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Streptococcus pyogenes Polymyxin B-Resistant Mutants Display Enhanced ExPortal Integrity

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The ExPortal protein secretion organelle in *Streptococcus pyogenes* is an anionic phospholipid-containing membrane microdomain enriched in Sec translocons and postsecretion protein biogenesis factors. Polymyxin B binds to and disrupts ExPortal integrity, resulting in defective secretion of several toxins. To gain insight into factors that influence ExPortal organization, a genetic screen was conducted to select for spontaneous polymyxin B-resistant mutants displaying enhanced ExPortal integrity. Whole-genome resequencing of 25 resistant mutants revealed from one to four mutations per mutant genome clustered primarily within a core set of 10 gene groups. Construction of mutants with individual deletions or insertions demonstrated that 7 core genes confer resistance and enhanced ExPortal integrity through loss of function, while 3 were likely due to gain of function and/or combinatorial effects. Core resistance genes include a transcriptional regulator of lipid biosynthesis, several genes involved in nutrient acquisition, and a variety of genes involved in stress responses. Two members of the latter class also function as novel regulators of the secreted SpeB cysteine protease. Analysis of the most frequently isolated mutation, a single nucleotide deletion in a track of 9 consecutive adenine residues in *pstS*, encoding a component of a high-affinity P_i transporter, suggests that this sequence functions as a molecular switch to facilitate stress adaptation. Together, these data suggest the existence of a membrane stress response that promotes enhanced ExPortal integrity and resistance to cationic antimicrobial peptides.

Numerous host-pathogen interactions depend upon the timely coordination of secretion and processing of multiple proteins, including adhesion proteins and toxins. For Gram-positive pathogens, this represents a significant problem because these bacteria lack a periplasmic space with which to spatially coordinate the interaction of nascently secreted polypeptides with the accessory factors required for their folding and processing. One solution found among certain species of streptococci and enterococci is to cluster secretion and accessory factors at a defined membrane microdomain known as the ExPortal (1–5). However, the factors that promote the integrity and organization of the Ex-Portal microdomain are not well understood.

The ExPortal has been most extensively analyzed in Streptococcus pyogenes, an important human pathogen and the causative agent of numerous diseases ranging from superficial infections of soft tissue (e.g., pharyngitis) to toxigenic diseases (e.g., scarlet fever) to those destructive of tissue (e.g., necrotizing fasciitis). Infection can also trigger several postinfection sequelae that may arise from an inappropriate immune response (e.g., rheumatic fever) (6). To cause this array of diseases, S. pyogenes secretes a multitude of extracellular toxins and cell wall-associated proteins (7). Many of these require accessory factors to facilitate their biogenesis into an active form or for their incorporation into the cell wall and presentation at the cell surface (8). Of the few secretion systems identified in Gram-positive bacteria, including twin-arginine translocation (TAT), ESX/type VII secretion, and the accessory SecA2 system, S. pyogenes encodes only the ubiquitous and highly conserved canonical Sec (SecA1) pathway (9). Components of the Sec translocon are enriched at the ExPortal microdomain (1, 2) along with accessory factors required for toxin biogenesis, including the HtrA protease/chaperone (1). Mutations that mislocalize HtrA away from the ExPortal result in significant defects in the postsecretion processing of certain secreted toxins, including the SpeB cysteine protease (1). These data suggest that ExPortal organization functions to facilitate the postsecretory interaction of substrate and accessory biogenesis factors. Similar studies showing that mislocalization of sortase away from the Ex-Portal in *Enterococcus faecalis* impairs the efficiency of pilus biogenesis (5) also support a critical role for the ExPortal in the spatial coordination of virulence factor biogenesis.

While the mechanism that underlies the structural organization of the ExPortal is not understood, some insight has come from the observation that its lipid composition is asymmetric with respect to the peripheral membrane and is enriched in its content of anionic phospholipids (10). Microdomains enriched for anionic phospholipids have been observed in numerous bacterial species (11–13) and have been implicated in multiple cellular functions, including cell division (11), and in maintenance of Sec translocons along a longitudinal spiral pattern in certain rodshaped bacteria (14). In the case of the *E. faecalis* pilus sortase (SrtC), mutations that reduce the density of positively charged residues of the cytoplasmic tail of its single transmembrane domain result in redistribution of SrtC from the ExPortal to the peripheral membrane (5). Together, these observations implicate protein-lipid charge interactions as an organizing principle re-

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sponsible for retention of at least some proteins at the anionic ExPortal microdomain.

In a similar fashion, redistribution of ExPortal anionic lipids also results in redistribution of ExPortal-retained proteins. The anionic character of the microdomain renders it a target of cationic antimicrobial peptides (CAPs), and at low sublethal concentrations, certain CAPs, including polymyxin B (PB) and human neutrophil peptide 1 (HNP-1), preferentially target the ExPortal over the peripheral membrane (15). At higher but still sublethal concentrations, these CAPs disrupt the ExPortal, resulting in the redistribution of both anionic lipids and retained proteins from a focal location at the ExPortal to a more uniform circumferential distribution around the peripheral membrane (15). As a consequence, this redistribution is associated with the inhibition of secretion of several toxins important for virulence (15). These data further support a role for protein-lipid interaction in ExPortal organization. However, how the anionic phospholipids themselves are organized and maintained as part of a discrete microdomain is not understood.

In the present study, we have exploited the ability of CAPs to disrupt the ExPortal in order to gain insight into factors that participate in ExPortal assembly and maintenance. Our approach was to first select for spontaneous mutants of *S. pyogenes* resistant to the lethal effects of PB and then to identify those that bound PB in a manner indistinguishable from the wild type (WT). This pool was then analyzed to identify mutants that displayed enhanced ExPortal integrity in the face of PB challenge. This strategy generated a variety of mutant classes, including those with altered lipid metabolism, nutrient acquisition, and stress responses. Together, these data demonstrate that numerous cellular processes are involved in promoting the integrity of the ExPortal.

MATERIALS AND METHODS

Escherichia coli strains, media, and growth conditions. Routine molecular cloning and plasmid propagation utilized *E. coli* DH5 α , which was cultured in Luria-Bertani medium at 37°C. When appropriate, antibiotics were added to media at the following concentrations: 500 µg ml⁻¹ erythromycin, 100 µg ml⁻¹ spectinomycin, and 15 µg ml⁻¹ chloramphenicol.

S. pyogenes strains, media, and growth conditions. Experiments utilized Streptococcus pyogenes HSC5 (16, 17) and mutant derivatives of this strain. The spontaneous PB-resistant (PBr) mutants derived in this study are listed in Table 1, and the full genome sequence data are listed in Table S6 in the supplemental material. Other mutants used, including the various insertion and deletion mutants in various core PBr genes, are listed in Tables S1 and S2. Gene deletion mutants were generated using standard techniques (18) except in the case of the $\Delta PtsI::cat$ mutant, whereby a promoterless chloramphenicol cassette (cat) was used to select for ptsI gene replacement upon plasmid integration at the nonpermissive temperature. Routine culture employed Todd-Hewitt medium (Difco) supplemented with 0.2% yeast extract (Difco) (THY medium). When indicated, strains were cultured in C medium (0.5% protease peptone no. 3 [Difco], 1.5% yeast extract [Difco], 10 mM K₂HPO₄, 0.4 mM MgSO₄, 17 mM NaCl). Salt-reduced C medium was identical, except that it lacked all added salts. For experiments involving carbohydrate supplementation, filter-sterilized 20% (wt/vol) stock solutions were used to add each of the indicated carbohydrates (Sigma) to a final concentration of 0.2% (wt/vol) in medium that had been sterilized in an autoclave, with the exception of raffinose, malate, and maltodextrin, which were prepared in 10%, 5%, and 0.4% (wt/vol) stock solutions, respectively, and added to sterilized 2× C medium which was further diluted with sterile water to reach finally 1× C medium. Unless otherwise indicated, all growth experiments utilized sealed culture tubes at 37°C under static conditions. Solid medium was

prepared by the addition of 1.4% Bacto agar (Difco) and was cultured anaerobically in sealed jars using a commercial gas generator (GasPak; BBL) as described previously (19). When appropriate, antibiotics were added at the following concentrations: 1 μ g ml⁻¹ erythromycin, 100 μ g ml⁻¹ spectinomycin, and 3 μ g ml⁻¹ chloramphenicol.

Selection of polymyxin B-resistant mutants. Wild-type HSC5 was cultured overnight in 50 ml THY medium. The following day, cultures were centrifuged at 6,000 × g for 5 min and the bacterial pellet was resuspended in 5 ml fresh THY medium. One-hundred-microliter aliquots (yielding approximately 5×10^7 CFU when plated under nonselective conditions) were plated onto protease indicator plates (solidified C medium supplemented with 2% milk) containing 400 µg ml⁻¹ (289 µM) polymyxin B (Sigma) and incubated at 37°C for up to 48 h under anaerobic conditions. Colonies that emerged were retested for growth on fresh polymyxin B-containing plates. Colonies verified as resistant were cultured overnight in THY medium and stored at -80°C with 20% glycerol for further analysis.

Resequencing analyses. Genomic DNA was extracted using the FastDNA Spin kit (MP Biomedical; catalog no. 116540-600) and subjected to DNA sequence analysis using the Illumina HiSeq 2000 platform (conducted by GTAC, Washington University, St. Louis, MO) by standard methods as described previously (16). Sequence reads were aligned to the reference HSC5 scaffold sequence using DNAstar SeqMan NGen 4.0.0 (DNAstar) using the default stringency parameters. Single nucleotide polymorphisms (SNPs) and single nucleotide insertions and deletions (InDels) were identified by setting the SNP filter to a minimum of 30% coverage and manually verified by inspection of the Illumina sequence data. On average, SNPs had ~80-fold coverage, but those with <15-fold coverage were manually verified by PCR and Sanger sequencing. Large deletions (deletions of >1 nucleotide) were either verified by realigning Illumina reads to a scaffold sequence altered to contain the deletion or manually verified by PCR and Sanger sequencing. Genomic loci follow the format L897_xxxxx, where xxxx are numbered. Gene names and annotations either are based on a consensus of all annotated S. pyogenes genomes available on www.BioCyc.org (20), or in the absence of consensus, are based on homology to other genomes-Streptococcus pneumoniae (fabT) and Bacillus subtilis 168 (gdpP [yybT]; yfmH; nupP [*yufP*])—or are based on annotated function (*luxR* family transcriptional activator).

Analysis of polymyxin B binding and disruption of SpeB secretion. Various mutant and wild-type S. pyogenes strains were cultured in C medium and challenged with a sublethal concentration of polymyxin B as described in detail elsewhere (15). Briefly, 30 µM polymyxin B was added to late-logarithmic-phase cells and allowed to grow for an additional 2 h before supernatants were collected and monitored for SpeB secretion by Western blotting using polyclonal anti-SpeB rabbit serum (Abcam; ab53403). For analysis of polymyxin B binding, late-logarithmic-phase cells were stained with dansyl-polymyxin B (catalog no. P13238; Invitrogen) at low (5 μ M) or high (30 μ M) concentrations and examined by fluorescence microscopy as described previously (15). Focal localization in images of cells was scored as staining at a unique focus, staining at multiple foci, or circumferential staining (staining was of homogeneous intensity around the cellular circumference) (5). A strain was considered to have an intact ExPortal if >80% of cells examined demonstrated focal binding. Each mutant was examined in a minimum of three independent experiments with examination of 1,000 total stained cells.

DNA techniques. Plasmid DNA was isolated via standard techniques and used to transform *S. pyogenes* or *E. coli* as described previously (21). In-frame deletion mutations in genes were generated using the vector pJRS233 (22) or pGCP213 (23), and insertion mutations were generated using the vector pSPC18 (24), to construct the mutagenic plasmids listed in Table S3 in the supplemental material. Each deletion or insertion allele was generated by PCR through a process of overlap extension PCR (25), and modified alleles were inserted directly into plasmids at the M13F and M13R universal primer binding sites utilizing the overlap extension PCR

TABLE 1 Spontaneous PBr mutants

	PB culture ^b		DD		Duadiated		Mutation ^e	
Strain ^a	Growth	SpeB	binding ^c	Locus ^d	protein ^d	Annotation ^d	Туре	Change
WT	-	-	+	NA	NA	NA	NA	NA
PBr 1.1	+	+	+	04725	PstS	Phosphate ABC transporter	InDel	fs-aa6
PBr 2.1	+	+	+	04725	PstS		InDel	fs-aa6
PBr 3.1	+	+	+	03260	EbsA	Pore-forming protein	SNP	I95N
				04415	ТорА	DNA topoisomerase I	InDel	fs-aa661
				04725	PstS		InDel	fs-aa6
PBr 3.2	+	+	+	00065	FtsH	Cell division protein	InDel	$\Delta 293$
				04720	PstC	Phosphate ABC transporter permease	SNP	Q126*
PBr 3.3	+	+	+	04725	PstS		InDel	fs-aa6
				09125	GdpP	GGDEF domain phosphodiesterase	InDel	fs-aa3
PBr 3.5	+	+	+	07215	FabT	Transcription repressor, fatty acid biosynthesis	SNP	S84L
				09125	GdpP		InDel	fs-aa218
				09180	YfmH	M16 family zinc protease	InDel	fs-aa140
PBr 3.6	+	+	+	02925	AgaS	Galactosamine-6P isomerase	SNP	G72D
				07145	ManN	Mannose-specific PTS IID protein	SNP	I75F
				07215	FabT	A A	SNP	*145L
				09125	GdpP		InDel	fs-aa76
PBr 3.7	+	+	+	00065	FtsH		SNP	V291I
				04725	PstS		InDel	fs-aa6
PBr 4.1	+	_	+	03620	ClpX	Clp protease ATP-binding subunit	SNP	O311*
PBr 4.4	+	+	+	04725	PstS		InDel	fs-aa6
				09125	GdpP		InDel	$\Delta 642 - 645$
PBr 4.5	+	+	+	03645	DeoB	Phosphopentomutase	SNP	S87Y
				04725	PstS	I I I I I I I I I I I I I I I I I I I	InDel	fs-aa6
PBr 6.3	+	+	+	07145	ManN		SNP	G166D
PBr 6.7	+	+	+	00060	Hpt	Hypoxanthine-guanine phosphoribosyltransferase	InDel	fs-aa20
				09125	GdpP	/F	InDel	fs-aa563
PBr 6.10	+	+	+	00055	TilS	tRNA ^{IIe} -lysidine synthase	InDel	$\Delta 428$
				00060	Hpt		InDel	ΔRBS-53
				07135	ManL	Mannose-specific PTS IIAB protein	SNP	N284K
PBr 9.21	+	_	+	03620	ClpX	······································	InDel	$\Delta 352 - 356$
				07215	FabT		SNP	H39N
PBr 10.6	+	+	+	07215	FabT		SNP	S84L
				07750	NA	Phage repressor protein	InDel	(-24)
				09125	GdpP	Thige represser protein	SNP	0547*
PBr 10 9	+	+	+	04415	TopA		InDel	fs-aa27
101100				04715	PstA	Phosphate ABC transporter permease	InDel	fs-aa35
				06715	Gmk	Guanylate kinase	SNP	L131I
PBr 10 11	+	_	+	01235	NanH	N-Acetylneuraminate lyase	SNP	[V120V]
1 D1 10111				03620	ClpX	1 Treedy medianimate 1/450	InDel	fs-aa178
				07215	FabT		SNP	A56V
PBr 10.14	+	+	+	04545	GuaA	GMP synthase	SNP	A248D
				04725	PstS		InDel	fs-aa6
PBr 10.15	+	+	+	00055	TilS		SNP	G428E
				00060	Hpt		SNP	(-9) RBS
PBr 11.1	+	_	+	032.60	EbsA		InDel	$\Delta 21-25$
				03620	ClpX		SNP	E263*
				04650	NupP	Guanosine ABC transporter permease	SNP	W209*
				07215	FabT	2oome me o transporter permease	SNP	(-9) RRS
PBr 11.2	+	+	+	00055	TilS		SNP	*429R
		,		00060	Hpt		SNP	(-7) RRS
				04415	TopA		SNP	R646*
PBr 11 3	+	+	+	05585	PteI	PED_protein phosphotransferase	SNP	A510V
1 DI 11.3		1	1	03383	LuxP	Transcriptional regulator LuxD family	SND	D693V
				09125	GdpP	manscriptional regulator, Luxic lanniny	SND	A574D
DBr 11 5	+	+	+	07125	Map		SINP	N284V
1 DI 11.3	Ŧ	T	Ŧ	0/133	Fba	Fructore bisphosphate aldolare	SINP	INZO4K
DBr 11 4	+	+	+	04415	Top A	raciose-bisphosphate aldolase	SINP	V20D
F DI 11.0	Ŧ	Ŧ	Ŧ	05595	TOPA DtoI		SINF	V 29D
				03383	F 151		SINF	D1041

^{*a*} WT is HSC5. Mutants whose designations begin with different numbers were isolated from independent experiments.

^b Ability (+) or inability (-) to form colonies (growth) and express SpeB protease activity (SpeB) when grown on protease indicator plates supplemented with 400 μ g/ml PB. ^c Focal (+) or undetectable (-) membrane binding when stained with 5 μ M dansyl-PB.

^d Genetic loci, the predicted proteins that an ORF encodes, and annotations are based on the genome of HSC5 (16). Loci follow the format L897_xxxxx, where xxxxx are numbered. NA, not applicable. Annotations are listed only at first instance. In subsequent examples, the field is left blank.

^{*e*} InDel, insertion/deletion; SNP, single nucleotide polymorphism; "fs-aa" followed by numeral, frameshift at the indicated codon; *, stop codon; number in parentheses, distance upstream of start codon; RBS, ribosome binding site; data in brackets, synonymous mutation; Δ, deletion. Note that *tilS* and *hpt* mutations result from a single alteration since the 3' end of *tilS* overlaps the RBS of *hpt*.

^f PEP, phosphoenolpyruvate.

cloning method described in detail elsewhere (26) using primers listed in Table S4. The fidelity of all molecular constructs and mutated chromosomal loci was confirmed by PCR and determination of DNA sequences (Genewiz) using oligonucleotide primers (IDT) of the appropriate sequences. Putative -35 and -10 promoter elements were identified using the online software BProm (Softberry, Inc., Mount Kisco, NY) (27), while ribosome binding sites (RBS) were manually inspected. The consensus binding site for the FabT transcriptional regulator was calculated using the online software RegPrecise (http://regprecise.lbl.gov) (28) from a consensus of 56 putative binding sites from a variety of streptococcal and lactococcal species. The *fab* operon from the *S. pneumoniae* strain D39 was used for comparison (29).

Analysis of transcript. Transcript abundances of select genes were analyzed by real-time reverse transcription-PCR (RT-PCR) as described previously (30). Briefly, for *speB* gene expression analysis, overnight cultured cells were diluted 1:25 or 1:10 into fresh C medium and harvested at mid-logarithmic phase (50% of final optical density at 600 nm [OD₆₀₀]), late logarithmic phase (75% of final OD_{600}), or stationary phase (100% of final OD_{600}) or 2 h after onset of stationary phase for mRNA extraction. For fatty acid and anionic lipid biosynthetic gene expression analysis, overnight-cultured cells were diluted 1:25 into fresh THY medium and harvested at early logarithmic phase (20% of final OD₆₀₀) for mRNA extraction. Select genes were analyzed by real-time RT-PCR normalized to recA using the primers listed in Table S5 in the supplemental material. speB transcript is displayed as threshold cycle $(-\Delta C_T)$, normalized to recA for each strain and time point, while fatty acid and anionic lipid biosynthetic genes are displayed as $\Delta\Delta C_T$, normalized to the wild type. RNA data shown are the means and the standard errors of the means derived from triplicate determinations of samples from at least three independent experiments.

Analysis of SpeB protease activity. The ability of various strains to express active SpeB protease was assessed on protease indicator plates (solidified C medium supplemented with 2% milk) (19) as an activity titer, determined as follows: cells cultured overnight in C medium were centrifuged and concentrated $20 \times$ to $200 \times$ into fresh C medium. A series of seven 10-fold dilutions were prepared, and 5-µl aliquots were spotted onto protease indicator plates and incubated at 37°C for 40 h under anaerobic conditions. CFU were quantified at dilutions which resulted in 3 to 30 CFU, and CFU were estimated to be 10-fold greater at each preceding dilution. Regions surrounding bacterial growth were monitored every 4 h for zones of clearance, which is indicative of SpeB proteolysis activity. Activity was quantified as the lowest dilution at which clearing occurred upon each observation, or if no clearing was observed, the CFU value was given as 10-fold greater than the highest dilution plated, and data were plotted as 7-log₁₀ (CFU_{clear}). Activity data shown are the means and the standard errors of the means derived from triplicate determinations of samples and are representative of at least three independent experiments. As opposed to the fluorescein isothiocyanate (FITC)-casein cleavage assay, protease indicator plates were utilized to measure SpeB activity over time, since this assay is technically simple to perform and is easily scalable to allow for testing of multiple strains simultaneously in order to monitor expression and activation kinetics.

Growth rate and yield analyses. Indicated bacterial strains were backdiluted 1:50 into 1 ml of fresh C medium (unmodified or altered as indicated in the text). Cultures were mixed briefly by vortexing, 200 μ l was placed in triplicate into a 96-well plate (catalog no. 655-180; Greiner), and plates were sealed using transparent adhesive film (catalog no. 60941-078; VWR). Monitoring of growth at 37°C was performed in a Tecan Infinite M200 Pro plate reader. During growth, the plate was shaken every 5 min for 30 s, followed by a 5-s wait period and measurement of the OD₆₀₀. Data were normalized by subtracting the OD₆₀₀ value of a blank well (200 μ l uninoculated C medium or C medium supplemented with ascorbic acid, which is oxygen sensitive and darkens the medium over time), and the first derivative of the log₁₀ [(1,000) OD₆₀₀] of each time point was input into Prism (GraphPad). The 13 time points (60 min) corresponding to the maximum slope were determined to define the time of peak growth. This peak growth time (approximately 15 to 30% of maximum OD_{600}) was used to calculate growth rate $(t_{1/2})$ as follows: normalized data were input into Prism. Next, the defined 60-min time period of exponential growth was fit to a growth equation $(Y = Y_0 e^{kt}; Y = OD_{600}, Y_0 = OD_{600}$ at time zero, k = exponential rate constant, and t = time) using nonlinear regression curve analysis to determine the exponential rate constant (k), which was then used to calculate the doubling time. Percent growth yield in sugar-supplemented versus unmodified medium was determined using normalized OD_{600} values from stationary-phase cultures, calculated as $OD_{600+sugar}/OD_{600;unmod}$ and displayed as a percentage. The average doubling time and percent growth yield were calculated from each replicate from at least three independent experiments.

Polymyxin B challenge and heat challenge on solid medium. Strains cultured overnight in C medium were used to prepare a series of six 10fold dilutions for challenge studies. For heat challenge, 5 µl of each dilution was spotted onto solidified THY medium in duplicate and incubated at either 37°C or 43°C for 48 to 72 h under anaerobic conditions. CFU were quantified, and the efficiency of plating (EOP) was calculated as $(\log_{10} CFU_{43^{\circ}C}) - (\log_{10} CFU_{37^{\circ}C})$. For polymyxin B challenge, 5 µl of each dilution was spotted onto solidified C medium unmodified or supplemented with 40 μ g ml⁻¹ (29 μ M) polymyxin B. CFU were quantified, and EOP was calculated as (log₁₀ CFU_{+PB}) - (log₁₀ CFU_{-PB}). Data shown are the means and the standard errors of the means derived from at least four independent experiments. Determination of EOP proved to be a more reproducible and sensitive metric of resistance than a standard culture-based MIC assay, as continued exposure of cultures to higher concentrations of polymyxin B tended to select for variants with higher levels of resistance. It is important to note that due to the higher lipid content from the added milk, the concentration of PB required to inhibit the growth of wild-type S. pyogenes is \sim 10-fold greater on protease indicator plates than on C medium plates.

Statistical analyses. Differences between wild-type and mutant strains were tested for significance using the Dunnett test available in InStat (GraphPad) software where an asterisk indicates P < 0.01. For all tests, the null hypothesis was rejected for P > 0.05.

RESULTS

Selection of PB-resistant mutants. Resistance to CAPs in most species of Gram-positive bacteria is associated with neutralization of the cell's highly negative surface charge, reducing its affinity for binding positively charged CAPs. Common mechanisms include lysinvlation of anionic phospholipids by MprF (31) and D-alanine esterification of lipoteichoic acid by DltA (32). However, S. pyogenes does not encode mprF, making it highly susceptible to inhibition by most CAPs, including PB (15). This property was exploited to select for mutants capable of growth at a normally inhibitory concentration of PB. We also made use of the fact that disruption of ExPortal integrity by PB is associated with inhibition of secretion of the SpeB cysteine protease, whose expression can be conveniently assessed on indicator plates. To select for spontaneous PB-resistant mutants, cells from overnight cultures of the M14 serotype S. pyogenes HSC5 were plated onto protease indicator plates containing a concentration of PB (400 μ g ml⁻¹) that prevents formation of colonies. Approximately 2×10^{10} CFU (as determined on unsupplemented medium) representing 11 independent pools was plated onto PB-protease indicator plates. Candidate mutants capable of forming discrete colonies were subjected to several rounds of restreaking to identify those with a reproducible and stable resistance phenotype. A total of 25 distinct PB-resistant mutants (designated PBr mutants) were isolated (spontaneous mutation rate of $\sim 10^{-9}$), of which 21 also displayed SpeB activity (Table 1).



FIG 1 Core PBr genes cluster within 10 gene groups. (A) Independently isolated PBr mutants contain mutations within a set of 10 core gene groups. Genes involved in guanine metabolism other than *hpt* (including *guaA*, *deoB*, *gmk*, and *nupP*) were represented by single PBr mutants and are displayed separately from *hpt* for clarity. (B) Each PBr mutant contains between 1 and 4 mutated core genes.

Mutations in a core set of 10 gene groups confer resistance to **PB.** In order to identify the genetic alterations that gave rise to PB resistance, each of the PBr mutants was subjected to whole-genome resequencing and compared to the recently completed genome of the wild-type strain HSC5 (16). Sequence alignments of the 25 mutants revealed a total of 31 single nucleotide polymorphisms (SNPs) and 24 insertion/deletions (InDels) that were identified in the open reading frames (ORFs) or putative RBS of a predicted gene. Individual PBr mutants contained between 1 and 4 mutations (average 2.2 mutations/isolate [Table 1; see also Tables S6 and S11 in the supplemental material]). SNPs displayed a bias toward transitions over transversions (17 and 14, respectively), similar to what has been reported in other organisms (33). However, 30/31 SNPs resulted in mutation to either A or T, reflecting the low GC content of S. pyogenes (38.5%) (see Table S7). The majority of InDels consisted of single nucleotide deletions in predicted ORFs resulting in frameshifts, although a few contained larger deletions of up to 167 bp (see Table S8). Insertions were rare and were limited solely to 1-bp insertions (see Table S8). The majority of InDels occurred from loss of repetitive elements, particularly in homopolymeric stretches of adenines, although loss of larger and more complex repetitive elements also occurred (see Table S9). SNPs caused a variety of genetic lesions, including nonsynonymous mutations and premature stop codons, as well as the loss of stop codons that generated larger fusion proteins. In 3 independent cases, a single mutation affected the same 2 overlapping genes, *tilS* and *hpt*, as the final 3 codons in the 3' end of *tilS* overlap the putative RBS of *hpt* (PBr 6.10, 10.15, and 11.2 [Table 1]). Of the 55 total mutations identified, only one mutation occurred outside a predicted ORF or RBS (PBr 10.6, which contains 2 other SNPs) and only one PBr mutant contained a synonymous mutation (PBr 10.11, which also has an SNP and an InDel) (Table 1), suggesting that mutations without functional consequences were rarely associated with mutations that promoted resistance. Surprisingly, the majority of mutations were localized within a set of only 10 gene groups. These "core PBr genes" (defined as individual genes or sets of genes with closely related annotations that

were mutated in >1 PBr mutant) accounted for 49 of the 55 total SNPs and InDels identified (Fig. 1A). Each PBr mutant contained between 1 and 4 core PBr mutations (Fig. 1B). Furthermore, a single mutation in any one of a set of 4 core PBr genes conferred resistance in the absence of any other mutation, including *pstS* (PBr 1.1 and 2.1), *clpX* (PBr 4.1), *manN* (PBr 6.3), and *tilS/hpt* (PBr 10.15) (Table 1).

Construction of mutations in individual PBr genes. Given the rich diversity of mutations, resistance may have developed from loss- or gain-of-function mutations or from the combinatorial effect of multiple mutations. To gain further insight into the role of each core PBr gene, an attempt was made to construct individual loss-of-function mutations in the wild-type background. An initial attempt was made to construct in-frame deletions within each gene group, and this strategy yielded mutations in 5 PBr gene groups (gdpP, pstS, manLMN, ebsA, and clpX [Table 2]). Since recovery of a mutant using this method requires that it compete successfully with the wild type (18), an alternative replacement strategy to place a promoterless chloramphenicol resistance gene under the control of the native gene's promoter was employed. If the promoter is of sufficient strength, chloramphenicol selection will promote recovery of the null mutant. This approach was successful for an additional gene (*ptsI* [Table 2]). Mutations in *hpt* and *ftsH* were successfully constructed by a one-step duplication-inactivation method (18) that allows for direct selection for insertion of a spectinomycin resistance determinant (Table 2). For fabT, it was noted that all spontaneous PBr fabT mutations were also associated with mutations of either *gdpP* or *clpX* (PBr 3.5, 3.6, 9.21, 10.6, 10.11, and 11.1 [Table 1]), suggesting that the latter mutations may compensate for a deleterious effect of loss of *fabT* on growth rate or viability. This idea was then supported by our ability to generate an in-frame deletion in *fabT* in

 TABLE 2 Characterization of mutants with deletion or disruption of PBr core genes

		Growth rate	DB		SpeB secretion ^e		
Mutant	Strain ^a	(% yield) ^b	binding ^c	$ExPortal^d$	-PB	+PB	
WT	HSC5	50.1 (100)	+	Disrupted	+	-	
$\Delta ClpX$	GCP688	70.1 (97)	+	Intact	-	-	
ΔGdpP	GCP751	51.9 (119)	+	Intact	+/-	+/-	
$\Delta PstS$	GCP754	58.0 (99)	+	Intact	+	+	
Δ GdpP- Δ FabT	GCP766	66.0 (76)	+	Intact	+/-	+/-	
$\Delta ClpX-\Delta FabT$	GCP767	70.9 (87)	+	Intact	-	-	
Δ ManLMN	GCP771	51.7 (109)	+	Heterogeneous	+	-	
ΔEbsA	GCP784	53.4 (99)	+	Heterogeneous	+	-	
$\Delta PtsI::cat$	GCP953	68.5 (80)	+	Intact	+	+	
ΩHpt	GCP859	84.8 (68)	+	Intact	+	+	
ΩFtsH	GCP862	88.5 (67)	+	Intact	+/-	-	
ΩCon	GCP292	59.2 (96)	+	Disrupted	+	-	

^a Name of strain with the indicated mutant phenotype. See Table S1 in the

supplemental material.

^b Doubling time $(t_{1/2})$ in minutes in C medium calculated from peak growth at midexponential phase between 15 and 30% of maximum growth yield. Yield is percentage of WT value based on final OD₆₀₀.

 c Absence (–) or focal pattern (+) of binding when stained with 5 μM dansyl-PB. d Staining with 30 μM dansyl-PB results in a focal (intact) or dispersed (disrupted) pattern of binding or a mixture of cells with an intact or dispersed (heterogeneous) pattern of binding.

^{*e*} Level of secreted SpeB cysteine protease in culture supernatant detected by Western blotting following challenge during the late logarithmic phase of growth with (+PB) or without (-PB) 30 μ M PB and harvested 2 h later after the onset of stationary phase: +, \geq 80%; +/-, 50 to 80%; -, \leq 10%, all compared to WT. the context of deletion of either *gdpP* or *clpX* (Table 2) but not in wild-type cells. However, a similar strategy to generate a *topA* mutation in the context of deletion of *pstS* was not successful. Finally, deletion of *tilS* was not attempted since the mutations in the original mutants likely confer PB resistance by alteration of the *hpt* RBS (see below). Analysis of growth rates and yields of the resulting mutants was mostly consistent with the method required to generate each mutant. In-frame deletion mutants had either a growth rate or yield (or both) near that of the wild type, while those requiring selection (Δ PtsI::*cat*, Ω Hpt, and Ω FtsH) had both a longer growth time and a lower growth yield (Table 2).

PB resistance primarily arises through loss of function of core PBr genes. To gain insight into the contribution of individual core genes to the PB resistance phenotype, the panel of null mutants was tested for sensitivity to PB challenge. Using a quantitative efficiency-of-plating assay, both the wild-type (WT) strain and a previously described control strain with an intergenic insertion downstream of a gene not associated with PB resistance $(\Omega \text{Con} [24])$ demonstrated over 5 logs of sensitivity to PB (Fig. 2A). In contrast, the various core PBr mutants were highly resistant and typically had plating efficiencies on the order of 4 to 5 logs higher than that of the wild type (Fig. 2A). The exceptions were Δ ManLMN and Ω FtsH, which were as sensitive as the WT and Ω Con strains (Fig. 2A). To test the contribution of FabT toward PB resistance, Δ GdpP- Δ FabT and Δ ClpX- Δ FabT double mutants were complemented with plasmids expressing either GdpP or ClpX. Although empty vector (pABG5) had no effect on PB resistance in any strain, pGdpP and pClpX were able to restore PB sensitivity to wild-type levels in their respective single deletion mutant strains (Fig. 2B). In contrast, these plasmids had no effect on PB resistance for either Δ GdpP- Δ FabT or Δ ClpX- Δ FabT (Fig. 2B), demonstrating that loss of FabT alone is sufficient to confer PB resistance. PB binding and enhanced ExPortal integrity were then assessed by challenge of mid-logarithmic-growth-phase cultures with fluorescently labeled PB at a low concentration that results in a focal pattern of binding and at a high but still sublethal concentration that results in a dispersed pattern of binding in the WT strain. At low concentrations, dansyl-PB bound in a focal pattern to all mutants (Table 2). However, when examined at a concentration that disrupts the ExPortal in the WT strain, all mutants demonstrated enhanced ExPortal integrity. Of the mutants, 5 gained the ability to secrete the SpeB protease at concentrations that inhibit its expression in the wild type or Ω Con (Table 2), although those with mutations in gdpP or ftsH had an overall reduction in SpeB expression under all conditions. In contrast, those mutants lacking ClpX function (Δ ClpX and Δ ClpX- Δ FabT) failed to express SpeB even in the absence of PB (Table 2). Thus, while most spontaneous mutants have mutations in multiple genes, PB resistance and enhanced ExPortal integrity can be recapitulated by the loss of function in almost all core PBr genes. Exceptions are a lack of resistance for *ftsH* and *manLMN* null mutants. However, spontaneous ftsH mutations were always found in the context of *pstS* or *pstC* mutations (Table 1) and spontaneous manLMN missense mutations were found in isolation (e.g., PBr 6.3 [Table 1]). Taken together, these data indicate that the PBr phenotype for the latter mutations likely represents a gain of function, while the former may contribute to a combinatorial effect.

Altered expression of the Fab operon in PBr fabT mutants. In streptococci, the expression of the genes encoding enzymes involved in fatty acid biosynthesis is under the control of the tran-



FIG 2 Most core PBr genes confer resistance through loss of function. Polymyxin B resistance of deletion or insertional disruption mutants (A) and selected mutants transformed with plasmids expressing GdpP (pGdpP), ClpX (pClpX), or empty pABG5 vector (Vector) (B) was determined by efficiency of plating on solid medium. Stationary-phase cultures of indicated strains were serially diluted and plated onto solidified C medium either unmodified or containing 40 μ g ml⁻¹ polymyxin B. Efficiency of plating was calculated as log₁₀ CFU_{+PB}/log₁₀ CFU_{unmod}. Data shown are the means and the standard errors of the means derived from at least four independent experiments. Differences between wild-type and mutant strains were tested for significance using the Dunnett test (*, *P* < 0.01).

scriptional regulator FabT (34). Since the ExPortal has an asymmetric content of anionic phospholipids (10), any PBr genes affecting lipid biosynthesis are of particular interest. In addition, the fact that *fabT* mutations were common among PBr mutants (Table 1) but that it was possible to construct null *fabT* mutations only in the context of other mutations complicates understanding how mutation of *fabT* may contribute to PB resistance. Similar to Streptococcus pneumoniae (35), the genes for fatty acid biosynthesis are clustered together in S. pyogenes (Fab operon). The two operons are nearly identical in sequence, although several of the FabT binding sites differ in location relative to their promoters and the S. pyogenes sites lack the sequence diversity of S. pneumoniae (Fig. 3A). Consistent with its function as a repressor of transcription (34), deletion of *fabT* resulted in a 2- to 8-fold increase in transcription of the genes adjacent to each of the 4 FabT sites when monitored by real-time RT-PCR (Fig. 3B). This increase was strictly due to the loss of fabT, as mutation of the companion genes alone (Δ ClpX and Δ GdpP) did not result in a sig-



FIG 3 Mutation of the FabT transcriptional repressor results in upregulation of fatty acid biosynthetic genes but not anionic lipid genes. (A) The fatty acid biosynthesis (*fab*) region carried within *S. pyogenes* contains 3 predicted operons (thin black arrows) and 4 putative FabT binding sites upstream (black boxes) or within *fab* genes (hollow arrows). Genes interrogated for mRNA expression levels are shaded in gray. Predicted FabT binding sites are shown in comparison with *S. pneumoniae*, and nucleotide distances in relation to translational start codons (ATG or TTG) are indicated in parentheses. Distances between FabT binding sites and predicted -35 and -10 promoter elements and the ribosome binding site (RBS) are indicated in the diagram. The consensus 13-nucleotide palindromic FabT binding site is shown (Con), and deviations are indicated in lowercase. (B and C) The relative level of message of select genes in deletion mutants (B) and PBr mutants containing FabT SNPs (C) was determined using real-time RT-PCR and normalized to the wild-type strain. Data shown are the means and the standard errors of the means derived from triplicate determinations of samples from at least three independent experiments. The dashed lines indicate >2-fold or <2-fold changes. An asterisk indicates an undetectable level due to the absence of the appropriate gene in the deletion mutant.

nificant increase in expression of any Fab operon gene (Δ ClpX and Δ GdpP [Fig. 3A]). Fatty acid biosynthesis also produces precursors required for synthesis of anionic phospholipids. However, the loss of *fabT* did not impact expression of genes required for phosphatidylglycerol (*pgsA*) or cardiolipin (*cls*) synthesis (Fig. 3B). Interestingly, although *fabT* mutations were present in approximately one-fourth of the PBr isolates, they consisted entirely of missense mutations or alterations to the RBS; there were no frameshift, nonsense, or InDel mutations. This limited distribution of mutation type was in contrast to the collection of mutations obtained in other core genes like clpX and pstS (see Table S10 in the supplemental material). While this suggests that PB resistance required a gain of function in FabT, this was not supported by analysis of gene expression, which indicated that all fabT PBr mutations derepressed expression of Fab genes to levels nearly approaching those observed for fabT deletion mutants (Fig. 3C). The contribution of fabT alteration to the PBr phenotype is discussed in more detail below.



FIG 4 Loss of ClpX or Hpt confers heat resistance. Stationary-phase cultures were serially diluted, plated onto solid medium, and incubated at permissive (37°C) or restrictive (43°C) temperature. After 48 h, CFU were quantified and efficiency of plating was calculated as $\log_{10} \text{CFU}_{43^\circ}/\log_{10} \text{CFU}_{37^\circ}$. Data shown are the means and the standard errors of the means derived from at least four independent experiments. Differences between wild-type and mutant strains were tested for significance using the Dunnett test (*, P < 0.01).

Several core PBr mutants exhibit enhanced multistress resistance. In other bacterial species, mutations in several of the core PBr genes have been implicated in resistance to multiple stresses, including heat (36) and acid (37). To investigate whether the S. pyogenes core PBr mutations can confer resistance to additional stresses, the panel of PBr mutants was tested for resistance to heat stress. The WT strain used here (HSC5) is typical of most S. pyogenes strains and cannot grow at temperatures over 39°C. When examined quantitatively, the WT strain was unable to grow at an elevated temperature and had an efficiency of plating below the limit of detection for the assay (over 5 logs of sensitivity) when tested at 43°C versus 37°C (Fig. 4). In contrast, several of the core PBr mutants, including $\Delta ClpX$, $\Delta ClpX/\Delta FabT$, and ΩHpt , gained the ability to grow at this high temperature and had plating efficiencies that were reduced less than 1 log (Fig. 4). Other core mutants remained heat sensitive (Fig. 4). Examination of several spontaneous PBr isolates with mutations in *clpX* (PBr 4.1, 9.21, 10.11, and 11.1) or hpt (PBr 6.7, 6.10, 10.15, and 11.2) revealed that they were similarly heat resistant (data not shown). Since hpt functions in guanine metabolism, other PBr isolates with mutations in genes involved in guanine metabolism were tested, including deoB (PBr 4.5) and guaA (PBr 10.14), and these were also heat resistant, as was PBr 11.5 with mutations in fba and manL (data not shown). Thus, some mutations that promote PB resistance, most notably in *clpX* and in guanine metabolism genes, also can confer resistance to other types of stress.

A subset of PBr mutants display altered carbohydrate metabolism. Many core PBr genes encode putative transporters of nutrients, suggesting that mutation of these genes may confer a starvation state that is associated with PB resistance (Table 1). To test this, the WT strain and selected mutants were cultured in a carbohydrate-poor medium (C medium) that was supplemented with various sugars to determine if PB resistance is associated with defects in carbohydrate utilization. The sugars tested were selected on the basis of the predicted repertory of *S. pyogenes* HSC5 carbohydrate transporters (see Table S12 in the supplemental material), and it was considered that a sugar could be utilized if supplementation resulted in at least a 20% increase in growth yield. A selection of sugars not predicted to be utilized by *S. pyogenes* was included for comparison. Generally consistent with predictions, the



FIG 5 Some PBr mutants show altered sugar utilization. (A) Wild-type or deletion mutant strains were tested for sugar utilization by supplementing C medium with various sugars that HSC5 was either predicted (+) or not (-) to utilize based on genome annotation. (B) PBr mutants containing SNPs in *ptsI* or *manL/N* were tested for sugar utilization with select sugars in comparison to deletion mutants. Percent yield was calculated as $OD_{600+sugar}/OD_{600:unmod}$. Data shown are from triplicate determinations of samples from at least three independent experiments.

addition of 14 different sugars increased growth yields (Fig. 5A). Similarly to a recently described ptsI deletion mutant of S. pyogenes (38), Δ PtsI lost the ability to utilize 9 of these 14 sugars (Fig. 5A). Its growth was enhanced by glucose, the amino sugar Nacetylglucosamine (GlcNAc), and multimeric forms of glucose linked by $\alpha(1\rightarrow 4)$ glycosidic bonds (maltose, maltotriose, and maltodextrin) but not α, α -(1 \rightarrow 1) and β (1 \rightarrow 4) glycosidic bonds (disaccharides trehalose and cellobiose, respectively) (Fig. 5A). Since this mutant lacks a key component of the sugar phosphotransferase system (PTS), these data define the classes of PTSdependent and -independent sugars for S. pyogenes. However, growth profiles differed markedly for spontaneous ptsI PBr mutants. These mutants had no obvious defect in utilization of representative PTS (mannose and fructose) and non-PTS (glucose) sugars (PBr 11.3 and 11.6 [Fig. 5B]). Since both of these have missense mutations (Table 1), their contribution to PB resistance must not arise from a loss of function.

Another series of carbohydrate-related core PBr genes are those of the *manLMN* operon putatively associated with transport of mannose (Table 1). However, the PBr phenotype was not recapitulated in the loss-of-function Δ ManLMN mutant (see above). Thus, it was of interest to compare profiles of carbohydrate metabolism in this group of mutants. The Δ ManLMN mutant was able to utilize most of the same sugars as those utilized by the WT strain with the primary exception being that it was much less efficient in its ability to utilize mannose, suggesting that while Man-LMN is not the only mannose transporter, it makes a major contribution to mannose metabolism in *S. pyogenes* (Fig. 5A). This phenotype differed for all spontaneous PBr Man operon mutants. The *manN* mutants had only a slight defect in mannose utilization, while the *manL* mutants had no defect and achieved en-

TABLE 3 Growth rates in presence and absence of salts

	Growth rate (% yield) ^a
$\operatorname{Salt}(s)^b$	WT	$\Delta PstS^{c}$
None	42.0	ND
K ₂ HPO ₄ , MgSO ₄ , NaCl	53.8	57.2
K ₂ HPO ₄	58.3	99.5 (81)
MgSO ₄	49.8	ND
NaCl	47.2	68.6
K ₂ HPO ₄ , MgSO ₄	56.8	85.4 (81)
K ₂ HPO ₄ , NaCl	54.0	55.4
MgSO ₄ , NaCl	46.4	75.8
Na ₂ HPO ₄	57.5	59.2
KCl	41.1	ND
MgSO ₄ , KCl	45.7	ND

 a Growth rate $(t_{1/2})$ was measured in salt-reduced C medium supplemented with indicated salts.

 b Salts were added to salt-reduced C medium as described in Materials and Methods. c Final yields for $\Delta PstS$ reached approximately 100% of the WT value except for those indicated in parentheses or labeled ND (none detected), which were ${<}5\%$ of the WT value.

hanced growth yields equivalent to those of the WT (Fig. 5B). Since there was no direct correlation with a loss of mannose metabolism, this indicates that the PBr phenotype is a gain of function associated with the mannose membrane transporter complex.

A subset of PBr mutants display altered salt metabolism. Another nutrient acquisition pathway associated with PB resistance involves the uptake of inorganic phosphate (P_i). Mutations in a putative high-affinity P_i ABC-type transporter encoded by the Pst operon were the most commonly isolated mutations, appearing in 40% of the total number of PBr mutants isolated (Fig. 1A). The most common lesion was a frameshift in *pstS* (Table 1), which encodes the extracellular substrate-binding lipoprotein component of the transporter. Independent mutants were also isolated with mutations in *pstA* (frameshift) and *pstC* (nonsense) (Table 1), which encode membrane permease components. To test if these mutations resulted in alterations of salt metabolism that may correlate with PB resistance, the growth characteristics of WT and the various mutants were compared in media with an altered content of salts. In unaltered C medium, the Δ PstS mutant had a growth rate and yield equivalent to those of the WT (Table 2). In C medium prepared without its normal complement of salts (10 mM K₂HPO₄, 0.4 mM MgSO₄, 17 mM NaCl), the growth rate of the WT strain was unaltered; however, $\Delta PstS$ failed to grow (none detected [ND] [Table 3]). Although MgSO₄ had no effect on growth, addition of NaCl alone was sufficient to restore $\Delta PstS$

growth at a slightly reduced rate, while the addition of K₂HPO₄ by itself also restored growth, although at a substantially reduced rate and growth yield (Table 3). Furthermore, the addition of both K_2 HPO₄ and NaCl restored growth of Δ PstS to levels equivalent to all 3 salts (Table 3). Finally, swapping cations by adding KCl did not restore growth; however, the addition of Na₂HPO₄ restored growth to a rate and yield equivalent to those for complete medium (Table 3). Thus, full growth of Δ PstS was restored only either by the presence of K₂HPO₄ plus NaCl or by Na₂HPO₄, indicating a requirement for both Na⁺ and P_i. This same requirement for salts was observed for all spontaneous PBr PstS/A/Ccontaining mutants (Table 4), although pstS frameshift mutants did show a low level of growth following an extended lag period of several hours (data not shown). Taken together, this analysis indicates that an altered salt metabolism may lead to the acquisition of PB resistance.

Growth in low-salt medium selects for pstS revertants. Overnight cultures of the various PBr mutants containing the pstS frameshift mutation would occasionally demonstrate reduced but detectable growth in C medium lacking salts. To determine if this was due to reversion, 10 independent cultures from each of the 8 pstS-containing mutants were prepared in reduced-salt C medium and incubated for an extended time period. When examined after 48 h, a wild-type growth yield was obtained for every culture of each mutant (Table 5). When growth from these cultures was tested again in reduced-salt C medium, they now demonstrated a growth rate indistinguishable from that of the wild type (Table 5). DNA sequence analyses of each of these cultures revealed reversion to the wild-type *pstS* sequence (see Table S2 in the supplemental material). Furthermore, reversion was specific to *pstS*, as each individual revertant retained all other associated mutations corresponding to their parental PBr strain. In contrast, salt-revertant mutants were rarely obtained in strains with mutations in *pstC* or *pstA* and were never recovered from Δ PstS (Table 5). DNA sequence analysis revealed that the one revertant derived from PBr 10.9 gained a single nucleotide insertion at the site of the original 19-bp deletion within *pstA*, thus resulting in an 18-bp in-frame deletion and generating a PstA open reading frame lacking 6 amino acids, while the one revertant of PBr 3.2 contained an SNP within *pstC* reverting the nonsense mutation (premature stop codon) to a missense mutation (altered amino acid), thus restoring the PstC open reading frame (see Table S2). The *pstS* frameshift mutation (fs-aa6 [Table 1]) occurs at the terminal position of a track of 9 consecutive adenine residues. As will be discussed below, the frequency with which this mutation arises and reverts and its robust PB-resistant phenotype suggest that this homopoly-

TABLE 4 Growth rates for salt-dependent PstS, PstC, and PstA PBr mutants

Salts	Growt	Growth rate ^a										
		PstS mutant ^b :								PstC mutant	PstA mutant	
	WT	Δ	1.1	2.1	3.1	3.3	3.7	4.4	4.5	10.14	3.2^b	10.9 ^b
None ^c	46.1	ND	99.4	127.0	173.1	133.0	158.8	104.0	186.3	177.6	200.3	ND
K ₂ HPO ₄ , MgSO ₄ , NaCl	51.9	58.7	54.0	55.6	58.2	55.5	52.9	52.5	65.3	56.8	59.7	59.0

 \overline{a} Growth rate ($t_{1/2}$) was measured in salt-reduced C medium supplemented with indicated salts.

^b PBr mutants containing mutations within *pstS*, *pstC*, and *pstA* are described in Table 1 and in Table S6 in the supplemental material. ND, none detected.

^c Final yields for PBr mutants containing SNPs in *pstS* or *pstC* reached approximately 60% of the WT value after 24 h of culture in the absence of salts and 100% in the presence of salts.

TABLE 5 Auxotrophic P_{i} revertants recovered following 48 h of culture in salt-reduced C medium

Mutant ^a	$\operatorname{Growth}^{b}$	Rate ^c
PstS		
Δ	0/10	ND
1.1	10/10	46.3
2.1	10/10	49.0
3.1	10/10	45.3
3.3	10/10	48.0
3.7	10/10	45.7
4.4	10/10	53.8
4.5	10/10	48.7
10.14	10/10	48.6
PstC mutant 3.2	1/10	46.3
PstA mutant 10.9	1/10	50.2

^{*a*} PBr mutants containing mutations within *pstS*, *pstC*, and *pstA* are described in Table 1 and in Table S6 in the supplemental material. Pst revertant mutants are described in detail in Table S2.

 b PBr mutants were subcultured 1:1,000 from C medium into salt-reduced C medium. Ability to grow in salt-reduced C medium was scored in each of 10 replicate culture tubes after 24 h by measurement of OD₆₀₀.

 c Growth rate $(t_{1/2})$ of revertant mutants was measured in salt-reduced C medium. ND, none detected.

meric track functions as a molecular switch to facilitate adaptation to stressful environments.

Core PBr genes include novel regulators of SpeB expression. As discussed above, the processing of the secreted SpeB cysteine protease zymogen to its intact form depends upon an intact Ex-Portal. However, several of the mutants with enhanced ExPortal integrity had a defect in SpeB expression when examined on protease indicator plates (Table 1). Furthermore, several of the core PBr deletion mutants had decreased levels of the SpeB polypeptide in culture supernatant when examined at a time point when expression in the WT strain is maximal (Table 2). Expression of SpeB is highly regulated by factors that act at multiple levels, including transcription, secretion, and postsecretory folding and maturation (for a review, see reference 39). For example, we have previously reported that a mutant with an insertional disruption of the core PBr gene gdpP has a kinetic defect in the processing of 40-kDa SpeB zymogen to its 27-kDa active form, even though it expresses normal levels of the speB transcript and is not defective for secretion (30). To gain insight into how other core PBr genes affect SpeB expression, the panel of SpeB-defective mutants was compared to the gdpP deletion mutant (Δ GdpP) mutant constructed for this study. Examination of Ω FtsH revealed that it had an even greater kinetic defect than Δ GdpP when both are compared to WT and to a mutant with an accelerated pattern of expression (ΔV fr) that results from uncoupling *speB* transcription from its normal mode of growth phase control (Fig. 6A; see also Fig. S1 in the supplemental material). In contrast, Δ ClpX was completely deficient in expression of SpeB and more closely resembled a strain that expresses a catalytically inactive SpeB protein (SpeB_{C192S}) or one that lacks RopB (Δ RopB), the transcriptional activator required for SpeB expression (Fig. 6A; see also Fig. S1). Examination of speB transcript levels by real-time RT-PCR revealed a significant defect at the level of *speB* transcription for the Δ ClpX mutant, and to a lesser extent, the Ω FtsH mutant (Fig. 6B). In contrast, transcript levels were unaltered in Δ GdpP, indicating a posttranscriptional defect (Fig. 6B). Taken together, these data

indicate that several core PBr mutants (Ω FtsH, Δ ClpX, and Δ GdpP) have SpeB defects at the transcriptional and posttranscriptional level that are independent of ExPortal integrity.

DISCUSSION

It has only recently been appreciated that CAPs can have significant biological effects at sublethal concentrations, including an ability to alter patterns of protein secretion by targeting and disrupting the anionic phospholipid-enriched microdomain of the ExPortal (15). In the present study, we have used this property of CAPs to design a genetic screen to gain insight into the mechanism(s) by which *S. pyogenes* organizes and maintains the ExPortal. Analysis of the resulting collection of mutants implicated a role for an assortment of genes whose loss or gain of function influenced ExPortal organization, including those involved in nutrient acquisition, stress responses, and virulence gene regulation.

Since *S. pyogenes* exclusively relies upon the Sec system for secretion of virulence factors, understanding how the ExPortal is maintained and organized is of central importance for understanding how the Sec pathway has been adapted to support the secretion and processing of proteins which have numerous roles in pathogenesis. The observation that disruption of the anionic phospholipid-enriched microdomain leads to the redistribution of ExPortal components (15, 40) indicates that a key question involves the mechanism that clusters these lipids into a focal microdomain. At the concentrations found in physiological membranes, various lipid species only transiently associate into



FIG 6 Several core PBr genes regulate SpeB expression. (A) SpeB protease activity was determined for various strains by plating serial dilutions onto protease indicator plates and monitoring for protease activity over time. Activity data shown are the means and the standard errors of the means derived from triplicate determinations of samples and are representative of at least three independent experiments. (B) The relative level of *speB* transcript over the course of growth was determined using real-time RT-PCR. RNA data shown are the means and the standard errors of the means derived from triplicate determinations of samples from at least three independent experiments.

microdomains. The stabilization of these microdomains then becomes dependent on a number of different organizing factors. In eukaryotic cells, numerous proteins have been described that function to extend the half-lives of lipid microdomains in order to promote specific functionality, which typically involves endocytosis or cell signaling (41). In rod-shaped bacteria, the stable segregation of anionic phospholipids into functional helix-like domains is organized via interaction with the cytoskeletal protein MreB and its various homologs (11, 42–44). However, S. pyogenes, like all other streptococcal species, lacks any MreB-like protein (9). Similarly, how CAPs can disrupt these normally stable microdomains in the absence of significant membrane damage is also not understood. In model membranes, rather than dispersing anionic phospholipids, challenge with CAPs tends to promote the transition of anionic lipids from a liquid-disordered state to a more clustered liquid-ordered configuration (45). Thus, further analysis of how the various mutations identified in the current study can promote enhanced ExPortal integrity will provide fundamental insights into the mechanisms that support the segregation of lipids into functional domains.

Analysis of this collection of mutants should also contribute to our understanding of mechanisms of resistance to CAPs. Many bacterial species possess factors that confer CAP resistance that most commonly act by modification of negative surface charges to abrogate CAP binding. These include MprF, which lysinylates anionic phospholipids (31), and DltA, which acts via D-alanine esterification of teichoic acids. However, S. pyogenes does not encode MprF, and loss of DltA results in increased susceptibility to CAPs (32). Alteration of surface charge also did not appear to be a common mechanism among the PB-resistant mutants isolated in the current study, as their PB-binding properties were unaltered. Another common mechanism of resistance to antibiotics that target anionic phospholipids involves alterations in the biosynthesis pathways for these lipids. For example, resistance to daptomycin can result from mutations that alter residues in the enzyme Cls or PgsA or from mutations that affect their transcriptional regulators (46-51). In the current study, no mutations were identified in the anionic phospholipid biosynthetic genes or their presumed regulators. However, multiple PBr isolates were found to have a mutation in *fabT*, which encodes the master regulator of fatty acid biosynthesis in streptococci (52). The frequency and diversity of fabT mutations suggest that alterations to membrane composition may contribute to resistance.

An understanding of the role of *fabT* mutations in the PB resistance phenotype was complicated by the fact that null mutations could be recreated only in the context of mutations in either clpX or gdpP. However, both comutations could be restored through expression off a plasmid in the *fabT* deletion strains, demonstrating that loss of *fabT* was sufficient for resistance to PB. Although it is unclear why these particular comutations facilitate the genetic manipulation of *fabT*, it has been noted that the generation of fabT deletions in Streptococcus pneumoniae is notoriously difficult (34), suggesting that the comutations may restore some measure of fatty acid homeostasis. However, since the comutations by themselves did not alter transcription of the fatty acid biosynthesis operon, some other compensatory mechanism must be involved. The S. pneumoniae fabT mutants display a 2- to 4-fold increase in transcription of the fatty acid biosynthesis operon and an increase in the ratio of saturated to unsaturated fatty acids, as well as a predominance of longer fatty acid chains

(34). Alteration to pools of anionic phospholipids also may result since these fatty acids are precursors in their biosynthesis. This modified membrane composition may both contribute to enhanced ExPortal integrity and trigger a membrane stress response to a deleterious level. A damping of this may explain the compensatory function of the comutations, as ClpX and GdpP have both been associated with regulation of stress responses (37, 53–55). Interestingly, two independent PBr mutations altered the same serine residue at position 84 of FabT (S84L), a potential target of the Stk/Stp serine/threonine kinase/phosphatase of *S. pyogenes* (56, 57). Stk regulation of FabT alters membrane fatty acid composition (56) and contributes to resistance to several types of stress (57), further implicating a link between FabT, stress response, and ExPortal integrity.

An involvement in stress response may be a property shared by many of the other genes identified as associated with PB resistance. For example, transposon mutagenesis screens designed to isolate Lactococcus lactis mutants resistant to heat (36) or acid stress (37) yielded resistant mutants containing transposon insertions in many of the same genes that were identified here, including core PBr genes *pstS*, *hpt*, and *gdpP*. Also identified among the genes that were represented only once among the current collection of mutants were those involved in guanine metabolism, including *deoB* and *guaA*. A general characteristic of mutants with these insertionally inactivated genes is that they are resistant to a variety of different stresses, suggesting that their loss of function results in the activation of a general stress response pathway(s). Several global regulatory pathways that link metabolic stress to guanine metabolism have been identified, including (p)ppGpp and the stringent response (37) and the transcription regulator CodY (58). Other mutants represent pathways that link other aspects of nucleotide metabolism to general stress responses, including the second messenger cyclic-di-AMP phosphodiesterase GdpP (55, 59, 60), which has recently been linked to potassium import under osmotic stress (61-63). If triggering a stress response is a characteristic shared by other mutations in this collection, then it is possible that those involving the loss of function of a single subunit of a multicomponent membrane transporter may cause a membrane perturbation that induces a stress response. However, the collection also includes examples of membrane transporter mutations that were a gain of function, such as the manL/N mutants that conferred PB resistance while retaining their ability to transport mannose. In this case, it is possible that these mutations had a more peripheral effect that induced a stress response, including altering how the transporter interacts with other membrane proteins or causing it to mislocalize or cluster at an inappropriate location in the membrane.

Similar to the mannose transporter, mutations conferring PB resistance that were apparently the result of a gain of function occurred in FtsH, a membrane-bound protease that degrades SecY when it becomes uncoupled from other Sec translocon components (64), or upon protein secretion stress which can occur when the Sec translocon becomes jammed (65). As PB causes the redistribution of the Sec translocons and may also impart significant secretion stress, these *ftsH* mutations may result in the stabilization of SecY. Interestingly, both spontaneous *ftsH* mutations localized to a specific region of the gene, immediately downstream of the region encoding its Walker B motif. Combined with the observation that the FtsH null mutant was not PB resistant, the localization of these mutations to a specific motif may suggest that

one of FtsH's other known regulatory functions (66) contributes to PB resistance.

An unusual core PB-resistant gene identified was the EbsA gene. Although deletion of EbsA was sufficient to confer PB resistance, the Δ EbsA mutant displayed disrupted ExPortal staining and was unable to secrete SpeB following PB challenge. Spontaneous EbsA mutants were identified twice in the screen (PBr 3.1 and 11.1), both of which maintained nearly full-length ORFs (I95N and $\Delta 21-25$, respectively), and were both isolated in the context of several additional comutations. Therefore, either the spontaneous EbsA mutants may impart a gain of function to EbsA or the comutations may impart a synergistic effect with respect to ExPortal integrity, as PBr 3.1 maintained SpeB expression upon PB challenge. EbsA was originally identified in *Enterococcus faecalis* as being required for expression of enterococcal binding substance and necessary pheromone-inducible plasmid conjugation (67). Although the mechanism is still unclear, EbsA has homology to pore-forming proteins associated with phage endolysins and is therefore predicted to affect cell wall metabolism. The recent association of peptidoglycan metabolism and ExPortal organization supports this model (40).

From the class of null mutations in membrane transporters, perhaps the most interesting case involves the Pst P; uptake transporter. Although the genes encoding three of its five components were represented in the collection, pstS was the most commonly mutated gene and was found altered in 8 of the 25 PBr mutants. Significantly, each isolate possessed the exact same mutation even when isolated from multiple independent experiments, a frameshift in the codon for the lysine residue at position number 6. This repetitively occurring mutation was most likely due to the fact that pstS carries a homopolymeric stretch of 9 consecutive adenines encoding 3 lysine residues which may be subject to slipped-strand mispairing during DNA replication (68). The AT-rich S. pyogenes HSC5 genome (38.5% GC content) contains 25 homopolymer A or T tracks of \geq 9 bp. However, the frequency of mutation at this homopolymer track far surpassed that of any other locus in the PBr isolates (see Table S9 in the supplemental material), demonstrating a significant association between slippage at this locus and PB resistance. As expected for a slippage mechanism, wild-type pstS revertant mutants were readily isolated at high frequency under selective pressure. Thus, strand slipping at this locus may represent a form of phase variation whereby S. pyogenes can transiently enter a P_i-starved state to induce a PB resistance response and then revert to wild type when the selective pressure is removed. Additional evidence to support this hypothesis comes from the observation that despite the availability of two codons encoding lysine (AAA and AAG), the homopolymeric 9-adenine track is conserved in 14 out of 17 of the available sequenced S. pyogenes genomes, while the remaining three contain an 8-adenine track (AAA-AAA-AAG), in which the guanine residue is replaced at the terminal adenine position. In contrast, S. mutans also contains a polylysine track in *pstS* but circumvents slippage with the sequence AAA-AAG-AAA-AAA. The ability of Na⁺ to restore growth of the $\Delta PstS$ mutant suggests that the NptA Na⁺/P_i cotransporter (L897_06955) may sustain viability in the absence of PstS, except under conditions of extreme P_i limitation. Since the pst mutations are all loss-of-function mutations, they could confer PB resistance either through the inappropriate assembly of the transporter's remaining components or through an effect on phosphate metabolism. For lactic acid bacteria, alterations of phosphate metabolism have been implicated in regulation of central metabolism and stress resistance (69, 70).

In addition to potential roles in regulation of stress responses, several of the core PBr genes were found to have a role in regulation of the SpeB cysteine protease. This protease can account for >95% of the total protein found in the culture supernatant (71). As a relatively nonspecific protease, it was anticipated that SpeB would be required for PB resistance, as has been reported for other CAPs (72). However, several of the PB-resistant mutants had a defect in SpeB expression associated with mutations in gdpP and *clpX*. It has previously been reported that inactivation of *gdpP* is associated with decreased SpeB activity associated with a kinetic delay in processing the SpeB zymogen to its active form (30, 59). The recently described second messenger molecule cyclic-di-AMP is the substrate of the GdpP phosphodiesterase and has been linked to regulation of cell wall and cell envelope homeostasis (55, 59, 73–75), suggesting an overlap of these activities with regulation of SpeB expression. In addition, DarR, a recently identified cyclic-di-AMP-responsive transcription regulator, controls expression of fatty acid biosynthesis genes in Mycobacterium smegmatis (76), consistent with other data implicating regulation of membrane lipid composition, stress response, PB resistance, and ExPortal integrity. In contrast to *gdpP*, previous transposon mutagenesis screens for SpeB regulators have failed to identify *clpX* (30), which is likely due to a polar effect on the gene immediately downstream, engB (L897_03625), encoding YsxC, a GTPase essential for ribosome assembly (77, 78). In addition to the regulation of toxin expression in other Gram-positive pathogens (79, 80), regulation of several stress responses has also been recognized as one of the multiple functions of ClpX, a component of the ClpXP AAA+ protease (54). Together, these data are consistent with the idea that many core PBr genes confer enhanced ExPortal integrity via activation of stress responses.

Intriguingly, like ExPortal proteins, several of the proteins encoded by core PBr genes have been found to localize to specific subcellular regions in other bacteria. These include FtsH, which localizes to the septum in *B. subtilis* (81); PtsI, which clusters at the poles in *E. coli* and *B. subtilis* (82–84); and ClpX, which clusters in cytoplasmic foci in multiple bacteria (85–88) and whose localization is crucial for its function (89). Further work will address whether there is similar compartmentalization of proteins encoded by core PBr genes, along with the contribution of fatty acid metabolism, nutrient acquisition, and stress responses in how *S. pyogenes* maintains its ability to secrete toxins in response to challenge with CAPs.

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