Expression profiling of hypothetical genes in *Desulfovibrio vulgaris* leads to improved functional annotation

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ABSTRACT

Hypothetical (HyP) and conserved HyP genes account for >30% of sequenced bacterial genomes. For the sulfate-reducing bacterium *Desulfovibrio vulgaris* Hildenborough, 347 of the 3634 genes were annotated as conserved HyP (9.5%) along with 887 HyP genes (24.4%). Given the large fraction of the genome, it is plausible that some of these genes serve critical cellular roles. The study goals were to determine which genes were expressed and provide a more functionally based annotation. To accomplish this, expression profiles of 1234 HyP and conserved genes were used from transcriptomic datasets of 11 environmental stresses, complemented with shotgun LC–MS/MS and AMT tag proteomic data. Genes were divided into putatively polycistronic operons and those predicted to be monocistronic, then classified by basal expression levels and grouped according to changes in expression for one or multiple stresses. One thousand two hundred and twelve of these genes were transcribed with 786 producing detectable proteins. There was no evidence for expression of 17 predicted genes. Except for the latter, monocistronic gene annotation was expanded using the above criteria along with matching Clusters of Orthologous Groups. Polycistronic genes were annotated in the same manner with inferences from their proximity to more confidently annotated genes. Two targeted deletion mutants were used as test cases to determine the relevance of the inferred functional annotations.

INTRODUCTION

The application of genome sequencing and sequence annotation to numerous bacterial species has yielded a ‘road map’ for several avenues of research. These include the incorporation of gene expression changes at both the mRNA and protein levels (1–5) with physiological and metabolic studies to deduce the behavior of the microbe as a whole, a field now called Systems Microbiology. Other approaches to discern function include genetic manipulations such as gene/protein tagging for the identification and visualization of protein complexes (6–8) and deletion mutagenesis (9–12) for confirming the function(s) of a given gene or protein. One aspect that has come to light through the sequencing of more than 780 completed bacterial and archeal genomes (http://www.ncbi.nlm.nih.gov/genomes/lproks.cgi) with an additional 2179 ongoing (http://www.genomesonline.org/gold.cgi), is that approximately one-third of all of the genes within a given genome are typically predicted to encode hypothetical (HyP) and conserved HyP

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The authors wish it to be known that, in their opinion, the second and third authors should be regarded as joint Second Authors.

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genes (13). HyP proteins are defined as those with no significant sequence similarity (i.e. homology) to any characterized or uncharacterized predicted proteins, while conserved hypothetical (CHyP) proteins are those that have significant similarity to a predicted protein in another species or strain without direct evidence of the expression of the gene as defined by TIGR (http://cmr.jcvi.org/tigr-scripts/CMR/CmrHomePage.cgi).

This lack of similarity to well-characterized genes and proteins increases the interest in these groups of sequences, as they may well be important to specialized cellular physiology and metabolism, may possess unique functions, or complete tentatively assigned pathways. In the case of HyP, their potential existence is only supported by the output of gene prediction programs such as GLIMMER (14,15). More recently however, computational tools for determining plausible functions of gene products have been developed. These include methods such as genomic context analysis (16–19) and phylogenomic profiling (20,21). Such tools are still of limited application for HyP or CHyP due to the lack of sequence homology for the gene or protein of interest within a well-characterized, sequenced genome. Hence, a working knowledge of the function(s) of the gene products encoded is expected to lead to a greater mechanistic understanding of the metabolic capabilities of a target microorganism and possibly to a better physiological knowledge of other organisms through gene sequence and neighbourhood association.

As a first step to the identification of gene function, it is important to determine if the genes are actually expressed, if expression results in production of a functional protein, and under which conditions the genes and proteins may be differentially regulated or influenced. Experiments to obtain expression information have applied a combination of transcriptomic and proteomic methods. In Haemophilus influenzae, 54 HyP genes of a total 1744 genomic loci were found to be expressed with both methods of analysis and used to assign specific functions for 16 genes and general roles for another 27 genes (13).

More recently, the metal-reducing bacterium Shewanella oneidensis MR-1 was the focus of three investigations (22–24). In this organism, 40% of the sequenced genome consists of genes predicted to encode HyP and CHyP. The first study of S. oneidensis MR-1 by Kolker and co-workers (24) showed the actual transcription of >500 HyP and CHyP genes with at least a general functional assignment for 240 of them. In a related study by Elias et al. (22) of only HyP, comprehensive MS-based proteomics data were queried across many culture conditions, and the results confirmed the existence of 262 predicted proteins. Additionally, inferences were made for the subcellular localization and function from differential expression in these discreet culturing conditions (22). A study of the CHyP in this same organism by Elias et al. (23) confirmed protein production from 758 such genes with improved functional assignments that were also inferred in part from the culturing conditions of expression (23).

The sulfate-reducing bacterium (SRB) Desulfovibrio vulgaris Hildenborough has a sequenced genome (25), and several physiological and metabolic studies have taken advantage of this information (3,26–29). This bacterium is well known as a model SRB (30) and is known to reduce and immobilize metals and radionuclides (31–33) making it of interest for bioremediation efforts. Additionally, D. vulgaris Hildenborough was recently the focus of an effort to assign putative functions to predicted HyP from proteomics data obtained through LC-MS/MS analysis of cultures grown with different electron donors (34). These assignments were based not only on proteomics data but also relied on a number of computational methods and resulted in the re-annotation of 20 CHyP genes and the confirmation of gene expression for 129 HyP genes.

In the current study, we have incorporated the previously published results and expanded the analyses. Transcriptomics data derived from 11 different stresses as well as corresponding shotgun LC-MS/MS proteomics data from selected stress conditions were used to query the expression of each HyP or CHyP gene. Some of these data have already been published as part of a genome-wide transcriptional response to a particular stress condition. These published data include responses to heat shock (26), high salt (28), nitrate or nitrite exposure (35,36), stationary versus exponential growth (2), high pH (37), effect of deleting the Fur gene (10) and O₂ exposure (38). We have incorporated the HyP and CHyP microarray and proteomics data from these studies as well as currently unpublished results. The latter conditions include peroxide stress in the wild type and a mutant lacking the perK gene; a strain lacking pDV1, the 202 kb native plasmid found in D. vulgaris Hildenborough, versus the wild-type D. vulgaris that has pDV1; a co-culture with a methanogen versus D. vulgaris alone, acidic conditions, cold stress, chromium exposure, NaCl adaptation and Fe(II)-limitation. The microarray datasets for all HyP and CHyP genes in published studies have been made publicly available (http://genomics.lbl.gov/supplemental/DvH-hypotheticals-090223/).

These data were compiled and the genes categorized by basal expression rates by the presence versus the absence of differential gene expression in response to a particular stress, by whether the response was specific to a single stress or seen in multiple stress conditions and by the operon status of the genes, monocistronic or polycistronic. For the latter, differential expression in a given stress that coordinated with the rest of the operon was also considered. Additionally, bioinformatic information such as Clusters of Orthologous Groups (COG) and subcellular location were used for functional annotation. This may be the first such comprehensive investigation utilizing mRNA microarrays and proteomics to infer a more robust functional annotation of HyP and CHyP genes from such a large number of stress conditions.

MATERIALS AND METHODS

Biomass production

Desulfovibrio vulgaris Hildenborough (ATCC 29579) was grown in a defined lactate (60 mM)/sulfate (50 mM) medium, LS4D (28), under a variety of different stress conditions as have been reported (28,35,36). The chilled
samples were harvested via centrifugation, flash frozen in liquid nitrogen and stored at −80°C until analysis.

**Culture maintenance**

*D. vulgaris* cultures from the American Type Culture Collection (ATCC) were grown to mid log phase in 1 liter of LS4D, checked for purity by the appearance of anaerobic SRB colonies on LS4D plates as well as the absence of colonies on aerobic glucose plates, dispensed into 2 ml cryogenic vials (Nalgene) with 0.5 ml 30% (vol/vol) glycerol and frozen at −80°C until used as previously described (28). To minimize phenotypic drift from repetitive culturing, all experiments used cells that were started from frozen stocks and were fewer than three subcultures from the original ATCC culture. All experiments, inoculations and transfers were performed in an anaerobic glove chamber (Coy Laboratory Products Inc., Grass Lake, MI) with an atmosphere of ~5% CO₂, 5% H₂ and 90% N₂.

**Microarray transcriptomic and data analysis**

DNA microarrays using 70-mer oligonucleotide probes covering 3482 of the 3534 annotated protein–coding sequences of the *D. vulgaris* genome that were constructed as previously described (39). Of the 52 genes not found on the microarrays, 14 were either HyP and CHyP (under Expression Category; Supplementary Tables 5 and 6). Total RNA extraction, purification and labelling with the fluorophore Cy5-dUTP (Amersham Biosciences, Piscataway, NJ) were performed independently on each cell sample using previously described protocols (38). Genomic DNA was extracted from *D. vulgaris* cultures and hybridized as previously described (36). Microarray data analyses were performed using gene models from NCBI. All mRNA changes were assessed with total genomic DNA (gDNA) as a control for each of the experimental and control hybridizations. Log₂ ratios of mRNA to gDNA hybridized to gene oligonucleotides and z-scores were computed as previously described (9). A mean log₂ ratio cut off across time points of ≥1.2 and an accompanying z-score ≥2, respectively, were used to identify the genes whose expression changed significantly.

\[
Z = \frac{\log_2 (\text{Treatment} / \text{Control})}{\sqrt{0.25 + \sum \text{variance}}}
\]

**Proteomics and proteomic data analyses**

*Shotgun LC-MS/MS analysis.* Sample preparation, chromatography and mass spectrometry for shotgun LC–MS/MS proteomics were performed as described previously (35). Briefly, frozen cell pellets from triplicate 50 ml cultures were thawed and pooled prior to cell lysis. Cells were lysed via sonication in lysis buffer, composed of 4 M urea with 500 mM triethylammonium bicarbonate, pH 8.5 (Sigma-Aldrich, St Louis, MO), and the clarified lysate was used as total cellular protein. Sample denaturation, reduction, blocking, digestion and labelling with isobaric reagents were performed according to the manufacturer’s directions (Applied Biosystems, Framingham, MA). Strong cation exchange chromatography was used to separate the iTRAQ-labelled samples into 21–23 salt fractions. Fractions were desalted using C₁₈ MacroSpin Columns (Nest Group, Southborough, MA), dried and separated on a PepMap100 C₁₈ reverse phase nano-LC-MS column (Dionex-LC Packings, Sunnyvale, CA) using an Ultimate HPLC with Famous Autosampler and Switchos Micro Column Switching Module coupled with an ESI-QTOF mass analyser (QSTAR® Hybrid Quadrupole TOF, Applied Biosystems) as previously described (26). A 2 h gradient from 0 to 25% acetonitrile was used. Two product ion scans were collected from each cycle with a 1 s accumulation time. A threshold of 50 counts was required for ions to be selected for fragmentation. Parent ions and their isotopes were excluded from further selection for 1 min, with a mass tolerance of 100 ppm.

Collected mass spectra were analysed using Analyst 1.1 with ProQuant 1.1, ProGroup 1.0.6 (Applied Biosystems) and MASCOT version 2.1 (Matrix Science, Inc, Boston, USA). A FASTA file containing all the putative open reading frame (ORF) sequences of *D. vulgaris*, obtained from Microbes Online (http://www.microbesonline.org) (40), was used to form the theoretical search database along with the common impurities trypsin, keratin, cytochrome c and bovine serum albumin. The same search parameters were used in both programs; namely, trypsin was designated as the cleavage enzyme, a maximum of one missed cleavage was allowed, mass tolerances of 0.1 for mass spectrometry and 0.15 for tandem mass spectrometry were allowed, andcharge states from +2 to +4 were searched. Only proteins identified by at least two unique peptides in at least one of the data sets at greater than 95% confidence by both ProQuant and MASCOT were considered.

**AMT tag analysis.** Whole cells lysis via bead beating and whole cell lysate tryptic digestion were performed as described previously (22,23). Separation of insoluble (i.e. membrane bound/associated) from soluble proteins in whole cell lysates was achieved with ultracentrifugation (356 000 × g, 10 min, 4°C) as described elsewhere (22,23). The capillary LC system and controller, in-house manufactured mixer, capillary column selector and sample loop for manual injections as well as separations are also as previously described (22,23).

All samples were analysed as previously described (22,23,41). The collision-induced dissociation (CID) tandem mass spectra from the LC-ion trap mass spectrometer measurements were analysed with SEQUEST (42) using the protein sequences deduced from the *D. vulgaris* Hildenborough genome sequence (25). All samples were analysed by a 9.4-T FTICR-MS (Bruker Daltonics) as described previously (43). Mass spectra were acquired with ~10⁵ resolution.

High-stringency constraints were used in the filtering of the data to maximize peptide identification confidence as described previously (22,23). All peptides were required to be fully or partially tryptic. In order to
gauge the confidence of MS peak matching from the FTICR data to the SEQUESTTM result, an algorithm to determine the quality of the match score, termed the ‘discriminant score’, was used (23). This scoring system computed a measure of confidence for each observation of each peptide via an extension of the approach described by Aebersold and co-workers (44). This incorporated the predicted central normalized elution time (NET) values instead of filtering out low-confidence peptides solely using observed and predicted NET values. It also utilized several SEQUESTTM scoring parameters (peptide cleavage state, difference in observed and computed mass, difference in observed and predicted NET and other indicators) to compute a confidence score for each peptide identification. This eliminated a fixed score threshold, e.g., SEQUESTTM Xcorr value of two, to filter peptides for inclusion in a database. The advantage was that a discriminant-based score is less likely to discard peptide identifications than a score based upon threshold criteria. Incorporation of NET data improved peptide identification confidence by ~10% compared to not using elution information (45). At least one high-confidence ‘unique’ peptide (i.e., mapping to only one possible parent protein) and a total of two peptides were required for protein identification in each database.

The FTICR data were processed using the PRISM data analysis system as described previously (22). Since the separation systems for both FTICR and LCQ analyses were identical, peptide confirmation was based on both the calculated (from the mass tag database) and measured mass (from the FTICR analysis) of the peptide matching to within 6 ppm and the elution times matching to within 5%.

Expression categorization of HyP and CHyP proteins

Each HyP or CHyP gene was identified and sorted as monocistronic or part of a polycistronic operon. This distinction allowed for inferences in the latter as to functional annotations by basing the expression responses to stresses and association with characterized genes in the same operon.

Each of these two groups of genes was then categorized solely on the basis of the microarray expression profiles. The first category was divided into those genes that exhibited ‘high expression’, where the basal expression levels were within the top 1/8th (12.5%) of all gene transcript levels, and those with ‘normal expression’, where basal expression was below the 12.5%. The basal gene expression level was determined by calculating the mean Log2 ratio of mRNA to gDNA hybridization intensities normalized as described earlier for all microarray experiments. With this method, the more negative Log2 value (e.g. −14.9) indicated a smaller degree of absolute expression while a less negative number (e.g. −10.5) indicated a more highly expressed gene (9). Genes were categorized as ‘not expressed’ if their second highest observed mean Log2 ratio of mRNA to gDNA hybridization intensity on any individual array was < −14.0, an arbitrary cutoff determined by visual inspection of probability density distributions for HyP + CHyP genes compared to annotated genes. Each of the basal expression groups was then further subdivided into the following differential gene expression categories: (1) expressed genes that lacked differential expression in response to any of the stress conditions, (2) those that showed differential expression to only one stress, (3) those that showed differential expression to multiple stresses ‘multiple stress response proteins’ and (4) the category ‘not expressed’ included those genes that showed no expression under any of the tested conditions.

Differential gene expression in response to either single or multiple stresses was determined by the observation of a minimum |log2 R| value ≥ 1.2 and a corresponding |z-score| ≥ 2 as compared to the control condition. Because samples were analysed at several time points after the induction of the stress condition, we established that this differential level of expression had to be observed in at least 20% of the time points to be further considered. If these parameters were met for only one of the stress conditions, then the gene was placed into the ‘single stress response’ category. If the gene met these criteria in two or more of the 11 stress conditions, then they were placed in the ‘multiple stress response’ category. In either case, the current annotation was retained for genes not meeting this criterion since every conceivable growth condition was not tested, making it premature to classify these predicted genes as ‘non-coding gene region’.

Deletion mutagenesis

Specific HyP or CHyP genes were selected for targeted deletion based upon the microarray datasets. These genes included the monocistronic HyP gene DVUA0095 that is on the native plasmid of the organism and responded only to chromate stress. The second was a pair of genes (DVU0303 and DVU0304) currently annotated as an operon on the main chromosome. The expression of these HP genes was predicted and found to be directly or indirectly influenced by the Fur regulon (10) and exhibited differential expression in virtually all stress conditions tested.

Deletion cassette construction. Deletion cassettes were constructed by a method similar to the molecular bar-coding methods described for Saccharomyces cerevisiae (46, 47). PCR primer sets were designed to amplify ∼800 bp up- and downstream of the selected ORF with unique barcode sequences between the common sequences and Km’ sequences (Supplementary Table 1). The PCR mixtures, marker exchange procedures, transformation and mutant selection procedures including Southern-blot analyses were performed as previously described (10). The three segments; up- and down-stream of the gene of interest as well as the Kanamycin cassette were individually amplified by PCR and then ligated by a fourth SOEing PCR. This mutagenic cassette was then introduced into D. vulgaris via electroporation, where a double recombination event replaced the target gene with the drug-resistance gene.
Physiological assessment of mutants
The deletion mutants were tested for growth compared to wild-type D. vulgaris under the same conditions used in the original stress experiments from which the microarray and proteomic datasets were generated, along with growth in LS4D medium at pH 7.2 (28) as a control. Amendments and modifications for the stress conditions included the addition of 250 mM NaCl (salt stress), lowering the pH to 5.5 (acid stress), addition of 100 mM or 150 mM sodium nitrate (nitrate stress), 1 mM or 2 mM sodium nitrite (nitrite stress) and 0.2 mM, 0.4 mM or 0.45 mM potassium chromate (chromate stress). Optical density (A600) measurements were taken periodically up to 400 h in duplicate cultures.

Sequence analysis
Protein sequence similarity was determined using FastBLAST (48) with an e-value threshold of 0.01 and an effective database size equal to 2.23 x 10^7. D. vulgaris Hildenborough protein sequences obtained from RefSeq (release 28 March 2008) (49) were searched against the non-redundant protein database from NCBI (as of May 15, 2008) (ftp://ftp.ncbi.nih.gov/blast/). Operon predictions (50) and COG (51) assignments were based on MicrobesOnline (40) data from the 7 April 2008 release [including the November 2007 release of the NCBI Conserved Domain Database (52)].

Homology searches and putative functional assignments
Several publicly available in silico tools were utilized in an attempt to assign a more detailed putative function to each of the HyP or CyHP genes that were expressed according to the microarray experiments. The first tool used was PSORTb version 2.0.4 (http://www.psort.org/psortb/) that was set for Gram negative organisms (53,54). This tool predicts the subcellular location of a given protein by estimating the presence and number of trans-membrane helices, the presence of signal pathway motifs as well as other parameters. These tools were used along with the final localization estimate in conjunction with the microarray and proteomic data to give the most accurate functional annotation possible. The second method was TMHMM (55) (http://www.cbs.dtu.dk/services/TMHMM-2.0/) that predicts the number of trans-membrane helices and determines if the protein is inside or outside the cytoplasm of the cell and was used to corroborate the findings with PSORTb. Other bioinformatics tools such as those used previously (34) were attempted for several HyP and CHyP genes other than those already reported (34), but the results were ambiguous and so they were not pursued for this work.

Statistical comparisons of basal expression distributions
Probability density plots were created in the statistical software environment R (http://www.r-project.org/) with probability densities estimated by smoothing with a Gaussian kernel. Statistical tests for differences in expression level distributions were computed in R using the two-sided Mann–Whitney test with a continuity correction in the normal approximation for the P-value.

RT-PCR HyP and CHyP basal expression
Eight genes were selected for reverse transcription PCR (RT-PCR) in order to provide a biological verification of the microarray results. These genes were selected across the range of the average of the basal expression range with emphasis on the lower end so that if all these genes were expressed according to RT-PCR, then the assumption could be made that most if not all of the other genes were expressed as well. The positive control was the constitutively expressed dsrC gene (DVU2776) with an average basal expression rate of −9.7, which would place it above the top 1/8th percentile cutoff of −11.8 so as to be placed in the ‘highly expressed category’.

RESULTS AND DISCUSSION
Global detection of HyP and CHyP gene-expression products
The sequenced genome of D. vulgaris shows an expected 887 HyP and 347 CHyP genes for a total of 1234 possible gene products. In general, mRNA was confidently detected for 1212 of these genes using microarrays, thus indicating actual expression of the gene (Table 1). Additionally, shotgun LC–MS/MS and AMT tag proteomic analysis further confirmed the expression of 786 proteins from HyP or CHyP genes. This represents the detection of gene expression for over 99% of the annotated HyP and over 95% of the CHyP genes with a complementary 471 (53%) of the HyP and 306 (88%) of the CHyP genes detected at the protein level (Table 1; Supplementary Table 3). In comparison, a recent report detailed the expression of 129 HyP and CHyP D. vulgaris genes via proteome analysis, with possible functional reassignments using in silico approaches (34).
Reverse transcription PCR was conducted in order to corroborate the findings of the microarray and proteomic analysis. Eight HyP and CHyP genes were selected with five being at the lower end of the basal expression range. Two genes (DVU1526 and DVUA0044) were included as negative test cases since all data to date suggests that DVU1526 is not expressed, while DVUA0044 was not on the microarrays and was not detected by either proteomic method. The well-annotated DVU2776 (dsrC) served as the positive control. The RT–PCR revealed that all eight of the test genes yielded bands at the predicted sizes along with the positive control, while the two negative test genes yielded no bands (Figure 1). These data were consistent with the microarray and proteomic data. Furthermore, because five of the eight test cases were among the lowest recorded basal expression rates, the results give an increased confidence that at least the large majority of genes deemed to be expressed according to the microarray and proteomics data are correct.

Neither proteomic approach gave evidence for protein synthesis from 448 transcribed genes. We sought to determine whether any bias existed that might explain this omission such as, was there a general difference in the expression of well-annotated genes versus the HyP and CHyP? Basal expression statistical profile comparisons were performed to answer this question. Overall, HyP and CHyP displayed lower gene-expression levels than well characterized proteins ($P = 1.5 \times 10^{-12}$, two-sided Mann–Whitney test; Figure 2A). This was not surprising since the core metabolic genes required for survival are unlikely to be among the HyP and CHyP and so might be expected to be expressed at or above the average levels. However, on an individual basis, the HyP and CHyP genes have appeared amongst the most highly differentially expressed genes under particular stress conditions (2,10,26,28,36,38).

With respect to confident identification of HyP and CHyP by proteomics, those that were identified showed a higher gene-expression level than those not identified ($P = 2.9 \times 10^{-5}$, two-sided Mann–Whitney test; Figure 2B). Again, this is as expected, since the most abundant proteins should be preferentially identified. Additionally, a determination of any bias in the proteomic data revealed that in a comparison of HyP vs CHyP, monocistronic vs polycistronic, proteins above or below 100 amino acids in length and highly expressed vs all others, both methods underrepresented proteins <100 amino acids in length (Figure 3A and B). Given this information, one possibility was that there was a lack of tryptic cut sites yielding no peptides for mass spectrometric detection. However, a query for the presence of either a lysine (K) or arginine (R) revealed that this was the case in only 3 of the 448 genes that did not have a protein confidently identified by either method (Supplementary Table 4). Curiously, 12 HyP and CHyP genes were confidently identified at the protein level but not with mRNA. Further investigation found that only five of these genes were on the microarrays (DVU0522, DVU1148, DVU1748, DVU2022, DVUA0050) while the remaining seven genes (DVU0509, DVU0797, DVU0833, DVU1852, DVUA0052, DVUA0088, DVUA0145) were not (Supplementary Tables 5 and 6). This suggests that there may have been an issue with the microarray hybridization for these five genes or that transcription was below detection.

Given this information and that one of the goals of this study was to assign a functional annotation to as many of the HyP and CHyP genes as possible, a separation of the polycistronic from monocistronic genes was performed. Such classifications allow the evaluation of the hypothesis that more of the polycistronic HyP and CHyP genes would be expressed compared to the monocistronic ones. We assumed a greater likelihood for a more accurate functional annotation if the gene were in a predicted operon and displayed similar stress response patterns to more confidently annotated genes in that same operon. Hence, monocistronic and polycistronic genes were treated separately for the remainder of the study.

### Table 1. HP and CHP genes with evidence of expressions

<table>
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<tr>
<th>Current annotation</th>
<th>Number of possible genes&lt;sup&gt;a&lt;/sup&gt;</th>
<th>Transcript identified&lt;sup&gt;b&lt;/sup&gt;</th>
<th>Protein identified&lt;sup&gt;c&lt;/sup&gt;</th>
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</table>

<sup>a</sup>Computational identification of putative open reading frames were as previously described (25,63).
<sup>b</sup>Transcript evidence obtained from microarray experiments reported by VIMSS/ESP efforts (2,10,26,28,36,38).
<sup>c</sup>Proteins identified by shotgun LC–MS/MS and/or AMT tag proteomics as previously described (35,64,65).
Figure 2. Comparison of the average basal gene expression levels as quantified by microarray analysis. (A) A comparison of the 1212 expressed HyP and CHyP genes (solid line) versus the 2278 better annotated genes (broken line) showed that the former displayed significantly lower expression levels overall; $P = 1.5 \times 10^{-12}$, two-sided Mann–Whitney test. (B) The 774 HyP and CHyP gene-expression products detected at the protein level (broken line) were significantly more abundant than the 438 proteins not confidently identified by either proteomics method (solid line); $P = 3.0 \times 10^{-5}$, two-sided Mann–Whitney test.

Figure 3. Comparison of HyP and CHyP detection at the protein level using the (A) AMT tag and (B) shotgun LC–MS/MS approaches. The bars are additive between the number of proteins not detected (black bar) and those detected (grey bar). A comparison of each of the subgroups with both methods showed a bias against the detection of proteins under 100 amino acids in length.
Categorization of differential expression patterns under stress conditions

The first step in characterization of the 882 HyP and 330 CHyP genes that were transcribed was to categorize them according to observed expression patterns under one or more of the cultivation/stress conditions tested. Collectively, 45% of all of the expressed HyP and CHyP genes were found to be polycistronic with the remaining 56% being monocistronic (Table 1). However, it is interesting that despite a greater number of monocistronic genes (687) compared to polycistronic genes (548), the distribution amongst the expression categories was quite similar (Figure 4). In each case, only 0.3% of the monocistronic and 0.2% of the polycistronic genes were highly expressed with no observed differential transcription. Highly expressed genes, differentially expressed or not, comprised 7.3% and 6.2% of these groups, respectively. Similarly, 10.9% of the monocistronic and 11.7% of the polycistronic genes that were not highly expressed responded to a single stress, while a considerably higher percentage (73.9% and 71.5%, respectively) responded to multiple stress conditions. This similarity in the categorizational proportions between the two groups of genes continued in the ‘expressed’ (6.4% and 8.4%, respectively) and not expressed (1.5% and 2.2%, respectively) categories. For each of the polycistronic genes not observed to be expressed (12 cases), the other genes in the operon were expressed under some condition. Therefore, these were not cases of an operon not being expressed, but rather particular genes not showing expression. Reasons for these patterns are not clear. In fact, a preliminary assumption was that more of the monocistronic genes would not be expressed than genes within operons, since the latter would be more likely to be co-expressed with the rest of the operon. However, this was not the case since the percentage of genes in each category was similar as detailed above.

Expression profiling and putative functional assignment to monocistronic genes

Monocistronic genes are arguably more difficult to functionally annotate, because the reference point of neighbouring genes is absent. However, the 81 monocistronic genes that responded to a single stress did give clues as to function, 65 encoded a HyP with the remaining 16 encoding a CHyP (Supplementary Table 5). In order to demonstrate the observed expression profiles, the stress responses for a randomly selected set of genes responding to a single stress condition are shown (Figure 5). It is interesting that even among these essentially unknown genes, several are found to be responsive to the same stress. Several exhibited differential expression when transcripts in stationary phase were compared to those in exponential cells or when acid treated culture transcripts were compared to base treated. In contrast, there were cases where multiple HyP genes responded specifically to one stress, as in the case of chromate exposure. Both DVUA0095 (Figure 5B) and DVU1338 were upregulated by chromate exposure while DVU2436 was downregulated (Supplementary Table 5). No polycistronic genes were solely influenced by chromate. Based on these findings and the observation that D. vulgaris lacking pDV1 is less tolerant of chromate exposure (M. Fields, personal communication), DVUA0095, located on pDV1, was targeted for deletion to ascertain the cellular response to chromate in this mutant.

For the purposes of functional annotation, each of the HyP genes has been renamed to reflect the stress response influencing its expression along with any other in silico features as determined by the use of COGs, TMHMM and PSORTb (Supplementary Tables 5 and 6). For example, DVUA0095 was found to be up-regulated upon exposure to chromate. Results from in silico analyses predicted that it possesses three transmembrane helices but no signal peptide motifs, with a final score of 9.46.
that it is associated with the cytoplasmic membrane. Given the lack of a signal peptide or assigned COG, we infer that the protein resides in the cytoplasmic membrane with the bulk of the protein facing the cytoplasm. Hence, this protein has been re-annotated to be a 'chromate-induced, cytoplasmic membrane protein'. For others where no such structural features were predicted, the genes were simply renamed e.g. 'acid-induced protein' or 'heat-repressed protein'. The remaining monocistronic genes that did not display a differential response to any of the stresses have simply been renamed as e.g. 'expressed protein in D. vulgaris' or 'expressed cytoplasmic membrane protein'.

Expression profiling and putative functional assignment of polycistronic genes
The polycistronic HyP and CHyP genes often have a reference point for their plausible biochemical function based on their location within operons that include ORFs with more characterized, orthologous genes in other bacteria. One of the main criteria used to assign function to the HyP or CHyP genes was the similarity of the differential expression pattern of the gene to that of other genes within the operon. An additional criterion was the degree of nucleotide overlap with other genes within the operon that could suggest transcriptional coupling. Good examples of such scenarios were found in areas of the genome apparently containing temperate bacteriophages or their remnants, such as DVU2488–DVU2729 containing several predicted operons. Some temperate phages may be induced by catastrophic stress conditions or as cells enter the stationary phase of growth. In fact, stresses from high heat and the stationary phase of growth resulted in the differential expression of the greatest
number of polycistronic genes that are likely to encode phage functions. One example is the apparent increased expression of a seven-gene operon likely to be involved in temperate phage activity during stationary phase versus exponential growth (Figure 6A). It is prudent to note that other changes in culture conditions usually coincide with these events, such as sulfide and acetate accumulation with a concomitant change in pH and a decrease in the specific growth rate.

Another example of HyP and CHyP genes within temperate phage operons was the four-gene operon of DVU0192 (adenine-specific DNA methyltransferase), DVU0194 (terminase) along with the HyP genes DVU0193 and DVU0195. In this case, all four genes responded coordinately to the onset of stationary phase compared to exponential growth (Figure 6B), suggesting that the two HyP genes could be involved in phage DNA metabolism.

Other cases were not as straightforward, such as with the predicted operon of DVU1639–DVU1642 (Supplementary Table 6) that contains two HyP and two CHyP genes. DVU1639 showed increased expression only to stationary phase whereas DVU1640 was not expressed. DVU1641 and DVU1642 were both up-regulated in multiple stresses including showing Fur-regulation. These results question the validity of the operon assignment. The three transcribed genes were renamed based upon the microarray stress response data. Hence, DVU1639 was reannotated as a ‘stationary phase induced protein’, whereas DVU1641 and DVU1642 were reannotated as a ‘Fur influenced, multiple stress induced protein’ and a ‘multiple stress induced outer membrane protein’, respectively. The original annotation for DVU1640 remained.

**Fur-influenced regulation of HP and CHP genes**

*D. vulgaris* possesses three Fur regulator paralogs DVU0942 (Fur), DVU3095 (PerR) and DVU1340 (Zur) (56). While little information is available on the latter in *D. vulgaris*, the global regulation roles of Fur and PerR have been explored in more depth. Gene deletions within *D. vulgaris* are available for the putative global regulators Fur (10) and PerR (unpublished results), and transcriptional analyses in a few stress conditions have been performed. In a number of bacteria, the Fur system regulates the uptake of ferric iron (57,58). Although Fur has also been shown to control the synthesis of specialized Fe(III) chelators known as siderophores (57,59,60), there is no evidence for siderophore production in *D. vulgaris*. In contrast, Fur is suggested to play a regulatory role in oxidative stress, motility, virulence and acid tolerance (57,58). PerR, the second of the potential global regulators, has been best studied in Bacillus subtilis (61,62). Experimental evidence supported a role for the PerR system in the cellular response of *D. vulgaris* to oxidative stresses such as peroxide or metal ion limitation through increased expression of ruberythrin and rubredoxin genes (10). Bender and co-workers (10) reported that the HyP gene DVU2681 was reported to have the greatest increase in transcription of all genes in the Fur deletion mutant, while 6 of the 21 genes showing strong increases in transcription in the absence of Fur were HyP or CHyP genes.

In the current work, a number of HyP and CHyP genes appeared to be either differentially transcribed in Fur or PerR deletion mutants compared to the wild-type strain, or displayed stronger transcriptional responses in the deletion mutant. Expression changes occurring in the deletion mutants were inferred to result from the altered genetic background, and, in particular, when the PerR deletion mutant was exposed to a 0.1% O2 stress or the Fur deletion mutant was exposed to high salt or high nitrate. A small proportion of genes apparently were influenced by both Fur and Per (Table 2). Hence, in these cases, the prefix of ‘Fur-influenced’, ‘Per-influenced’ or ‘Fur- and Per-influenced’ was added to the annotation where appropriate (Supplementary Tables 5 and 6). While such numbers of proteins are probably not directly linked to either of these global regulatory systems, it is conceivable that indirect, cascading regulators could affect this number of genes.
Table 2. Apparent HyP and CHyP influence by the global regulators Fur and Per

<table>
<thead>
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<th>Fur</th>
<th>Per</th>
<th>Fur and Per</th>
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<td>Highly expressed, multiple-stress response</td>
<td>3</td>
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Validation of putative assignments with targeted deletion mutagenesis

In order to test the relevance of the stress responses recorded via microarrays and proteomics, as well as the inferred functional annotations applied to the HyP and CHyP genes in D. vulgaris during this study, two targeted deletions were constructed. A two-gene operon, DVU0303-DVU0304, predicted to be part of the Fur regulon (56) showed altered transcription rates in all conditions tested (Supplementary Table 6). The second targeted deletion was the monocistronic gene DVUA0095 that increased in expression only upon exposure to chromate (Figure 5B; Supplementary Table 5). Both deletion mutants and the wild type grew similarly in the control unamended medium containing 30 mM sodium lactate and 60 mM sodium sulfate at pH 7.2, pH 5.5 and when amended with 250 mM NaCl (Figure 7A–C). These results were as predicted for ΔDVUA0095 since it responded only to chromium exposure. However, the growth results of Δ(DVU0303-DVU0304) was unexpected since both genes originally increased in transcription at pH 5.5 and in 250 mM NaCl. However, such a lack of correlation between gene expression and cellular fitness in an imposed treatment is not uncommon (46).

Other stress conditions included amendments of 100 mM NaNO₃, or NaNO₂ at 1 mM or 2 mM. Expression of DVU0303 in wild-type cultures increased with NaNO₃ or NaNO₂ (35,36). A slightly increased sensitivity of Δ(DVU0303-DVU0304) with these stressors (Figure 7D–F) was observed. The interpretation of changes in mutant growth rate and extent may suggest a functional role in tolerance to the treatment or a general metabolic perturbation that impedes the production of cell material. Surprisingly, the ΔDVUA0095 mutant also grew more poorly relative to the wild type when exposed to the various nitrogen species (Figure 7E and F). Reasons for this phenotype are not clear, but could include oxidative stress and subsequent protein or DNA damage due to the unstable nature of nitrite.

The final stress conditions used to test the deletion mutants were three concentrations of K₂CrO₄ at 0.2 mM, 0.4 mM and 0.45 mM. When wild-type cultures were challenged with 0.45 mM K₂CrO₄, the transcriptional analysis showed large increases in expression of DVUA0095 as well as for DVU0303-DVU0304. At the lower concentration of 0.2 mM K₂CrO₄, there were no apparent effects on the growth of any of the strains (Figure 7G). At 0.4 mM K₂CrO₄, ΔDVUA0095 exhibited a lag phase of 75 h, compared to 25–30 h in ΔDVU0303–DVU0304) and the wild type (Figure 7H), suggesting that the removal of this gene interfered with the response to this level of K₂CrO₄. At 0.45 mM K₂CrO₄, both ΔDVUA0095 and ΔDVU0303-DVU0304 showed extended lag phases of 190 and 175 h, respectively, approximately twice that of the wild type (Figure 7I).

CONCLUSIONS

The main purpose of this study was to validate the expression of D. vulgaris genes annotated as HyP and CHyP and to infer additional functions when possible. Overall, 98% of the HyP and CHyP genes were found to be transcribed via microarrays with 63% also being translated. Among these, many displayed specific transcriptional responses to single stresses whereas others showed responses to multiple treatments. Some of these genes were also shown to be influenced by or in regulons of, the global regulatory systems of Fur and/or PerR. The fact that these genes actually produce proteins increases the possibility that they may play some role in the responses to environmental perturbations.

Assessment of the ΔDVUA0095 and Δ(DVU0303–DVU0304) mutants confirmed a likely role for DVUA0095 in cellular responses to chromate as inferred from the microarray results. However, further complexity exists as revealed by some surprising phenotypes of the mutants as described earlier. Ongoing and future work is anticipated to include a detailed analysis of many HyP and CHyP genes, particularly those that displayed responses to only a single stress. This will include gene tagging with eventual protein complex elucidation, assessment of gene deletions and additional stress treatments. Through these efforts, a better understanding of the physiological and metabolic aspects of bacteria with sequenced genomes such as D. vulgaris will be achieved. This type of methodical analysis of unknown genes may well be useful as a means to derive more meaningful functional annotations in other organisms where a few or many RNA microarray and/or proteomic datasets exist. Reanalysis of these data can range from simply comparing HyP and CHyP transcript or proteome abundances in different culturing conditions to the more discreet stress conditions such as were used to generate the data for the present work. In addition, decreasing the lists of HyP and CHyP genes will improve annotations through inter-organismal comparisons.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.
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Conflict of interest statement. None declared.

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