



GlpR Is a Direct Transcriptional Repressor of Fructose Metabolic Genes in *Haloferax volcanii*

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ABSTRACT DeoR-type helix-turn-helix (HTH) domain proteins are transcriptional regulators of sugar and nucleoside metabolism in diverse bacteria and also occur in select archaea. In the model archaeon *Haloferax volcanii*, previous work implicated GlpR, a DeoR-type transcriptional regulator, in the transcriptional repression of *glpR* and the gene encoding the fructose-specific phosphofructokinase (*pfkB*) during growth on glycerol. However, the global regulon governed by GlpR remained unclear. Here, we compared transcriptomes of wild-type and $\Delta glpR$ mutant strains grown on glycerol and glucose to detect significant transcript level differences for nearly 50 new genes regulated by GlpR. By coupling computational prediction of GlpR binding sequences with *in vivo* and *in vitro* DNA binding experiments, we determined that GlpR directly controls genes encoding enzymes involved in fructose degradation, including fructose bisphosphate aldolase, a central control point in glycolysis. GlpR also directly controls other transcription factors. In contrast, other metabolic pathways appear to be under the indirect influence of GlpR. *In vitro* experiments demonstrated that GlpR purifies to function as a tetramer that binds the effector molecule fructose-1-phosphate (F1P). These results suggest that *H. volcanii* GlpR functions as a direct negative regulator of fructose degradation during growth on carbon sources other than fructose, such as glucose and glycerol, and that GlpR bears striking functional similarity to bacterial DeoR-type regulators.

IMPORTANCE Many archaea are extremophiles, able to thrive in habitats of extreme salinity, pH and temperature. These biological properties are ideal for applications in biotechnology. However, limited knowledge of archaeal metabolism is a bottleneck that prevents the broad use of archaea as microbial factories for industrial products. Here, we characterize how sugar uptake and use are regulated in a species that lives in high salinity. We demonstrate that a key sugar regulatory protein in this archaeal species functions using molecular mechanisms conserved with distantly related bacterial species.

KEYWORDS transcription factor, glycerol, fructose metabolism, catabolite repression, glucose metabolism, archaea

Many archaea are extremophiles, able to thrive in habitats of extreme salinity, pH, and temperature. These biological properties are ideal for applications in biotechnology. However, widespread use of archaea in the bioconversion process is limited by gaps in knowledge of the metabolism and associated regulatory networks of

Received 24 April 2018 Accepted 6 June 2018

Accepted manuscript posted online 18 June 2018

Citation Martin JH, Sherwood Rawls K, Chan JC, Hwang S, Martinez-Pastor M, McMillan LJ, Prunetti L, Schmid AK, Maupin-Furlow JA. 2018. GlpR is a direct transcriptional repressor of fructose metabolic genes in *Haloferax volcanii*. *J Bacteriol* 200:e00244-18. <https://doi.org/10.1128/JB.00244-18>.

Editor William W. Metcalf, University of Illinois at Urbana Champaign

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these microorganisms. *Haloferax volcanii*, first isolated from the Dead Sea, is obligately halophilic, requiring at least 0.7 M NaCl for growth, but surviving up to 5 M (1, 2). Among the archaea, this organism is highly tractable genetically (3), making it an excellent model organism to lay the groundwork for future success in metabolic engineering of extremophiles.

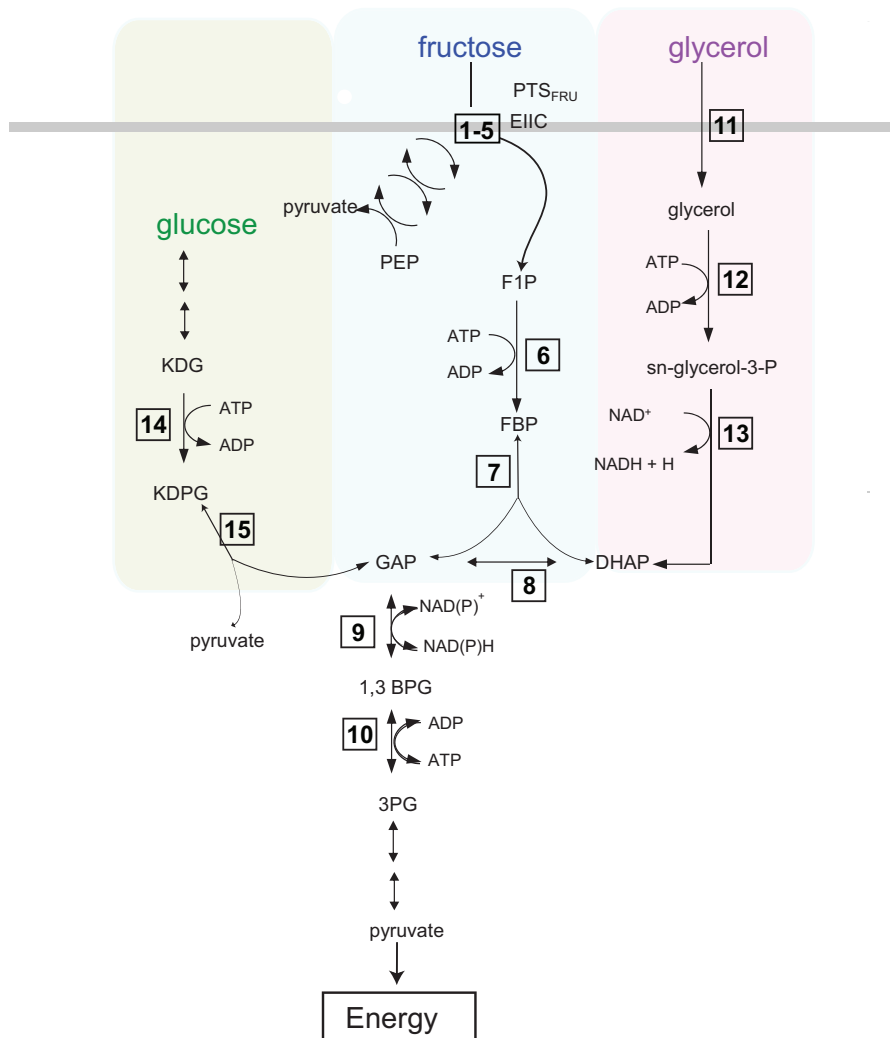
H. volcanii catabolizes a wide variety of carbon sources, including glycerol, fructose, glucose, xylitol, and chitin, among others (4). Fructose, glucose, and glycerol are taken up and degraded via three different metabolic pathways. A bacterium-like phosphoenolpyruvate (PEP)-dependent phosphotransferase system (PTS) specifically and actively transports fructose into the cell (Fig. 1) (5). Upon concomitant uptake and phosphorylation of fructose to F1P, the catabolism of fructose proceeds by a modified Embden-Meyerhof-Parnas (EMP) pathway, where F1P is converted by a series of enzymatic reactions to dihydroxyacetone phosphate (DHAP) and glyceraldehyde phosphate (GAP) (4, 5). In contrast to fructose, glucose is oxidized via an unusual archaeon-specific semiphosphorylated Entner-Doudoroff (spED) variant pathway (4, 6) (Fig. 1). Glycerol uptake by *H. volcanii* is hypothesized to occur through a putative glycerol facilitator (7). Once glycerol enters the cell, glycerol kinase (8, 9) phosphorylates glycerol to form glycerol-3-phosphate, which is transformed to GAP via the DHAP intermediate (7). All three carbon source pathways funnel GAP into the common lower shunt of the EMP pathway to form pyruvate (4).

Selective degradation of glycerol over glucose is regulated by catabolite repression in *H. volcanii* (9) and appears to be regulated in part by GlpR, a member of the DeoR family of transcription factors (10). While rare in archaea, DeoR homologs are widespread in bacteria and commonly function as specific regulators of carbon source uptake and catabolism, often playing a role in catabolite repression. Examples include catabolism of deoxyribonucleoside (DeoR), glycerol (GlpR), xylitol (XytR), and maltose (DeoT) in Gram-negative bacteria (11–15), as well as lactose (LacR), fructose (SugR), and mannitol (MtlR) in Gram-positive bacteria (16–18). Frequently, a phosphorylated catabolic intermediate relieves repression by dissociating the transcription factor from C/A-rich DNA operator binding sites (11, 12, 16, 18, 19). Although DeoR homologs typically function as tetramers (20), the pattern of cooperative binding to operators is complex and differs across organisms. The regulator can bind to multiple widely separated operators with a dyadic symmetry, while in other cases, the DeoR-type regulators bind adjacent operators with tandem symmetry (12, 13, 21, 22). For example, in *Escherichia coli*, GlpR binds up to four operators in the promoter and open reading frame of the gene encoding glycerol 3-phosphate dehydrogenase, each bound with differing affinity (22). Binding to multiple operators strongly represses transcription, facilitated by DNA looping and bending by the nucleoid-associated protein HU (22).

Our previous work demonstrated that GlpR of *H. volcanii* represses fructose and glucose catabolic enzyme-coding genes during growth on glycerol (10). The regulated enzymes include phosphofruktokinase 1 (*pfkB* product) and 2-keto-3-deoxyglucokinase (*kdgK1* product), which play key roles in fructose and glucose catabolism, respectively (5, 10) (Fig. 1). The genes of the *glpR-pfkB* operon are cotranscribed (5, 10). In contrast, studies in the closely related species *Haloferax mediterranei* have demonstrated that GlpR is an indispensable activator of the phosphoenolpyruvate-dependent phosphotransferase system (PTS) gene cluster for fructose utilization when grown on nutrient-rich medium and then supplemented with fructose (23). F1P is the hypothesized effector molecule of GlpR in both *Haloferax* species (10, 23), but this has not been demonstrated by experimental evidence. The complete regulon of GlpR and its function during growth on glucose also remain unclear. Here, we use a combination of *in vivo*, *in vitro*, and *in silico* methods to determine the effector molecule for *H. volcanii* GlpR, the global regulon of genes directly bound and regulated, and the consensus DNA binding motif.

RESULTS

Genome-wide expression analysis suggests a specific function for GlpR in the regulation of carbohydrate degradation. To determine the global GlpR regulon and



reaction	gene	enzyme	reference
1	<i>ptfC</i> , HVO_1499	Fructose-specific PTS enzyme IIC	Pickl et al., 2012
2	<i>ptfA</i> , HVO_1498	PTS enzyme IIA	Pickl et al., 2012
3	<i>ptsHI</i> , HVO_1497	PTS histidine phosphotransfer protein	Pickl et al., 2012
4	<i>ptfI</i> , HVO_1496	PTS enzyme I	Pickl et al., 2012
5	<i>ptfB</i> , HVO_1495	PTS enzyme IIB	Pickl et al., 2012
6	<i>ptfK</i> , HVO_1500	1-phosphofructokinase	Pickl et al., 2012
7	<i>tba</i> , HVO_1494	Fructose-1,6-bisphosphate aldolase	Pickl et al., 2012
8	<i>tpiA</i> , HVO_2105	Putative triose phosphate isomerase	NA
9	<i>gap</i> , HVO_0481	Glyceraldehyde phosphate dehydrogenase	Brasen et al., 2014
10	<i>pgk</i> , HVO_0480	Phosphoglycerate kinase	Brasen et al., 2014
11	<i>glpF</i> , HVO_1542	Putative glycerol facilitator	Rawls et al., 2011
12	<i>glpK</i> , HVO_1541	Glycerol kinase	Sherwood et al., 2009
13	<i>glpA1B1C1</i> , HVO_1538-1540	Glycerol-3-phosphate dehydrogenase	Rawls et al., 2011
14	<i>kdgK1/kdgK2</i> , HVO_0549/HVO_A0328	2-keto-3-deoxy-D-gluconate kinase	Rawls et al., 2010
15	<i>kdgA1</i> , HVO_0952	2-keto-3-deoxy-D-phosphogluconate aldolase	Sutter et al., 2016

FIG 1 Schematic of carbon source uptake and degradation pathways in *H. volcanii*. The semiphosphorylative Entner-Doudoroff pathway for glucose degradation is shown at the left (green box), the modified Embden-Meyerhoff pathway for fructose use is shown in the center (blue box), and glycerol degradation (purple box) is shown at the right. The thick gray line represents the membrane boundary between the outside and inside of the cell. Numbered reactions correspond to genes and enzymes with corresponding references to experimental evidence in the table at the bottom. Reactions with multiple arrows represent pathways with multiple reactions not of focus in this study. PEP, phosphoenolpyruvate; PTS_{FRU}, fructose-specific phosphotransferase transport system; F1P, fructose-1-phosphate; FBP, fructose-1,6-bisphosphate; GAP, glyceraldehyde-3-phosphate; 1,3BPG, 1,3-bisphosphoglycerate; 3PG, 3-phosphoglycerate; DHAP, dihydroxyacetone phosphate; KDG, 2-keto-3-deoxy-gluconate; EIIIC, enzyme IIC; NA, not applicable.

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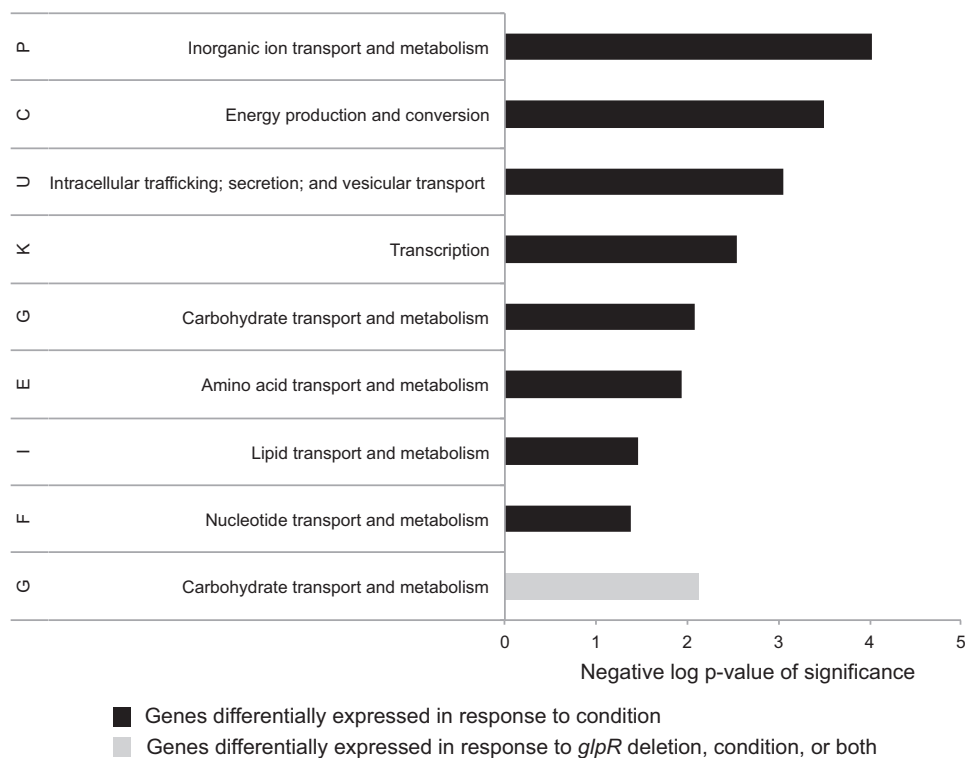


FIG 2 Statistical enrichment of differentially expressed genes in archaeal clusters of orthologous genes (arCOG) categories. Enriched arCOG categories are listed on the y axis. The x axis depicts the inverse log *P* value of significance of enrichment in each functional category according to the hypergeometric distribution. Genes differentially expressed in the parent strain in response to glucose versus glycerol are represented by the black bars; those differentially expressed in the Δ *glpR* mutant strain are in gray. Full annotations are given in Table ST2.

how it responds to various nutrients, *H. volcanii* parent and Δ *glpR* mutant strains were grown on substrates of two distinct pathways, glycerol (to analyze a C_3 substrate) and glucose (to analyze a spED substrate). Total RNA was isolated from log-phase cells, and the transcriptomes were analyzed by microarray hybridization. Differential gene expression was deemed significant by statistical analysis of the detected transcripts, as outlined in Materials and Methods.

We first determined the response of wild-type cells (parent strain H26) to a carbon/energy source. Over 600 genes were found to be differentially expressed in a comparison of the two distinct growth substrates of glucose and glycerol (see Table ST1 in the supplemental material). These 600 genes were significantly enriched in gene functions for certain archaeal clusters of orthologous gene (arCOG) categories (Wolf et al. [24]). In particular, the arCOG category of carbohydrate transport and metabolism (G) was significantly enriched, consistent with the shift in carbon/energy source (Fig. 2). Signal transduction (e.g., histidine kinases of two-component systems), noncarbohydrate transporter (e.g., cation ABC transporters), and transcription (e.g., Lrp and TATA-binding protein [TBP] families) were also found to be arCOG categories that were overrepresented among the differentially expressed genes. These results reveal the global transcript changes that occur in cells using substrates of the spED (glucose) versus glycerol metabolic pathways are primarily associated with metabolism and regulation (Table ST2 and Fig. 2).

To determine the global impact of GlpR on the cell, transcript levels in the parent and Δ *glpR* mutant strains were directly compared by statistical analysis (see Materials and Methods). Genes regulated by GlpR were identified (48 total) and found to be enriched only in carbohydrate transport and metabolism functions (arCOG category G [Fig. 2]), suggesting a specific role for GlpR in these processes. Further clustering of these 48 differentially expressed genes according to expression on the two different

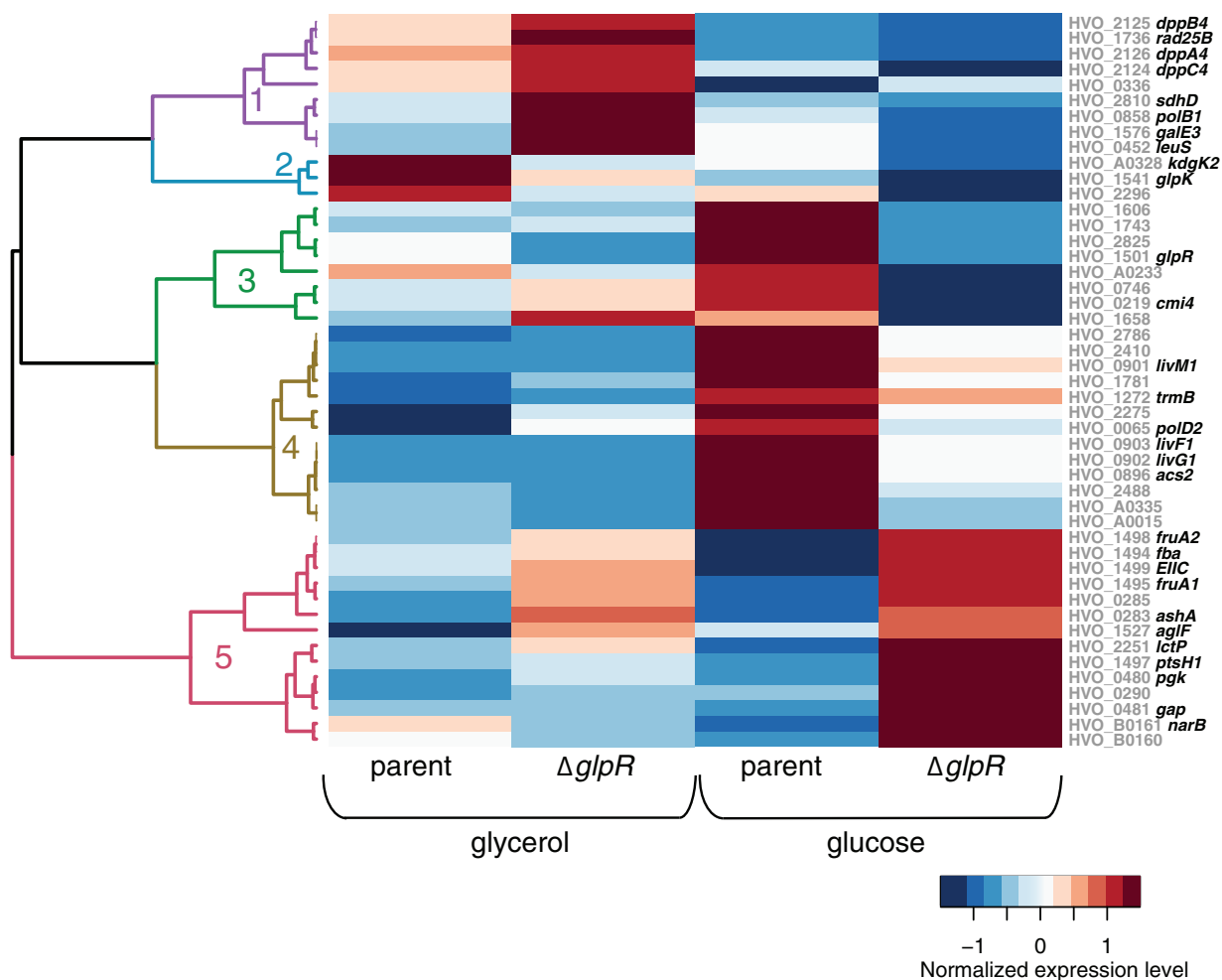


FIG 3 GlpR activates and represses genes in response to sugar. The heat map represents the results of hierarchical clustering of genes differentially expressed in the *glpR* deletion strain versus the parent strain. Each row represents mean and variance scaled expression values for each gene with significant differential expression in response to *glpR* deletion. Each gene is labeled at the right side of the heat map with its unique identifier in gray text and common name in bold black text. Each column represents the expression for that gene in each strain and under each growth condition. The dendrogram at the left is colored and numbered according to the clusters of coexpressed genes. The scale bar at the lower right indicates the colors that represent the \log_{10} gene expression levels. Clustering details and full gene annotations for each cluster are listed in Table ST3.

carbon sources revealed five patterns of GlpR-dependent gene regulation (clusters 1 to 5 [Fig. 3 and Table ST3]). Genes in cluster 1 were repressed in a GlpR-dependent manner during growth on glycerol but relatively unaffected by the deletion of *glpR* on glucose (Fig. 3, purple branches on dendrogram). Genes in cluster 1 encode putative functions in peptide transport and succinate dehydrogenase. Genes in cluster 2 were mildly activated by GlpR on glycerol and glucose (Fig. 3, blue branches). Cluster 2 included the glycerol kinase gene (*glpK*) and the spED pathway enzyme 2-keto-3-deoxygluconate kinase gene (*kdgK2*) (6). Genes in clusters 3 and 4 were activated by GlpR during growth on glucose (Fig. 3). Nearly half of the genes in cluster 3, including a putative transcription factor (HVO_0219), were also overexpressed in the $\Delta glpR$ background during growth on glycerol (Table ST3). Cluster 4 included genes encoding homologs of the TrmB transcription factor and a putative branched-chain amino acid transporter. Genes in cluster 5 were repressed by GlpR under both glycerol and glucose conditions. Genes in this cluster encoded fructose uptake and degradation functions (Fig. 3) (5). Together, these clustering results suggest bifunctional and multifactorial regulation by GlpR.

Of the 48 genes identified by microarray analysis to be differentially expressed in a GlpR-dependent manner, members of only a few transcriptional units were up- or downregulated ≥ 2 -fold due to the *glpR* deletion under either glucose or glycerol

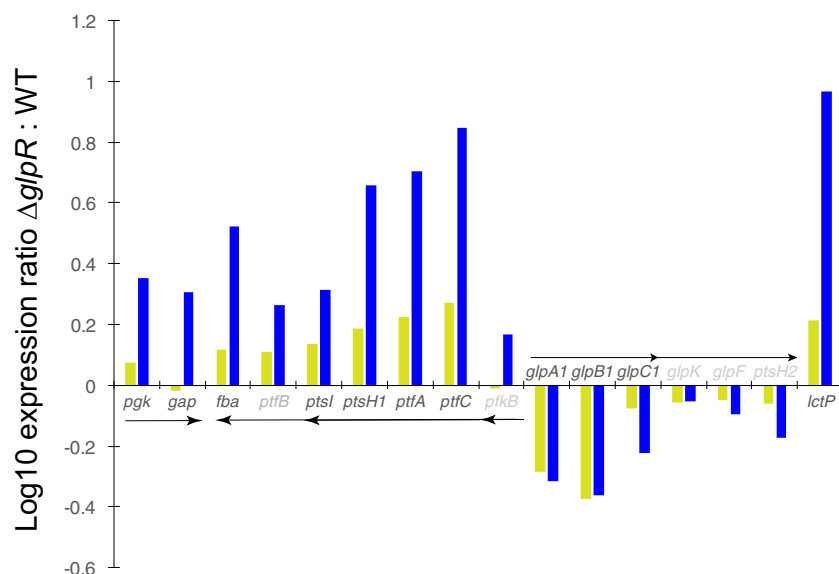


FIG 4 At least a 2-fold change in gene expression for fructose and glycerol degradation pathways is observed in $\Delta glpR$ mutant cells. Bars represent the \log_{10} fold change (mean from three biological replicates) in gene expression in $\Delta glpR$ mutant versus parent strain cells during growth on glycerol (yellow) versus glucose (blue). Arrows represent genes expressed in operons, with multiheaded arrows representing experimentally observed differential transcripts (5, 7, 10). Gene names in black text represent those at least 2-fold differentially expressed. Those in gray text did not meet the 2-fold cutoff but are expressed in operons with those expressed at least 2-fold. Full annotations and gene unique identifiers are given in Tables ST1 and ST3.

conditions (Fig. 4). These transcripts included the monocistron *lctP*, which was predicted to encode a permease for the uptake of carboxylic acids (e.g., succinate, malate, and lactate) based on homology and structural modeling (Fig. SF1). The remaining operons were associated with (i) fructose metabolism and transport (*pfkB-ptfCA-ptsH1-ptfB-fba*), (ii) the common trunk of carbohydrate metabolism (*gap-pgk*), and (iii) glycerol uptake and degradation (*glpA1B1C1-glpK-glpF-ptsH2*) (Fig. 4). Under the conditions tested, the expression of genes required for glycerol metabolism was mildly affected by the deletion of *glpR*, as the $\Delta glpR$ -based differential expression of *glpA1B1C1* (encoding glycerol dehydrogenase) barely met the 2-fold cutoff criterion, while that of *glpK* (encoding glycerol kinase) was statistically significant but did not meet the 2-fold cutoff (Fig. 4 and Table ST1). In contrast, the fructose metabolism and transport operon was strongly regulated. The members of this operon were also the only differentially expressed genes in genomic synteny with *glpR* itself. The magnitude of GlpR-mediated regulation during growth on glucose was generally greater than that for glycerol, although gene expression behavior in the $\Delta glpR$ mutant under both conditions was strongly correlated ($\hat{\rho} = 0.914$) (Fig. 4). Taken together, these data suggest that during growth on glucose and glycerol (all in the absence of fructose), (i) the primary role of GlpR is to repress genes encoding functions associated with the uptake and metabolism of fructose and (ii) GlpR plays a minor role in regulating glycerol and di-/monocarboxylic acid pathways (Fig. 4).

GlpR purifies as a tetramer stabilized by fructose-1-phosphate binding. To examine GlpR properties *in vitro*, GlpR was fused to a C-terminal His₆ tag and purified to homogeneity from *H. volcanii* by tandem Ni²⁺ affinity and gel filtration chromatography (see Materials and Methods). A high salt concentration (2 M NaCl) was used in the purification scheme to maintain halophilic protein structure. The purity of the preparation was based on total protein staining (Sypro Ruby and Coomassie blue) and anti-His tag immunoblotting analysis of GlpR separated by SDS-PAGE (Fig. 5A). GlpR was found to be associated as a homotetramer of 121 kDa, as observed by gel filtration chromatography, compared to the theoretical mass of 29 kDa for the subunit based on amino acid sequence (Fig. 5B).

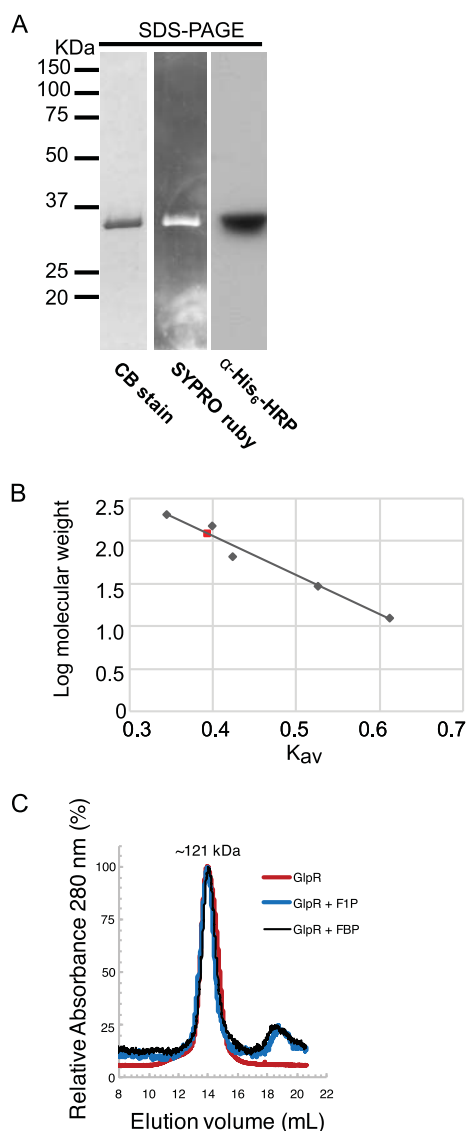


FIG 5 GlpR purifies as a tetramer. (A) Preparations of GlpR protein are pure and migrate close to the predicted monomeric molecular mass of ~29 kDa under denaturing conditions. Molecular weight ladder is shown at right. Staining methods are labeled beneath each lane (CB, Coomassie blue; α -His₆-HRP, horseradish peroxidase-conjugated His tag antibody no. HRP-66005 from Proteintech). (B) Gel filtration chromatography standard curve of GlpR under native conditions. The red square represents GlpR tetramer (~121 kDa); gray diamonds represent the standard curve. Regression line shows $R^2 = 0.96976$; $y = -4.6131x + 3.9009$. (C) Gel filtration chromatography profile of GlpR under native conditions (red line, where 100% A_{280} is 0.213) and native conditions supplemented with F1P (blue line) or FBP (black line) (where 100% A_{280} is 0.015).

To determine the effector molecule for the GlpR mechanism of action, the purified GlpR was examined for ligand interactions that promote protein stability by differential scanning fluorimetry (DSF). In the absence of a small-molecule ligand, the GlpR melting temperature (T_m) was 65°C (Table 1). In contrast, upon incubation of GlpR with fructose-1-phosphate (F1P), a metabolite in fructose degradation, the GlpR T_m increased 4°C \pm 0.1°C, suggesting that F1P significantly stabilized the tertiary structure of GlpR (Table 1). The 13 other small molecules tested displayed minimal, if any, stabilization of the tertiary structure of GlpR (Table 1). The quaternary structure of the GlpR homotetramer was relatively stable during gel filtration in the presence of F1P and fructose-1,6-bisphosphate (FBP) (Fig. 5C). Together, these results suggest that GlpR functions as a stable homotetramer, with F1P as a specific ligand. Together with the *in*

TABLE 1 DSF screen used to identify low-molecular-weight ligands that bind and stabilize purified 1 μ M GlpR-His₆^a

Small molecule (100 μ M) (abbreviation)	Δ Temp ($^{\circ}$ C) ^b
D-Fructose	-0.7 ± 0.6
Fructose-1-phosphate (F1P)	$4.0 \pm <0.1$
Fructose-1,6-bisphosphate (FBP)	$0.0 \pm <0.1$
Fructose-6-phosphate (F6P)	$-0.3 \pm <0.1$
D-Glucose	$0.0 \pm <0.1$
Gluconate	$0.0 \pm <0.1$
2-Keto-3-deoxygluconate (KDG)	-0.3 ± 0.6
2-Keto-3-deoxy-6-phosphogluconate (KDGp)	-0.3 ± 0.6
Glycerol	-0.3 ± 0.6
<i>sn</i> -Glycerol-3-phosphate	$0.0 \pm <0.1$
Dihydroxyacetone phosphate (DHAP)	-0.3 ± 0.6
Glyceraldehyde-3-phosphate (GAP)	$0.0 \pm <0.1$
Pyruvate	-0.7 ± 0.6

^aThe melting temperature of native GlpR-His₆ with no ligand was 65 $^{\circ}$ C.

^bValues represent the mean of the results from at least 3 biological replicate trials \pm standard deviation.

in vivo expression data, these *in vitro* data are consistent with the model that, upon the addition of fructose to glycerol- and glucose-grown cells, F1P levels would elevate, bind GlpR, and alter the DNA binding activity of this transcription factor. As in the case of FBA (the first metabolic step after group translocation of fructose by the PTS system), *fba* promoter (*Pfba*) transcription would be primarily stimulated by the derepression of GlpR.

GlpR binds the promoter regions of genes associated with carbon metabolic enzyme-coding genes. Genes differentially expressed in response to *glpR* deletion are presumed to include those directly bound by GlpR and those regulated indirectly. To identify direct targets, promoters of genes differentially expressed due to the deletion of *glpR* (Fig. 3 and 4) and of a gene previously shown to be GlpR regulated (*kdgK1* [10]) were selected and tested for GlpR binding assays *in vitro* and *in vivo*. Titration gel shift assays indicated that 75 nM purified GlpR was sufficient to shift DNA sequences from the *PglpR-pfkB* region, with 200 nM GlpR enabling a complete bandshift (Fig. 6A). As expected from previous evidence (7), binding was not observed for the *PkdgK2* region (Fig. 6A). However, *PkdgK1* binding was unexpectedly not observed (Fig. 7), despite previous evidence that the Δ *glpR* mutation is required for its repression (7), suggesting indirect regulation of this promoter. GlpR binding to *PglpR-pfkB* was abrogated in the presence of a 60-fold excess of unlabeled competitor DNA (Fig. 6B), which together

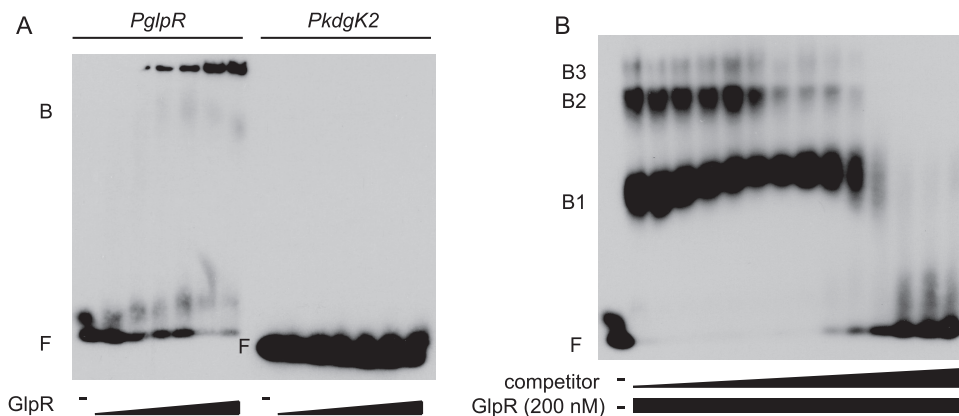


FIG 6 GlpR-promoter binding is specific. (A) Electrophoretic mobility shift assays (EMSA) with biotin-labeled *PglpR-pfkB* promoter DNA (left) or *PkdgK2* (right) incubated in the absence (–) or presence of increasing concentrations of GlpR (triangles; each lane corresponds to 25, 50, 75, 100, 150, 200 nM GlpR). F, free DNA; B, bandshift. (B) Binding of GlpR to the *PglpR-pfkB* region is specific. *PglpR-pfkB* DNA was incubated in the presence (rectangle) and absence (–) of GlpR and increasing concentrations of unlabeled competitor probe (triangle; 0 to 120 nM). F, free DNA; B1, bandshift 1; B2, bandshift 2; B3, bandshift 3.

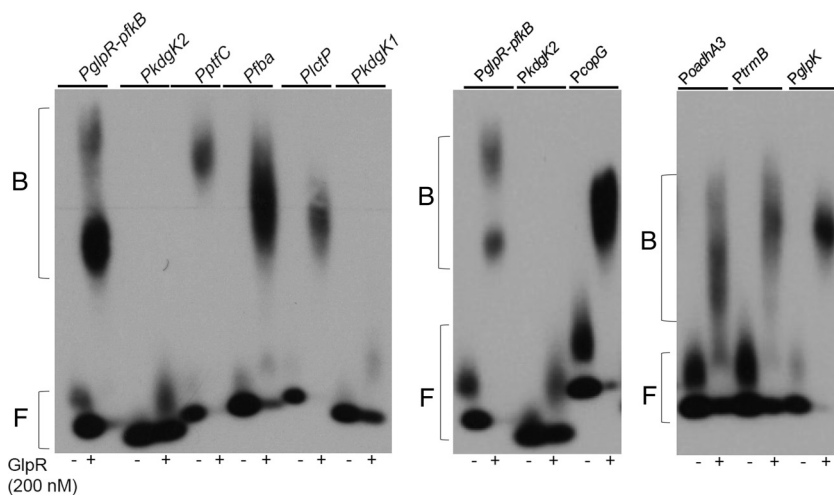


FIG 7 GlpR binds directly to enzyme-coding gene promoters. DNA from the promoter regions indicated at top were incubated with (+) or without (–) purified GlpR in EMSA reactions. These reaction mixtures were incubated in the absence of other additives. *PglpR-pfkB* and *PkdgK2* reactions were run on each gel as a positive and negative control, respectively. F, free DNA; B, bandshift.

with the absence of binding at *PkdgK1* and *PkdgK2*, suggests a specific interaction between GlpR and its own promoter. F1P was found to reduce the level of GlpR binding to the *PglpR-pfkB* promoter, supporting the model that fructose metabolism and transport genes are derepressed upon GlpR binding to this ligand (Fig. SF2).

Gel shift binding assays demonstrated that GlpR also bound to seven additional promoter regions of genes that were differentially expressed upon the deletion of *glpR*, including regions upstream of PTS system genes (*ptfC*), *fba*, *lctP*, and *glpK* and other transcription factors (Fig. 7). Binding at four of these promoters was corroborated *in vivo* during growth on various sugars by chromatin immunoprecipitation (ChIP) coupled to PCR (Fig. SF3). Control experiments and structural modeling indicated that the epitope tagging of GlpR used for these *in vivo* binding assays did not impact the activity of the protein (Fig. SF4). Together with the gene expression data, these *in vitro* and *in vivo* binding assays suggest a specific binding interaction between GlpR and target promoters, and hence direct regulation of the expression of operons involved in carbohydrate metabolism and transcriptional regulation.

Multiple C/A-rich motifs are detectable upstream of GlpR-regulated genes. In electrophoretic mobility shift assay (EMSA) experiments, we observed up to three shifted GlpR-DNA complexes, suggesting multiple GlpR binding sites within the *PglpR-pfkB* promoter region (Fig. 6B and 7). Multiple GlpR binding sites in target promoters have also been observed in other organisms, where GlpR also functions as a tetramer (18, 20). To determine a consensus binding sequence for GlpR, two strategies of *de novo* motif prediction were performed. In one search, sequences from GlpR-regulated promoters conserved across species of the *Haloferax* genus were used as input (e.g., *Pfba* and *PglpR*; see Materials and Methods [25]). In the second search, sequences upstream of genes differentially expressed in the *H. volcanii* Δ *glpR* mutant strain were used as input (Tables ST1 and ST4). In the two searches, highly similar C/A-rich motifs were identified throughout the genome, with the first search identifying a 26-bp motif that encompassed the motif identified in the second search (Fig. 8A and Table ST4). Putative binding motifs were detected upstream of eight of the genes that were differentially expressed in response to *glpR* deletion (Fig. 3, 4, and 8B). GlpR also bound to DNA regions upstream of seven of these eight genes *in vitro* and/or *in vivo* (Fig. 6, 7, 8B, and SF3). Most of these direct GlpR targets with binding motifs were located within the *glpR*-to-*fba* gene cluster (Fig. 8C). Within intergenic regions upstream of *glpR*, *ptfC*, and *fba*, multiple binding motifs flanked the putative BRE/TATA promoter sequences (Fig. 8C), corroborating the microarray evidence for GlpR repression of these genes

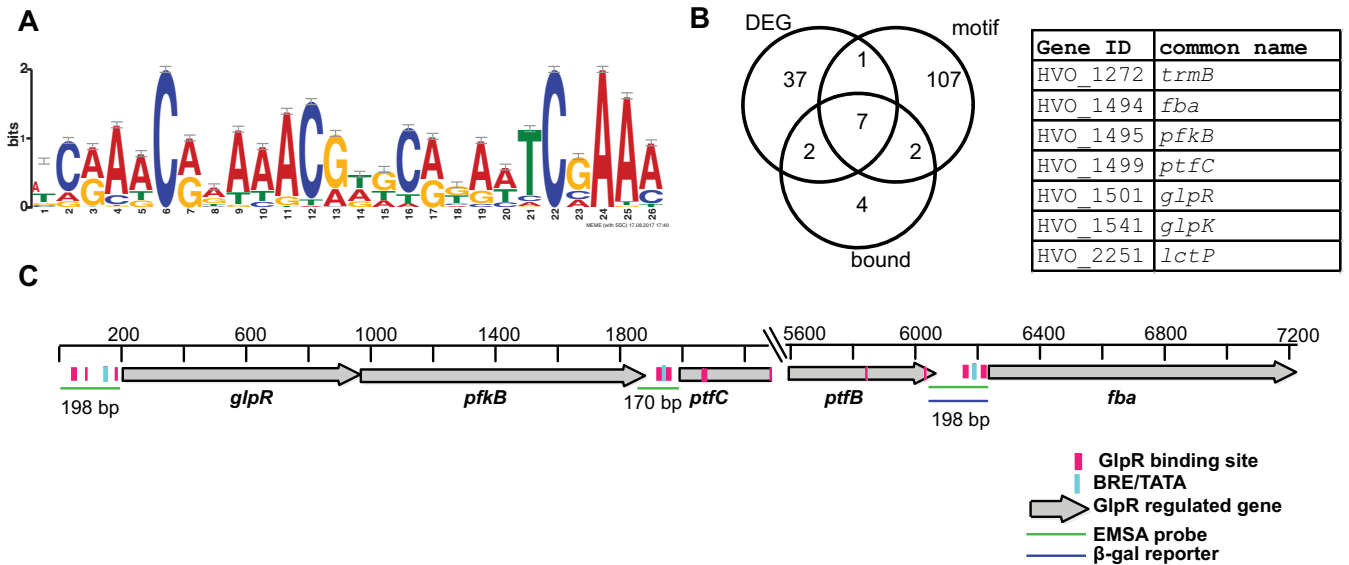


FIG 8 GlpR binds a C/A-rich motif located upstream of genes involved in fructose uptake and degradation as well as transcription. (A) Motif logo for computationally identified GlpR binding site (see Table ST4 for a full listing of motifs identified across the genome). (B) Venn diagram representing the number of genes differentially expressed (DEG) in the $\Delta glpR$ knockout mutant whose upstream regions are also directly bound by GlpR and contain the identified binding motif sequence upstream. The identities of the 7 genes in the three-way intersection of the diagram are given in the table at the right. ID, identification. (C) Locations of motifs identified in the *glpR-fba* operon. The scale bar indicates the distance from the start of the region of interest (400 bp upstream of *glpR*). See the key at the bottom right for colors and shapes. Wider pink binding-site boxes represent the full 26-bp motif, while the narrower boxes represent the shorter motif.

during glycerol and glucose growth (Fig. 3 and 4). Interestingly, sites within the *ptfC* and *ptfB* coding regions were also detected, suggestive of repression, and consistent with reports in bacteria of DNA wrapping by tetrameric GlpR at multiple binding sites within gene bodies (22). Together, these motif data suggest that GlpR directly regulates genes encoding functions in fructose degradation and uptake, perhaps by binding to multiple C/A-rich motifs for transcriptional repression.

Reporter assays verify that C/A-rich motifs are required for GlpR-mediated regulation. Guided by the microarray results and computational motif detection, we reasoned that the C/A-rich motif may serve as a GlpR *cis*-regulatory binding recognition sequence to repress target genes in the absence of fructose. To test this hypothesis, we performed reporter assays with wild-type and mutagenized *Pfba* translationally/transcriptionally fused to β -galactosidase (Fig. 9A). The expression of fusions encompassing the full *Pfba* promoter region (pJAM140) was significantly higher in the H26 parent control strain grown on fructose than that grown on glucose or glycerol (Fig. 9B; *P* values of significance listed in Table ST5). Consistent with a GlpR repression mechanism, the expression of *Pfba* from pJAM140 was significantly higher in the $\Delta glpR$ mutant than in the parent (H26) and GlpR-HA (JM2) control strains when grown on glycerol or glucose (Fig. 9B). Furthermore, *Pfba* expression from pJAM140 during growth on fructose was not significantly different between the $\Delta glpR$ mutant and control strains (Fig. 9B). Across conditions, *Pfba* activity from pJAM140 was indistinguishable in the $\Delta glpR$ mutant strain (Table ST5) and was not significantly different between the parent (H26) and control (JM2) strains (Table ST5). Together, these data suggest that GlpR is required for the repression of *Pfba* during growth on glucose and glycerol. This repression is relieved in a fructose-dependent manner.

The effect of promoter mutations on GlpR-dependent repression were compared to *Pfba* expression driven by the full promoter region (pJAM140) during growth on glycerol across the $\Delta glpR$ mutant, H26, and JM2 strains. Repression was partially relieved by the deletion of promoter distal sequences (pJAM3404) (Fig. 9A), although the expression of *Pfba* from pJAM3404 was still significantly higher in the $\Delta glpR$ mutant background than in the parent H26 strain (Fig. 9C and Table ST5). In contrast, site-

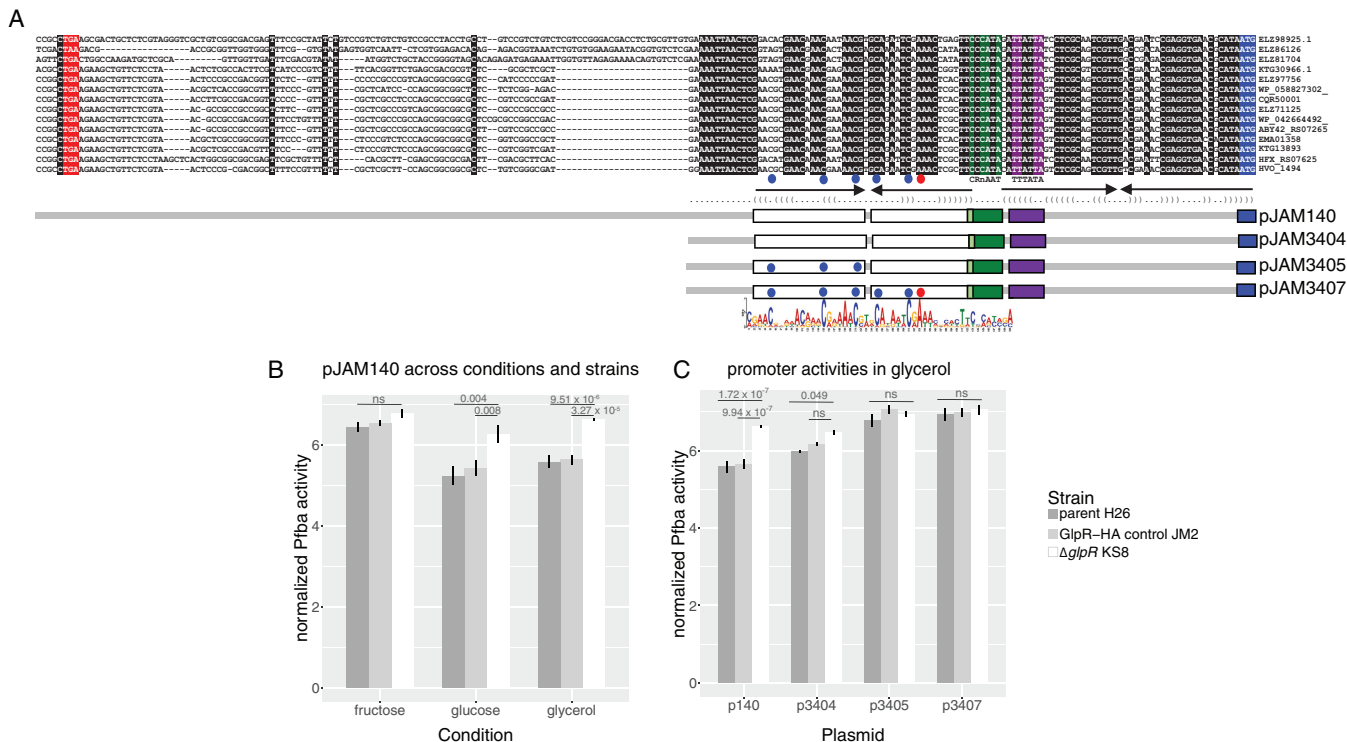


FIG 9 A conserved GlpR-dependent operator motif identified upstream of *Haloferax fba* (encoding fructose bisphosphate aldolase) is required for repression on glucose and glycerol. (A) Multiple-DNA-sequence alignment of the intergenic region 3' of *Haloferax fba* genes and the corresponding plasmids used to examine *H. volcanii fba* expression in a β -galactosidase reporter gene assay. A sequence logo representation of the consensus sequence of the GlpR-dependent operator region identified by computational analysis (see also Fig. 8) is indicated at the bottom. Database identifiers for the *Haloferax fba* genes are indicated on the right (homolog or accession number), including *H. volcanii* DS2 (HVO_1494), *H. gibbonsii* ARA6 (ABY42_RS07265), *H. mediterranei* ATCC 33500 (HFX_RS07625), *H. massiliensis* (CQR50001), *Haloferax* sp. strain SB3 (KTG13893), *H. denitrificans* ATCC 35960 (EMA01358), *H. sulfurifontis* ATCC BAA-897 (ELZ97756), *Haloferax* sp. strain ATB1 (WP_042664492), *Haloferax* sp. strain Q22 (WP_058827302), *H. prahovense* DSM 18310 (ELZ71125), and *H. elongans* ATCC BAA-1513 (ELZ86126). Conserved nucleotides are highlighted: black (general), green (TFB-responsive element BRE, CRnAAT consensus), purple (TATA box, TTTAWA consensus), blue (translation start codon), or red (stop codon of upstream gene), where W is A or T, R is A or G, and n is any nucleotide base. Inverted repeat elements predicted using the RNAfold web server (<http://rna.tbi.univie.ac.at/cgi-bin/RNAWebSuite/RNAfold.cgi>) are indicated by arrows and parentheses. Plasmids pJAM3405 and pJAM3407 have site-directed mutations, as indicated by C \rightarrow T (blue dots) or A \rightarrow T (red dots). Plasmid pJAM140 carries *Pfba* DNA corresponding to GenBank accession no. CP001956.1 (positions 1364060 to 1364252). (B) Bar plot of GlpR-dependent *Pfba* expression based on β -galactosidase reporter (*bgah*) gene assay. *H. volcanii* parent (H26, dark-gray bars), GlpR-HA control strain (JM2, light-gray bars), and KS8 (Δ *glpR* mutant, white bars) strains ectopically expressed the reporter gene from the full-length *H. volcanii Pfba* promoter region (pJAM140), as indicated in panel A. Strains were grown to log phase in glycerol, fructose, or glucose minimal medium, lysed, and analyzed for β -galactosidase activity, as indicated. Each bar reports normalized β -galactosidase activity (least-square means, log transformed). Error bars represent the standard error of the mean. Overbars report *P* values of significance from Tukey *post hoc* tests (see Materials and Methods). All pairwise significance tests were performed but not shown for simplicity. *P* values of all tests and raw data (including basal expression levels from empty vector control plasmid pJAM2715) are given in Table S75. (C) Bar plot and significance of ectopic expression of the reporter gene from various *Pfba* promoter region fragments (pJAM140, pJAM3405, and pJAM3407 [see also panel A]) across strains. Each bar reports normalized β -galactosidase activity (least-square means, log transformed). Error bars represent the standard error of the mean. Raw data, including basal expression levels from empty vector control plasmid (pJAM2715), are given in Table S75.

directed mutagenesis of specific conserved C and A residues in the promoter distal C/A motif (pJAM3405) led to significant derepression of *Pfba* in H26 and JM2 control strain backgrounds, such that expression differences between the Δ *glpR* mutant and control strains were no longer detectable (Fig. 9A and C). Site-directed mutagenesis of the downstream half-site had no additional effect on activity (expression in pJAM3405 versus pJAM3407 was not significantly different regardless of strain background; Table S75). *Pfba* activity was significantly different between pJAM3404 and pJAM3407 constructs in the Δ *glpR* mutant background but not between pJAM3405 and pJAM3407. Furthermore, activity was indistinguishable between the parent (H26) and control (JM2) strains across plasmids (Table S75), providing an additional control for the assay. Taken together, these data suggest that GlpR represses the *Pfba* promoter during growth on glycerol by binding to the C/A-rich motif. The promoter is derepressed in a GlpR- and fructose-dependent manner. These data support our model that (i) GlpR is a repressor of fructose metabolism when fructose is not present in the environment and (ii) the two C/A-rich motifs of the *fba* promoter region are required for GlpR-dependent regulation.

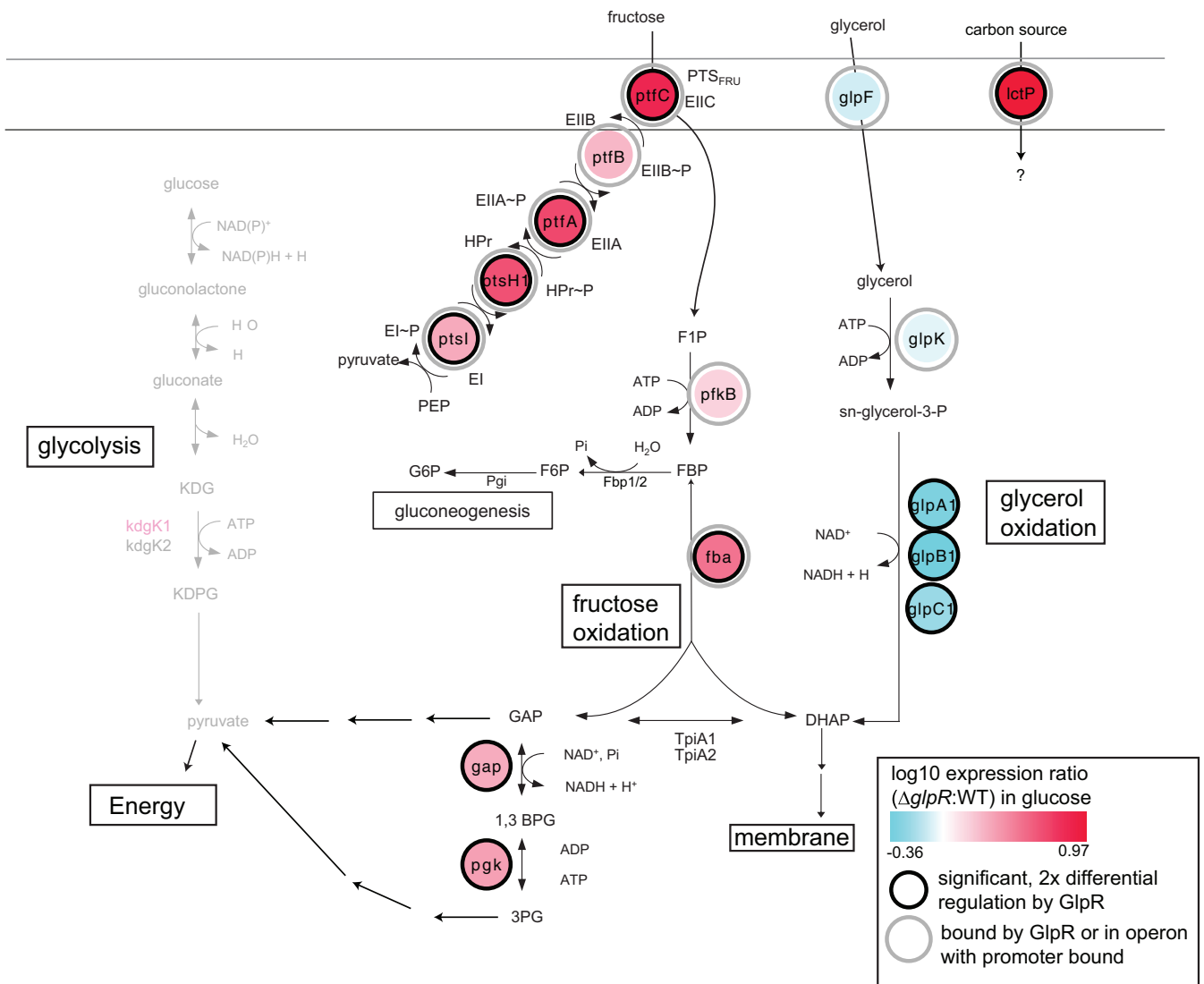


FIG 10 Metabolic reconstruction integrated with the GlpR regulatory network. Pathways for fructose, glucose, and glycerol degradation are shown overlaid with differential gene expression data in the $\Delta glpR$ mutant background and GlpR binding data (see key at bottom right). Abbreviations are as in Fig. 1. G6P, glucose-6-phosphate; Pgi, phosphoglucose isomerase; Fbp, fructose bisphosphatase.

DISCUSSION

Our previous work identified the GlpR transcription factor as a putative regulator of carbohydrate utilization pathways in the archaeal model organism *H. volcanii* (10). Here, we integrated evidence from gene expression and DNA binding experiments with computational motif detection and metabolic network analysis to demonstrate how GlpR may control these pathways (Fig. 10). Taken together, these data suggest that GlpR directly regulates genes encoding enzymes that function in fructose and glycerol uptake and degradation in response to nutrient availability (Fig. 10). In addition, GlpR may regulate other pathways through the direct regulation of other transcription factors. Specifically, our microarray and *in vitro* binding assays reveal that GlpR binds and regulates the promoter regions of genes *copG* and *trmB*, encoding putative transcription factors (Fig. 3, 4, and 7; also see Table ST1 in the supplemental material). A putative GlpR binding motif is also detected in the *trmB* promoter (Fig. 8B). The TrmB family of transcriptional regulators is conserved among archaea and has been shown to regulate genes involved in maltose, trehalose, and glucose uptake and metabolism (26–28). TrmB of *Halobacterium salinarum* regulates central metabolism and other

pathways in response to nutrient availability, particularly glucose (28–30). By homology, *trmB* (HVO_1272) of *H. volcanii* might play a role in transcriptional regulation of genes encoding metabolic enzymes or other cellular functions indirectly regulated by GlpR.

In *H. volcanii* and possibly other sugar-degrading halophiles, fructose is degraded via a modified EMP pathway, glucose is oxidized via the spED pathway, and glycerol is degraded via glycerol-3-P and DHAP (Fig. 1) (4). Fructose biphosphate aldolase (FBA) is thought to act as a central control point in the gluconeogenesis and fructose degradation pathways (4, 5), and here, we have identified a new GlpR-regulated promoter (*Pfba*) that is distinct from previously identified promoters expressing the *glpR-pfkB-ptfCA-ptsH11-ptfB* operon (Fig. 8 and 9) (5). In *H. volcanii*, GlpR regulation of carbon source degradation/uptake pathways appears to end at 3-phosphoglycerate (3PG; Fig. 10), suggesting constitutive or alternative regulation of the lower shunt of glycolysis in response to the carbon source. Thus, the data reported here are consistent with the hypothesis that GlpR regulates the expression of genes encoding enzymes in the uptake and early steps of degradation of alternative carbon sources during growth on glucose or glycerol (Fig. 10). This expands our knowledge regarding the regulation of central metabolism in archaea.

The function of GlpR in *H. volcanii* appears to differ from that of the GlpR homolog in *H. mediterranei*, a closely related species (23, 25). Cai and colleagues (23) performed a global analysis of GlpR function after growth of *H. mediterranei* on complex medium supplemented with fructose or glucose. By this approach, GlpR was found to be required to activate transcript levels of *glpR-pfkB* and *ptfCA-ptsH11-ptfB* (and not *fba*) in the presence of fructose. Among these genes, only the promoter upstream of *ptfCA-ptsH11-ptfB* (*PptfC*) was directly activated by GlpR during growth on fructose. In contrast, in *H. volcanii*, our study found that GlpR is a direct repressor of several genes encoding enzymes in the fructose uptake and degradation pathway during growth on glucose and glycerol, including *fba*.

Specifically, here, we provide *in vivo* and *in vitro* evidence that GlpR directly represses genes encoding fructose uptake and degradation (encoded by *glpR-pfkB*, *ptfC*, *fba*, [5]) and the transport of alternative carbon sources (*lctP*) but directly activates genes in glycerol metabolism (*glpK* [7]). *lctP*, which forms an apparent operon with the adenine phosphoribosyltransferase gene *hpt*, encodes a putative permease that bears structural similarity to 1-carboxylate permeases from bacteria (Fig. SF1). *LctP* is likely important for metabolism. Although GlpR regulation most strongly influences the expression levels of the fructose pathway and *lctP* genes (Fig. 3 and 4), our *in vitro* gel shift assays and motif analysis corroborate the binding of GlpR to promoter regions encoding the glycerol pathway (e.g., *glpK*) (Fig. 7, 8, and 10). We conclude that GlpR is a direct regulator of multiple carbohydrate utilization pathways.

We provide evidence for the mechanism by which GlpR accomplishes this regulation. GlpR purifies as a tetramer (Fig. 5), which may bind to C/A-rich motifs *in vivo*, repressing genes encoding fructose uptake and degradation functions (e.g., FBA) during growth on glucose and glycerol (Fig. 3, 4, 8, and 9). GlpR binds F1P *in vitro* as the specific small-molecule effector (Fig. 5 and Table 1), which likely disengages GlpR from DNA (Fig. SF2), relieving repression when F1P becomes available in the presence of fructose (Fig. 4 and 9). A homotetrameric arrangement is common for bacterial DeoR-type regulators (20, 31, 32). The F1P effector is a metabolic intermediate of fructose catabolism at an important regulatory point of the modified EMP pathway in halophilic archaea (4). *In vitro*, F1P may dissociate GlpR-DNA complexes at fructose metabolic operator regions (*glpR-pfkB*) (Fig. SF2). *In vivo*, growth on fructose derepresses *Pfba* in a GlpR- and C/A sequence-dependent manner. Together, these data are consistent with the bacterial DeoR family mechanism in which a phosphorylated sugar intermediate dissociates the transcription factor (TF) from DNA (11, 12, 16, 18, 19). For example, in the Gram-positive bacterium *Corynebacterium glutamicum*, F1P binds the DeoR-type transcription factor SugR, causing derepression of the PTS system for uptake of fructose and sucrose (18, 33). F1P is an effector that governs central metabolism and redox control in a variety of bacteria (33, 34). In summary, the results

presented here suggest that *H. volcanii* GlpR functions as a direct transcriptional regulator of carbohydrate metabolic pathways using a mechanism conserved with bacterial DeoR family transcription factors.

MATERIALS AND METHODS

Strains, media, and plasmids. The strains and plasmids used in this study are summarized in Table ST6 in the supplemental material. The oligonucleotide primers used for PCR are listed Table ST7. *Escherichia coli* TOP10 was used for routine recombinant DNA experiments. *E. coli* GM2163 was used for isolation of the plasmid DNA transformed into *H. volcanii* strains according to standard methods (35). The GlpR-hemagglutinin (HA) epitope-tagged strain (JM2) was constructed from the H26 parent strain using the *pyrE2*-based “pop-in/pop-out” method (36, 37). The HA tag was inserted in frame at the 3′ end of the *glpR* (*HVO_1501*) gene locus in the *H. volcanii* chromosome. Structural modeling, control Western blots, and promoter-reporter fusion assays were used to indicate that the HA tag did not interfere with wild-type activity of GlpR (Fig. SF4 and 9). *E. coli* strains were grown at 37°C in Luria-Bertani medium supplemented with ampicillin (100 $\mu\text{g} \cdot \text{ml}^{-1}$) and kanamycin (50 $\mu\text{g} \cdot \text{ml}^{-1}$) as needed to maintain the plasmids. *H. volcanii* strains were grown at 42°C in Casamino Acids (CA), ATCC 974 (tryptone-yeast extract) medium, and minimal medium (MM) with glycerol, glucose, or fructose, as indicated in the text and figures. Medium formulae were according to *The HaloHandbook* (35), with the following exception: glycerol, glucose, or fructose was the sole carbon source at 20 mM. *H. volcanii* media were supplemented with novobiocin (0.1 $\mu\text{g} \cdot \text{ml}^{-1}$), 5-fluoroorotic acid (5-FOA; 50 $\mu\text{g} \cdot \text{ml}^{-1}$), and uracil (10 and 50 $\mu\text{g} \cdot \text{ml}^{-1}$ for growth in the presence and absence of 5-FOA, respectively) as needed. Uracil and 5-FOA were solubilized in 100% (vol/vol) dimethyl sulfoxide (DMSO) at 50 $\text{mg} \cdot \text{ml}^{-1}$ prior to addition to the growth medium. Solid medium included agar at 15% (wt/vol). All liquid cultures were aerated with orbital shaking at 200 rpm. Cells were freshly inoculated in ice from -80°C glycerol stocks onto solid medium using a toothpick. Isolated colonies were used for inoculum into initial liquid cultures. Cells were subcultured twice to logarithmic-growth phase prior to culture for RNA extraction. Cell growth was monitored by an increase in optical density at 600 nm (OD_{600} ; where 1 OD_{600} unit equals approximately 1×10^9 CFU $\cdot \text{ml}^{-1}$ for all strains used in this study). All experiments were performed at least in triplicate.

Microarray hybridization. H26 and KS8 (*H26 ΔglpR*) were grown aerobically in glycerol MM or glucose MM in 10-ml cultures at 42°C and 200 rpm shaking to logarithmic-growth phase (OD_{600} 0.3 to 0.5). Total RNA was extracted from three biological replicate cultures, each from three pooled 10-ml technical replicate cultures, under each condition using TRI-reagent (Sigma-Aldrich), according to the manufacturer’s instructions. RNA quality was ensured using an Agilent Bioanalyzer 2100. Double-stranded cDNA libraries were created from the extracted RNA using the Superscript cDNA synthesis kit, according to the manufacturer’s instructions (Invitrogen). One microgram of cDNA from each biological replicate was labeled with Cy3 dye and hybridized to NimbleGen 12 × 135-k feature single-color custom microarray slides (Roche NimbleGen, Inc.), with each 135-k array containing 98% of the annotated genes in the *H. volcanii* genome (56). Microarray hybridization and scanning were conducted at a FSU-NimbleGen-certified facility (The Florida State University, Tallahassee, FL). For each gene, 96 replicate data points were measured (32 replicate probes per gene per array, with 3 biological replicate hybridizations per sample).

Microarray data analysis. Raw spot intensities were first normalized within arrays using RMA (38) and then normalized to quantiles of the distribution across arrays using the ANAIS online server (<http://anais.versailles.inra.fr/normalization.html>) (39). The four replicate probes corresponding to the same gene were averaged following normalization using the R limma package (40). Processed data were analyzed for significant differential expression using 2-factor analysis of variance (ANOVA) comparing expression between (i) strains (wild type [WT] versus ΔglpR mutant) and (ii) growth conditions (glycerol versus glucose). Genes with a *P* value ≤ 0.01 were considered significant in differential expression in response to the *glpR* deletion, carbon source, or both. Intensity data are reported for biological triplicate experiments, which were averaged for each gene. Log_{10} expression ΔglpR -to-WT ratios are reported in Table ST1.

Normalized intensities (mean = 0 and standard deviation = 1 for each gene across strains and conditions) are reported in Fig. 3. Differentially expressed gene lists were subjected to hierarchical clustering using Pearson complete linkage correlation and the tree cut at five branches. The clustering results are presented in the heat map in Fig. 3. These analyses were conducted in the R statistical coding environment using the packages stats (41), dendextend (42), and gplots (43). Gene annotations (24, 44, 56) for members of each cluster are presented in Table ST3. Significant enrichment in gene functional categories according to the archaeal clusters of orthologous genes (arCOG) ontology (24) was determined using the hypergeometric test in the R statistical analysis environment, as previously described (45, 46), with the code freely available at https://github.com/amyschmid/histone_arCOG.

Computational prediction of GlpR binding sequence. GlpR-regulated genes were tested for a binding motif in two orthogonal searches. In the first search, protein homologs of *H. volcanii* FBA (*HVO_1494*), GlpR (*HVO_1501*), and PtfC (*HVO_1499*) were retrieved for members of the *Haloflex* genus by the Basic Local Alignment Search Tool using BLASTP (protein-protein BLAST) (47). The DNA sequences in the 5′ direction of the genes encoding these homologs were retrieved using the graphics tool within the NCBI nucleotide portal (<https://www.ncbi.nlm.nih.gov/nucleotide/>). In the second search, 400-bp sequences upstream of *H. volcanii* genes differentially expressed in response to *glpR* deletion (Table ST1) and bound directly by GlpR in binding assays (Fig. 6, 7, SF2, and SF3) were used as input. In both searches, *de novo* motif detection was performed using the MEME Suite version 4.12.0 (48), with the

following parameters: any number of repeats, max width of 20 to 26 bp, and 3 output motifs. The strongest resultant motif (the motif from the second search is encompassed by the first) is reported in Fig. 8A. Compared to shuffled sequences, this motif was highly significant (Wilcoxon signed-rank test, $P < 5.291 \times 10^{-14}$). MAST and FIMO algorithms from the MEME Suite were used to scan the *H. volcanii* genome (uid12524 version 210 within the GenBank Bacteria Genomes and Proteins database) for additional putative targets of GlpR binding. The results of genome scanning are given in Table ST4.

Transcriptional reporter construction, assay, and statistical analysis. A plasmid-based reporter system was used to analyze transcription from promoter regions of *Pfba* and control promoters by fusion to the *Haloferax alicantei*-derived *bgaH* encoding β -galactosidase (10). All plasmids used in this study are listed in Table ST6, and all primers are listed in Table ST7. For the construction of pJAM140, the region of interest was amplified from *H. volcanii* DS70 genomic DNA by PCR. PCR products were fused to *bgaH* using XbaI and NdeI sites of plasmid pJAM2678. Plasmid pJAM3404 was generated by PCR using pJAM140 as the template. PCR based site-directed mutagenesis was applied to generate the plasmids pJAM3405 and pJAM3407 containing the variants of *Pfba*. In brief, pJAM3404 was used as the template to construct the plasmids by reverse PCR. The PCR amplicon was treated with DpnI restriction enzyme (NEB) followed by T4 polynucleotide kinase, as recommended by the supplier (NEB). The product was circularized by T4 ligase (NEB) and transformed into *E. coli* TOP10 competent cells. The constructed plasmids were confirmed by PCR method and sequencing (Eton Bioscience, San Diego, CA).

The β -galactosidase activity of each construct was assayed quantitatively in logarithmic-growth phase, as described in reference 49. The resultant activity values were normalized to protein concentration as estimated by the Bradford assay (50) and background values, in which no substrate (*o*-nitrophenyl- β -D-galactopyranoside [ONPG]) was added to the reaction mixture. For each strain across plasmids and conditions, 7 to 19 biological replicate trials were performed.

The resultant data were subjected to log transformation, then analyzed for statistical significance in two separate two-factor ANOVAs, each with an underlying linear regression model. *Post hoc* analysis of residuals indicated that log transformation was necessary to uphold normality and homoscedasticity assumptions of ANOVA. In the first test, β -galactosidase activity (dependent variable) from the pJAM140 plasmid was compared across strains (JM2, KS8, and H26; first independent variable) and across conditions (glucose, glycerol, fructose; second independent variable). In the second test, β -galactosidase activity was compared across plasmids (pJAM140, pJAM3405, and pJAM3407) and strains under glycerol conditions. *P* values from pairwise Tukey *post hoc* tests of significant differences between means of groups determined by ANOVA to have main effects on the dependent variable are reported in Fig. 9 and Table ST5. Given the unbalanced nature of the underlying data (i.e., different numbers of replicates across groups), least-square means of log-transformed data are reported in Fig. 9. Analyses were conducted in the R coding environment using the packages *car* (51), *Rmisc*, and *lsmeans* (52). Bar graphs were generated using the package *ggplot2* (53).

Protein purification. A His₆ tag fusion strategy was used that incorporated tandem Ni²⁺-affinity and size-exclusion chromatography. High salt concentration (2 M NaCl) was maintained throughout the purification to maintain halophilic protein structure. The purity of the preparation was based on total protein staining (Sypro Ruby and Coomassie blue) and anti-His tag immunoblotting of the GlpR protein separated by reducing SDS-PAGE. *H. volcanii* KS8-pJAM124 strain (Δ *glpR* mutant carrying *glpR*-His₆ gene in *trans*) was grown to stationary phase (4×1 -liter cultures in 2.8-liter Fernbach flasks). Cells were harvested by centrifugation ($6,700 \times g$ for 10 min, 4 °C). Cell pellets were resuspended in 60 ml Tris-salt-low-imidazole buffer (2 M NaCl, 50 mM Tris-HCl [pH 8.0], 40 mM imidazole). The cells were lysed by passage through a French press ($4 \times$, 2,000 lb/in²). Cell lysate was clarified by centrifugation ($20,000 \times g$ for 30 min, 4 °C) and sequential filtration through 0.8- μ m and 0.45- μ m cellulose acetate filters (Fisher Scientific). Sample was applied at a flow rate of $5 \text{ ml} \cdot \text{min}^{-1}$ to a HisTrap high-performance (HP) column (5 ml; GE Healthcare) equilibrated with Tris-salt-low-imidazole buffer and washed with ~ 5 column volumes of the same buffer. GlpR-His₆ was eluted from the column with high-imidazole buffer (2 M NaCl, 50 mM Tris-HCl [pH 8.0], 500 mM imidazole) at a flow rate of $1 \text{ ml} \cdot \text{min}^{-1}$, and 1-ml eluate fractions were collected. Fractions (9 to 13) containing GlpR-His₆ were combined and concentrated with Amicon Ultra 15-ml centrifugal filters (10-kDa molecular weight cutoff [MWCO]; Merck Millipore) by centrifugation at $3,900 \times g$ in a swinging bucket rotor (75 min total, 4 °C). The concentrated protein sample was filtered (0.45 μ m), and sample (500 μ l) was injected onto a gel filtration column (Superdex 200 HR 10/30; GE Healthcare) equilibrated with Tris-salt buffer (2 M NaCl, 50 mM Tris-HCl [pH 8.0]). Eluate was collected in 0.5-ml fractions at a flow rate of $0.3 \text{ ml} \cdot \text{min}^{-1}$. Fractions (29 to 34) containing GlpR-His₆ were pooled, concentrated, filtered, and reinjected onto the gel filtration column, as mentioned above. Protein quantification was performed using a bicinchoninic acid (BCA) assay, according to the supplier (Pierce, Rockford, IL). GlpR-His₆ purity was assessed by SDS-PAGE under reducing conditions and subsequent Coomassie blue staining.

Differential scanning fluorimetry. DSF was adapted from Niesen et al. (54), with the following modifications. Purified GlpR-His₆ (1 μ M) was mixed with 0.1 mM small molecule (Table 1) in Tris-salt buffer (2 M NaCl, 50 mM Tris-HCl [pH 8.0]) with $1 \times$ Sypro Orange (Invitrogen). The mixtures with Sypro Orange were maintained in the dark. Aliquots of the mixtures (40 μ l) were transferred to a 96-well PCR microplate and incubated for 10 min at room temperature, followed by a temperature gradient of 22 to 95 °C at $1^\circ \text{C} \cdot \text{min}^{-1}$ using a C1000 thermal cycler (Bio-Rad CX96 real-time system). Fluorescence was scanned for 5 s at temperature increments of 0.2 °C. Protein melting temperatures were calculated by melting curve fitting using CFX Manager 2.1 (Bio-Rad). The organic compounds examined for influence on GlpR by DSF included D-fructose 1-phosphate barium salt trihydrate (F1P; catalog no. S408697;

Aldrich), D-(−)-fructose (catalog no. CAS 57-48-7; Acros Organics), D-fructose 1,6-bisphosphate tetra(cyclohexylammonium) salt (FBP; catalog no. F0752; Sigma), D-glucose 6-phosphate dipotassium salt hydrate (F6P; catalog no. F1502; Sigma), D-(+)-glucose (catalog no. CAS 50-99-7; Acros Organics), D-gluconic acid sodium salt (gluconate; catalog no. G9005; Sigma), 2-keto-D-gluconic acid hemicalcium salt hydrate (KDG; catalog no. K6250; Sigma), 3-deoxy-2-keto-6-phosphogluconic acid lithium salt (KDGP; catalog no. 79156; Sigma), glycerol (catalog no. G5516; Sigma), sn-glycerol 3-phosphate bis(cyclohexylammonium) salt (G3P; catalog no. G7886; Sigma), dihydroxyacetone phosphate dilithium salt (DHAP; catalog no. D7137; Sigma), D-glyceraldehyde 3-phosphate solution (GAP; catalog no. 39705; Sigma), and sodium pyruvate (pyruvate, catalog no. P2256; Sigma).

Electrophoretic mobility shift DNA binding assays. DNA probes for EMSAs were generated by PCR using oligonucleotides, as indicated in Table S7, where one of the primers was 5′ end biotinylated (Integrated DNA Technologies). Prior to use, the 5′-end-labeled probes were purified by DNA agarose gel electrophoresis using a QIAquick gel extraction kit. GlpR-His₆ at the indicated concentrations (Fig. 6 and 7; Fig. SF2) was incubated with 1 nM 5′-biotinylated probe in binding buffer (2 M NaCl, 20 mM Tris-acetate [pH 6.0], 2 mM EDTA, 2 mM dithiothreitol [DTT], 15 mM MgCl₂, 100 μg · ml⁻¹ bovine serum albumin [BSA], 57 mM 2-mercaptoethanol, 10% [wt/vol] sorbitol) for 30 min at 42°C. After incubation, the reactions were run on a nondenaturing 0.8% (wt/vol) agarose gel in high-salt TBE buffer (pH 6.0; 220 mM Tris base, 945 mM boric acid, 5 mM EDTA) for 34 min at 140 V. After cooling and 30-min equilibration in two changes of ice-cold 0.5× TBE buffer (pH 8.3), DNA and nucleoprotein complexes were transferred to a nylon membrane (BrightStar Plus; Ambion) using the Trans Blot system (Bio-Rad) at 150 V for 6 h. DNA and nucleoprotein complexes were cross-linked to the membrane via UV radiation (UV Stratallinker 2400; Stratagene) and visualized using PhosphorImager (New England Biolabs). The sources of the metabolites used in EMSAs are detailed in “Differential scanning fluorimetry,” above.

Accession number(s). The microarray platform used in this study was deposited in the NCBI Gene Expression Omnibus (GEO) (55) under accession number [GPL21414](https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GPL21414). Raw and normalized microarray data generated in this study are freely available through NCBI GEO at accession number [GSE77589](https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE77589).

SUPPLEMENTAL MATERIAL

Supplemental material for this article may be found at <https://doi.org/10.1128/JB.00244-18>.

SUPPLEMENTAL FILE 1, XLSX file, 1.0 MB.

SUPPLEMENTAL FILE 2, XLSX file, 0.1 MB.

SUPPLEMENTAL FILE 3, XLSX file, 0.1 MB.

SUPPLEMENTAL FILE 4, XLSX file, 0.1 MB.

SUPPLEMENTAL FILE 5, XLSX file, 0.1 MB.

SUPPLEMENTAL FILE 6, XLSX file, 0.1 MB.

SUPPLEMENTAL FILE 7, XLSX file, 0.1 MB.

SUPPLEMENTAL FILE 8, PDF file, 5.3 MB.

ACKNOWLEDGMENTS

We acknowledge funding awarded through the U.S. Department of Energy, Office of Basic Energy Sciences, Division of Chemical Sciences, Geosciences and Biosciences, Physical Biosciences Program (grant DE-FG02-05ER15650) to J.A.M.-F.; the National Institutes of Health (grant R01 GM57498) to J.A.M.-F.; and the National Science Foundation (grants MCB-1615685, MCB-1417750, and MCB-1651117) to A.K.S.

We thank S. Shanker at the UC ICBR Genomics Core for Sanger DNA sequencing, Steven Miller at the USF microarray facility for technical assistance with transcriptomics data, and Peter Tonner for statistical advice.

We declare no conflicts of interest.

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