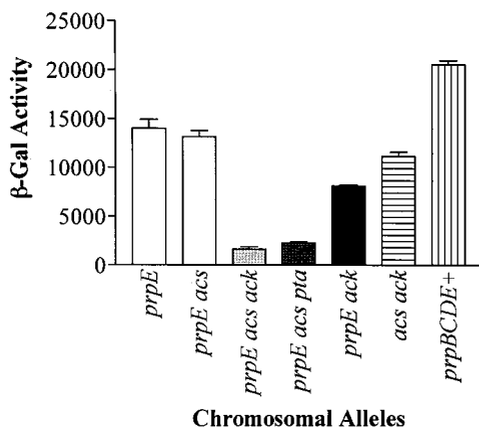


FIG. 4. Propionyl-CoA is a required precursor for the signal that triggers *prpBCDE* expression. β -Galactosidase specific activity is expressed in units per A₆₅₀ unit. The unit of activity for this enzyme is defined in Materials and Methods. The genes listed are *ack* (encodes acetate kinase), *pta* (encodes phosphotransacetylase), *prpE* (encodes propionyl-CoA synthetase), and *acs* (encodes acetyl-CoA synthetase). All mutations used were insertion alleles which are described in Table 1.

activity was reduced 89% relative to the level in the control strain JE4044 (2,286 \pm 206 versus 20,696 \pm 850 U/A₆₅₀ unit). Thus, the acetyl-CoA-synthesizing enzymes in *S. typhimurium* can activate propionate into propionyl-CoA. In addition, these results mimic those obtained with strains carrying mutations in *prpC*, indicating that PrpE (propionyl-CoA synthetase) activity is also required for expression of the *prpBCDE* operon.

It should be pointed out that a strain deficient in the synthesis of PrpE and Acs, but wild type for *ack* and *pta*, did not grow on propionate (14). In such a strain, expression of the *prpBCDE* operon was reduced only by a third relative to the level expressed by the wild-type strain (Fig. 4), and yet this strain failed to grow on propionate. To help explain this result, we argue that in the context of growth, Ack and Pta are poor substitutes for Acs or PrpE and that these enzymes cannot keep up with the demand of the pathway for propionyl-CoA to sustain cell growth. On the other hand, Ack and Pta can clearly generate enough propionyl-CoA to activate transcription of the *prpBCDE* operon. These results warn us about the risks of concluding that a high level of transcription of any gene necessarily means that the function of the corresponding gene product is sufficient for the cell to grow.

Although transcription of the *prpBCDE* operon in a *prpE ack* double mutant (strain JE4521) was only 39% of the level measured in the wild-type strain JE4044 (8,148 \pm 171 versus 20,696 \pm 850 U/A₆₅₀ unit), strain JE4521 grew well on propionate, indicating that, unlike Ack and Pta, Acs can satisfy the demand of the pathway for propionyl-CoA by itself and that the cell can grow on propionate. In the *acs ack* double mutant (strain JE4524), transcription of *prpBCDE* was high (11,196 \pm 785 or 50% of wild-type expression) and the strain grew well on propionate. These results were not surprising since strain JE4524 was wild type for *prpE*.

***cobB* function is required for transcription of the *prpBCDE* operon.** As alluded to in the introduction, one possible explanation for the inability of *cobB* mutants to grow on propionate was that CobB activity is required for the expression of the *prpBCDE* operon. To test this idea we measured the expression of the *prpC114::MudJ* transcriptional fusion in a merodiploid strain as a function of CobB. Data in Table 4 clearly show that *cobB* mutants fail to express the *prpBCDE* operon (Table 4). In the absence of CobB, expression of the operon was reduced 24-fold. This drastic reduction was due exclusively to the lack of CobB, since when a wild-type allele of *cobB* was provided in *trans*, transcription of the operon was restored to the levels observed in the *cobB*⁺ strain (Table 4). Plasmid pCOBB5 carries only a wild-type allele of *cobB* (35).

The ability of the strains used in these experiments to grow on propionate was consistent with the observed pattern of *prpBCDE* expression (Table 4).

Induction of the propanediol utilization (*pdu*) operon restores expression of *prpBCDE* and growth on propionate in *cobB* mutants. As mentioned above, we previously reported that induction of the propanediol utilization (*pdu*) genes by 1,2-PDL was sufficient to restore the ability of *cobB* mutants to grow on propionate as the carbon and energy source (34). Data in Table 4 show that induction of the *pdu* operon in a *cobB* mutant resulted in expression of the *prpBCDE* operon to levels observed in strains carrying a wild-type allele of *cobB* (Table 4). Consistent with previous phenotypic observations (34), introduction of the insertion element *pdu-8::MudA* into *cobB* mutant strain JE4265 completely eliminated the ability of the resulting strain (JE4359) to express the *prpBCDE* operon; hence, strain JE4359 failed to grow on propionate as the sole carbon and energy source in spite of the presence of 1,2-PDL in the medium (Table 4). These results further define our

TABLE 4. CobB or an uncharacterized Pdu protein is required for *prpBCDE* transcription in response to propionate or 1,2-PDL

Strain ^b	Relevant genotype	Mean β -galactosidase activity \pm SD (U/A ₆₅₀ unit) in cells grown in minimal medium ^a supplemented with:		Growth on Prp ^c	Growth on Prp, 1,2-PDL ^d
		No additions	1,2-PDL		
JE4334	<i>cobB</i> ⁺ DUP1033[<i>proA-pyrC</i>] <i>prpC114::MudJ</i>	663 \pm 51	770 \pm 20	Yes	Yes
JE4265	DEL1184 <i>cobB</i> DUP1033[<i>proA-pyrC</i>] <i>prpC114::MudJ</i>	28 \pm 7	750 \pm 37	No	Yes
JE4359	DEL1184 <i>cobB</i> DUP1033[<i>proA-pyrC</i>] <i>prpC114::MudJ pdu-8::MudA</i>	21 \pm 3	17 \pm 2	No	No
JE4402	JE4265/pCOBB5 <i>cobB</i> ⁺	692 \pm 50	ND ^e	Yes	Yes

^a NCE medium (7) containing glycerol (22 mM) and propionate (30 mM).

^b All strains were derivatives of strain TR6583 (*metE205 ara-9*).

^c NCE medium with 30 mM propionate (Prp) was the sole carbon and energy source.

^d NCE medium with 30 mM propionate (Prp) and 12 mM 1,2-PDL. The medium did not contain cobalamin to prevent 1,2-PDL degradation.

^e ND, not determined.

understanding of the role of CobB in propionate catabolism and provide a good example of pathway networking in the cell.

DISCUSSION

The work reported herein makes two important contributions to our understanding of the physiology of *S. typhimurium*: (i) it provides insights into the complex regulation of transcription of the *prpBCDE* operon, which encodes functions required for the catabolism of propionate in this bacterium, and (ii) it provides strong evidence for metabolic pathway integration.

Propionyl-CoA synthetase (PrpE), 2-methylcitrate synthase (PrpC), and CobB activities are needed to activate *prpBCDE* operon transcription. Data reported in this paper show that three proteins are required for expression of this operon, namely, PrpE, PrpC, and CobB or its alternative, an uncharacterized Pdu protein. All these proteins are needed in addition to PrpR, the sigma-54 activator encoded by *prpR* (13, 20).

Two of these proteins have documented biochemical activities associated with them: PrpE has propionyl-CoA synthetase activity (14) and PrpC has 2-methylcitrate synthase activity (9, 15, 32). The role of PrpE and PrpC in *prpBCDE* transcription can be rationalized on the basis of the proposed pathway for propionate catabolism in this bacterium (Fig. 1). On the basis of our data, we hypothesize that the catabolite of propionate that signals the presence of this fatty acid in the environment is either 2-methylcitrate or a derivative of it. This metabolite may be the coregulator needed by PrpR to activate transcription of *prpBCDE*.

Since PrpC catalyzes the synthesis of 2-methylcitrate from propionyl-CoA and oxaloacetate, it is clear that PrpE activity is central to the generation of one of the substrates for PrpC. That is why the effect of blocking propionyl-CoA synthesis in *prpC*⁺ strains mimics the effect that mutations in *prpC* have on the expression of the *prpBCDE* operon.

Role of the PrpD and PrpB proteins in *prpBCDE* expression. Our data on the involvement of PrpB in the regulation of the *prpBCDE* operon suggest that this putative 2-methylisocitrate lyase enzyme activity most likely does not play a role in the generation of the catabolites that signals the presence of propionate in the environment.

Although the data for PrpD are less clear, we also conclude that PrpD activity is not needed for the generation of the signal. The data show a significant reduction in *prpBCDE* expression in *prpD* mutants that suggests that PrpD activity may be required to generate the signal needed to activate *prpBCDE* transcription. However, this reduction should be interpreted

with caution. The same effect would be observed if the absence of PrpD had the net effect of reducing the amount of 2-methylcitrate made by PrpC, in which case, the observed reduction of *prpBCDE* operon expression in *prpD* mutants would be the result of suboptimal concentrations of the signal needed by PrpR to activate transcription of the operon. An additional argument supporting the idea that PrpD activity may not be needed to generate the signal comes from the experiments that analyzed the expression of the *prpC114::MudJ* fusion in a haploid strain with or without *prpC*⁺ in *trans* (Table 3). In these experiments, PrpC was sufficient to restore full expression of the fusion in a strain where expression of *prpD* was presumably eliminated by the MudJ element upstream of this gene. At this point, we cannot rule out the possibility that a small but sufficient amount of PrpD was synthesized in this strain. Such a small amount of PrpD would have to be sufficient to allow regulation at the wild-type level to occur.

One important difference between these experiments and the ones performed with strains carrying duplications of the *prpBCDE* operon should be kept in mind. In the experiments with the haploid strain, *prpC*⁺ was carried by a high-copy-number plasmid; thus, the concentration of PrpC protein was greater than the concentration afforded by a single chromosomal copy of *prpC* in the strain with the duplication. This high concentration of PrpC, we argue, is likely to generate sufficient 2-methylcitrate to make PrpR fully functional in the absence of PrpD.

Roles of the CobB and Pdu proteins in *prpBCDE* operon expression. The roles that the CobB protein and its alternative, Pdu protein, play in the transcription of *prpBCDE* are not obvious. The CobB and Pdu proteins may have one or more enzymatic activities needed for the generation of a metabolite required for *prpBCDE* transcription activation. Alternatively, they may be proteins without any enzymatic activity that play some role in either transcription activation or attenuation, or in a more complex scenario these proteins may have one or more enzymatic activities in addition to playing a more direct role in transcription activation and/or posttranscriptional regulation of the *prpBCDE* operon expression. How these proteins affect expression of the *prpBCDE* operon remains under investigation.

Integration of the 1,2-PDL and propionate catabolism pathways. Regardless of how the CobB and Pdu proteins affect transcription of the *prpBCDE* operon, we believe that the existence of these redundant functions reflects physiological strategies to ensure that synthesis of the propionate degrading

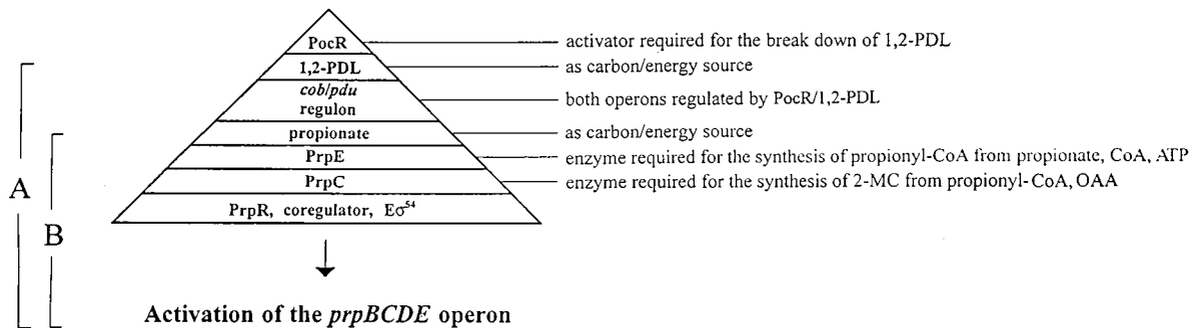


FIG. 5. Elements controlling *prpBCDE* operon expression in the presence or absence of propionate in the environment. Under all conditions, propionyl-CoA synthetase (PrpE) and 2-methylcitrate synthase (PrpC) activities are needed to synthesize the coregulator (2-methylcitrate or a derivative of it) required by PrpR to activate transcription of the operon. Under all conditions, core (E) RNA polymerase with sigma-54 (RpoN) is needed for expression. (A) Under conditions where 1,2-PDL is the sole source of carbon and energy, expression of the *prpBCDE* operon CobB activity is dispensable due to the synthesis of an uncharacterized protein encoded by the *pdu* operon. This Pdu protein compensates for the lack of CobB in *cobB* mutants. (B) Under conditions where propionate is the sole source of carbon and energy, CobB activity is required. 2-MC, 2-methylcitrate.

enzymes occurs under different environmental conditions. Since 1,2-PDL catabolism is likely to proceed via propionyl-CoA, it is clear that expression of the *prpBCDE* operon is needed for the cell to be able to use 1,2-PDL as a carbon and energy source.

It appears that the Pdu alternative for CobB function is a mechanism that the cell has evolved to ensure the timely synthesis of the PrpB, -C, -D, and -E enzymes to further degrade propionyl-CoA into central metabolites. Figure 5 illustrates how we think the expression of the *prpBCDE* operon is ensured when either 1,2-PDL or propionate is present in the environment. When 1,2-PDL is the sole carbon and energy source, it binds to the PocR protein (21, 22) and the PocR-1,2-PDL complex activates transcription of the *cob* or *pdu* regulon (23, 24). Since this Pdu function is available only upon induction of the operon by 1,2-PDL, *S. typhimurium* would be unable to use propionate as a carbon and energy source in an environment devoid of 1,2-PDL. One way to solve this problem would be to have a redundant function encoded outside of the *pdu* operon, with the expression of such a gene being independent of the presence of 1,2-PDL. In this case, the redundant function is encoded by *cobB*. Learning more about how the CobB and Pdu proteins affect *prpBCDE* operon expression will shed light on one mechanism used by this bacterium to integrate its metabolism.

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