Predicted Highly Expressed Genes of Diverse Prokaryotic Genomes

SAMUEL KARLIN* AND JAN MRÁZEK

Department of Mathematics, Stanford University, Stanford, California 94305-2125

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Our approach in predicting gene expression levels relates to codon usage differences among gene classes. In prokaryotic genomes, genes that deviate strongly in codon usage from the average gene but are sufficiently similar in codon usage to ribosomal protein genes, to translation and transcription processing factors, and to chaperone-degradation proteins are predicted highly expressed (PHX). By these criteria, PHX genes in most prokaryotic genomes include those encoding ribosomal proteins, translation and transcription processing factors, and chaperone proteins and genes of principal energy metabolism. In particular, for the fast-growing species Escherichia coli, Vibrio cholerae, Bacillus subtilis, and Haemophilus influenzae, major glycolysis and tricarboxylic acid cycle genes are PHX. In Synechocystis, prime genes of photosynthesis are PHX, and in methanogens, PHX genes include those essential for methanogenesis. Overall, the three protein familiesribosomal proteins, protein synthesis factors, and chaperone complexes—are needed at many stages of the life cycle, and apparently bacteria have evolved codon usage to maintain appropriate growth, stability, and plasticity. New interpretations of the capacity of Deinococcus radiodurans for resistance to high doses of ionizing radiation is based on an excess of PHX chaperone-degradation genes and detoxification genes. Expression levels of selected classes of genes, including those for flagella, electron transport, detoxification, histidine kinases, and others, are analyzed. Flagellar PHX genes are conspicuous among spirochete genomes. PHX genes are positively correlated with strong Shine-Dalgarno signal sequences. Specific regulatory proteins, e.g., two-component sensor proteins, are rarely PHX. Genes involved in pathways for the synthesis of vitamins record low predicted expression levels. Several distinctive PHX genes of the available complete prokaryotic genomes are highlighted. Relationships of PHX genes with stoichiometry, multifunctionality, and operon structures are discussed. Our methodology may be used complementary to experimental expression analysis.

Gene expression and protein abundances in prokaryotes are regulated at several levels: (i) initiation of transcription, promoter strength, promoter configuration, and transcription factors; (ii) transcription termination, mRNA stability, and turnover rates; (iii) codon usage; (iv) translation initiation and elongation; and (v) protein folding, degradation, and cellular localization. An accounting of high gene expression in prokaryotic genomes generally focuses on at least one of three criteria: (i) The gene possesses a potent promoter sequence sometimes associated with bent DNA and/or specific binding factors. However, the characterization of regulatory cis elements underlying gene transcription is largely an unresolved problem. (ii) The gene possesses a strong Shine-Dalgarno (SD) ribosome binding sequence, but recognition of SD sequences is not discriminating (10, 14, 34, 51, 53) (see also below). (c) The gene exhibits favorable codon usage; in rapidly dividing bacteria, this largely corresponds to the prevalent codon usage frequencies of ribosomal protein (RP) genes (20, 44, 54). Our approach to ascertaining gene expression levels relates to codon usage differences among gene classes. We show data suggesting that codon usage contributes importantly to setting the level of expression of the gene. Our data support the proposition that each genome has evolved a codon usage pattern accommodating "optimal" gene expression levels for most situations of its habitat, energy sources, and life style.

Gene codon preferences vary considerably within and between organisms (for reviews and perspectives, see references 25, 29, and 55). Variations in tRNA availabilities are interpreted by several authors as an important factor in generating codon biases of the "highly expressed genes" of yeast and *Escherichia coli* (24, 29, 32, 54, 55). Translational accuracy and efficiency and codon-anticodon interaction strength may contribute to codon choices (1, 38). Selective and nonselective substitutional biases operating during DNA replication, transcription, and repair also play key roles. Gene codon usages to some extent correlate with functional categories (29, 32), as exemplified by polypeptide synthesis and chaperone-degradation activities. Other factors that may influence codon choices include methylation effects of DNA, mRNA stability, tissue and cellular location, codon context, and species of origin (30).

It is generally recognized that in most prokaryotic genomes during exponential growth, RPs and translation and transcription processing factors (TF) are highly expressed. The major chaperone-degradation (CH) genes functioning in protein folding, trafficking, and secretion are also largely highly expressed (e.g., data in reference 65). The three classes RP, CH, and TF are consistent in that they record congruent high codon biases relative to the average gene, whereas the codon usage differences among these three gene classes are low. Specifically, for rapid division, many ribosomes are indispensable, augmented by abundant TF and CH proteins needed to assure properly translated, modified, and folded protein products. These proteins expedite and regulate cellular activities. From this perspective, we have used the three classes RP, CH, and TF as representative classes of highly expressed genes.

A gene is predicted highly expressed (PHX) if the gene has codon frequencies similar to the codon frequencies of the RP, TF, and CH genes but deviates significantly in codon usage from the average gene of the genome (see "definition I" below for precision). PHX genes in most prokaryotic genomes include, in addition to those for RP, TF, and CH proteins, the principal genes of energy metabolism and key genes involved in amino acid, nucleotide, and fatty acid biosyntheses. In the

^{*} Corresponding author. Mailing address: Department of Mathematics, Stanford University, Stanford, CA 94305-2125. Phone: (650) 723-2204. Fax: (650) 725-2040. E-mail: fd.zgg@forsythe.stanford.edu.

TABLE 1. Statistics for highly expressed genes in diverse prokaryotic genomes

Genome (doubling time)	Length (kb)	No. of genes ≥ 100 aa	No. (%) PHX	$\max E(g)$	Reference
Eubacteria					
Fast growing $(<1 h)$					
E. coli	4,639	3,898	306 (8)	2.66	6
H. influenzae	1,830	1,529	142 (9)	2.01	15
B. subtilis	4,215	3.612	148 (4)	2.34	37
V. cholerae	4,036	3,253	172 (5)	2.25	
V. cholorae long chromosome	2,963	2,393	158 (7)	2.25	
Moderately fast (90 min)	,	,			
D. radiodurans	3,284	2,923	337 (12)	2.56	69
D. radiodurans long chromosome	2,649	2,421	307 (13)	2.56	
Obligate intracellular parasites	,	,			
R. prowazekii (10 h)	1,112	770	42 (5)	1.20	2
C. trachomatis (3 h)	1,043	829	52 (6)	1.27	59
C. pneumoniae (3 h)	1,230	963	85 (9)	1.38	26
Surface parasites	,				
M. genitalium	580	446	27 (6)	1.25	17
M. pneumoniae	816	654	57 (9)	1.36	23
Spirochetes					
B. burgdorferi (11–12 h)	1,231	1,007	76 (8)	1.29	16
T. pallidum (33 h)	1,138	916	99 (11)	1.35	18
Pathogens	,				
H. pylori 26695 (4–12 h)	1,668	1,388	73 (5)	1.21	63
M. tuberculosis (24 h)	4,412	3,660	569 (16)	1.68	9
Cyanobacteria Synechocystis (6–18 h)	3,573	2,896	380 (13)	1.51	27
Deep-branching gram-negative thermophiles	<i>,</i>	,			
A. aeolicus	1,551	1,481	233 (16)	1.62	11
T. maritima	1,861	1,681	175 (10)	1.40	49
Archaea					
Methanogens					
M. jannaschii (10 h)	1,665	1,466	114 (8)	1.59	7
M. thermoautotrophicum (4 h)	1,751	1,640	160 (10)	1.38	57
A. fulgidus (4 h)	2,178	2,077	343 (17)	1.49	34
Hyperthermophiles					
P. horikoshii	1,739	1,992	179 (9)	1.47	33
P. abyssi	1,765	1,683	242 (14)	1.60	

Synechocystis genome, many genes important in photosynthesis, respiration, and glycolysis are PHX, and among methanogens, those genes essential for methanogenesis are PHX.

MATERIALS AND METHODS

Data. PHX genes are identified across the 22 complete prokaryotic genomes listed in Table 1. Information on genome sizes, doubling times, and life styles (e.g., parasite versus free living or extremeophilic versus mesophilic) are indicated where the information is available.

Codon usage differences between gene classes. We previously introduced a versatile way of assessing the codon biases of one group of genes (or a single gene) relative to a second group of genes (29, 32). Let *G* be a group of genes with average codon frequencies g(x, y, z) for the codon triplet (x, y, z) normalized for each amino acid codon family such that $\sum_{(x,y,z)=a} g(x,y,z) = 1$, where the sum extends over all codons (x,y,z) translated to amino acid *a*. Let f(x,y,z) indicate the average codon frequencies for the gene group *F*, normalized to 1 in each amino acid codon family. The codon usage difference of the gene family *F* relative to the gene family *G* (codon bias relative to *G*) is calculated by the formula

$$B(F|G) = \sum_{a} p_{a}(F) \left[\sum_{(x,y,z) = a} |f(x,y,z) - g(x,y,z)| \right]$$
(1)

where $\{p_a(F)\}\$ are the average amino acid frequencies of the genes of *F*. When no ambiguity is likely, we refer to B(F|G) as the codon bias of *F* with respect to *G*. The assessments implied by equation 1 can be made for any two gene groups from the same genome or from different genomes. Equation 1 can also be applied to a subset of amino acids (e.g., restricted to hydrophobic, charged, or aromatic types).

Measures of gene expression. Let B(g|S), as above, denote the codon usage difference of the gene g relative to the gene class S as formalized in equation 1. The following gene classes are paramount: C, all protein genes; RP genes; CH genes; and TF genes. Qualitatively, a gene g is PHX if B(g|C) is high while B(g|RP), B(g|CH), and B(g|TF) are low. Predicted expression levels with respect to individual standards are based on the ratios

$$E_{\rm RP}(g) = \frac{B(g \mid C)}{B(g \mid \rm RP)}, E_{\rm CH}(g) = \frac{B(g \mid C)}{B(g \mid \rm CH)}, \text{ and } E_{\rm TF}(g) = \frac{B(g \mid C)}{B(g \mid \rm TF)}$$
(2)

Combined, these produce the general expression measure

$$E = E(g) = \frac{B(g|C)}{\frac{1}{2}B(g|RP) + \frac{1}{4}B(g|CH) + \frac{1}{4}B(g|TF)}$$
(3)

Other weighted combinations can also be used. For any class of genes S, a measure $E_S(g)$ for expression level of a gene g relative to S is calculated by B(g|C)/B(g|S).

Definition I. A gene is PHX if the following two conditions are satisfied: at least two among the three expression values $E_{\text{RP}}(g)$, $E_{\text{CH}}(g)$, and $E_{\text{TF}}(g)$ exceed 1.05, and the general expression level E(g) is ≥ 1.00 .

It is instructive to plot B(g|C) versus B(g|RP), B(g|TF), or B(g|CH) for all individual genes g (encoding proteins of \geq 100-amino-acid [aa] length). This is done in Fig. 1 for the E. coli genome and for the Deinococcus radiodurans genome. The distribution of points reveals two horns. The upper left horn corresponds to the PHX genes, and the upper right horn is designated "alien" genes. Alien genes consist mostly of open reading frames (ORFs) of unknown function but also include genes encoding transposases, cryptic prophage sequences, and restriction or modification enzymes, which are often conjugatively transferred via plasmids. Other examples of alien genes in several genomes are genes associated with lipopolysaccharide biosynthesis and fimbrial-gene-like genes (29, 32, 47). The term "alien" was chosen because such genes with high codon bias might have been acquired through recent lateral gene transfer (39, 40, 50). The formal definition and an extensive analysis of alien genes in diverse prokaryotic genomes will be presented elsewhere. The focus of this paper concerns identification and interpretation of PHX genes across the 22 prokaryotic genomes at hand. Table 2 highlights the primary PHX genes of the gene classes TF and CH.

RESULTS AND DISCUSSION

Statistics of PHX genes in prokaryotic genomes. Implementation of definition I provides lists of PHX genes for each



FIG. 1. Genes of ≥ 100 codons in the *E. coli* (left) and *D. radiodurans* (right) genomes. Each gene is represented by a single point. Its position is determined by its bias relative to all genes [B(g|C)] and by its bias relative to the RP genes [B(g|RP)] (top). B(g|RP) is replaced by codon bias relative to the CH standard [B(g|CH)] (middle) and relative to TF standard [B[g|TF)] (bottom). RP genes are shown in red, CH genes are in blue, and TF are in green.

prokaryotic genome. The global statistics are displayed in Table 1. For genes encoding proteins of at least 100-aa length, the percentages of PHX genes across the different genomes range from 4 to 17%. In particular, the fast-growing *E. coli* and Haemophilus influenzae genomes (doubling time, ≤ 1 h) contain 8 to 9% PHX genes. Chromosome I of Vibrio cholerae contains about 7% PHX genes, and Bacillus subtilis contains only about 4% PHX genes. D. radiodurans, exhibiting the dou-

TABLE 2. PHX genes in most bacterial genomes

Gene class	Proteins
RP	All large-subunit and small-subunit RPs
СН	GroEL (HSP60), DnaK (HSP70), GrpE, HtpG (HSP90), ClpP, ClpB, ClpC, ClpX, FtsH, HslU, trigger factor, PPIase, thioredoxin reductase, thermo-
Protein synthesis processing	somes in archaeal genomes
factors	EF-Tu (Tuf), EF-G (Fus), EF-Ts, IF-2, IF-3, ATP-dependent RNA polymerase factors (β , β' , α), RpoB, RpoC, RpoA, transcription terminator-antiterminator protein (NusA, NusB, NusG), ribosome release factor (Rrf)

bling time of 1 to 1.5 h, carries about 12 to 13% PHX genes. The slow-growing Mycobacterium tuberculosis (24- to 36-h doubling time) devotes 16% of its genome to PHX genes, with more than 80 of these genes acting in fatty acid biosynthesis or degradation. More than 40% of PHX genes in M. tuberculosis are ORFs of unknown function. The fraction of PHX genes of the archaeal genomes of Methanococcus jannaschii (~10-h doubling time) is about 8%, and that for Methanobacterium thermoautotrophicum (4-h doubling time) is about 10%. Thus, the proportion of PHX genes does not correlate with growth rate (doubling time) or with genome size. There are no consistent PHX gene proportions among hyperthermophiles: Aquifex aeolicus, 16%; Thermotoga maritima, 10%; Pyrococcus abyssi, 14%; Pyrococcus horikoshii, 9%; and M. jannaschii, 8%. The pathogens Rickettsia prowazekii, Chlamydia trachomatis, and Helicobacter pylori contain only 5 to 6% PHX genes with expression levels (equation 3) only reaching 1.20. The archaea show many unidentified PHX genes ranging from 21% of all PHX genes in M. thermoautotrophicum to 47% in P. horikoshii. The highest expression levels are achieved for genomes of rapidly dividing organisms.

Special PHX genes of diverse prokaryotic genomes. The complete lists of the PHX genes of the current complete genomes corresponding to Table 1 are available on our ftp site, gnomic.stanford.edu/pub/highlyexpressed. In this section, we

highlight special PHX genes in several of the prokaryotic genomes.

(i) *E. coli.* (Table 3). The polynucleotide phosphorylase (Pnp) gene attains the top expression level (E = 2.66) among all *E. coli* genes. Pnp is a multifunctional enzyme fundamental in RNA processing and mRNA degradation. The Pnp gene is also the gene with the highest expression level in *Borrelia burgdorferi* and is PHX in *H. influenzae*, *V. cholerae*, *Synechocystis*, *M. tuberculosis*, *Treponema pallidum*, *Chlamydia pneumoniae*, *A. aeolicus*, and *T. maritima*.

Peptidyl-prolyl cis-trans isomerases (PPIases) accelerate the folding of proteins. Their catalytic activity promotes cis-trans isomerization of proline bonds in oligopeptides. There are up to nine PPIases defined by sequence similarity in E. coli. The survival protein SurA version of PPIase concentrates on enhanced folding of periplasmic and outer membrane proteins. Some PPIases are PHX in H. influenzae, V. cholerae, B. subtilis, E. coli, D. radiodurans, H. pylori, R. prowazekii, M. tuberculosis, T. pallidum, A. aeolicus, P. horikoshii, and P. abyssi. Trigger factor is a ribosome-associated chaperone (exhibiting a PPIase activity) that can substitute for DnaK (12) and that also contributes to protein export. Trigger factor and DnaK cooperate in the folding of newly synthesized proteins. Simultaneous deletion of trigger factor and DnaK is lethal under the usual growth conditions (61). Trigger factor is broadly PHX, as seen in E. coli, H. influenzae, B. subtilis, V. cholerae, D. radiodurans, Synechocystis, T. pallidum, C. pneumoniae, and A. aeolicus.

The GAPDH (glyceraldehyde 3-phosphate dehydrogenase) protein is multifunctional, acting primarily in glycolysis; in eukaryotes it can also structurally bind actin filaments and microtubules. The GAPDH gene is PHX in almost all eubacterial genomes and also in the archaea *M. jannaschii* and *P. abyssi*. The data on expression levels of associated subunits of a protein complex are often variable. For example, in the UvrABC complex of *E. coli*, we find unit B to be PHX, whereas units A and C are not. Subunits may have activity separate from that of the total protein, which is the case with UvrB (56). From this perspective, the UvrB protein is multifunctional. By contrast, genes encoding ferredoxin oxidoreductase subunits in the genomes of *A. aeolicus*, *M. jannaschii*, *P. horikoshii*, and *T. maritima* tend to be incorporated in a single PHX operon at about equal expression levels.

TABLE 3. Prominent E. coli PHX genes

Function class	Description	Specific PHX genes
1	RPs	All RPs ≥ 100 as in length
2	Chaperonins and/or protein degradation	HSP70 (DnaK), HSP90 (HtpG), trigger factor, PPIase, HSP60 (GroEL), GroES, HslU, FtsH, thioredoxin, polynucleotide phosphorylase (mRNA degradation), protease DO (<i>htrA</i>)
3	Transcription and translation factors	EF-Tu (<i>tufA</i> ; duplicated), EF-G (<i>fusA</i>), EF-Ts, ATP-dependent RNA helicase, DNA-dependent RNA polymerase β and β', σ ⁷⁰ (<i>rpoB</i> , <i>rpoC</i> , <i>rpoD</i>), transcription terminator factor (<i>rho</i>), transcription antiterminator (NusG, NusB), ribosome release factor (<i>rrf</i>)
4	Detoxification	SodA, C, catalase, alkyl hydroperoxide reductase, thiol peroxidase, Dps (DNA protection during starvation)
5	Recombination and repair	RecA, single-stranded DNA binding protein (ssb)
6	Aminoacyl tRNA synthetases	ileS, proS, leuS, glnS, serS, asnS, pheS (α chain), pheT (β chain), thrS, aspS, argS, metG, gltX, alaS, lysS, valS, glyS (β chain), glyQ (α chain), glutamine amidotransferase (glmS)
7	Energy metabolism: mostly gly- colysis and tricarboxylic acid (Krebs) cycle and several enzymes of anaerobic growth	Pyruvate dehydrogenase, g3pd (glyceraldehyde 3-phosphate dehydrogenase), triosephosphate isomerase, fructose 1,6-bisphosphate aldolase, dihydrolipoamide dehydrogenase, phosphoglycerate mutase, 6-phos- phofructokinase aldolase, phosphate acetyltransferase, transketolase, deoxiribose-phosphate aldolase, succinyl-CoA ^{<i>a</i>} synthetase α and β , malate dehydrogenase, ATP synthetase F1 α and F1 β
8	Electron transport	Cytochrome <i>o</i> ubiquinol oxidase I, III, C; flavodoxin (<i>fldA</i>); cytochrome <i>d</i> ubiquinol oxidase I, II; fumarate and nitrate reductase I, II; NADH dehydrogenase subunits N, L, I, G, F, C, D; ferredoxin (<i>fdx</i>); NADH-nitrate oxidoreductase (<i>nirB</i>); fumarate reductase (<i>frdA</i> , B, D)
9	Outer membrane proteins	ompA, ompC, ompF, ompX, nmpC

^a CoA, coenzyme A.

The DNA binding protein Dps (DNA protection during starvation; labeled PexB in reference 43) is PHX (E = 1.13). Dps protects DNA from highly reactive oxygen radicals by forming a hollow spherical complex, where it sequesters DNA and iron to keep the reactive oxygen away (42). Not only is Dps highly expressed during rapid division, as shown by two-dimensional (2-D) gel assessments, it has been evaluated as being among the most abundant proteins in stationary phase (42).

(ii) **B.** subtilis. PHX genes of the *B.* subtilis genome parallel the PHX genes of *E.* coli. These include mainstream glycolysis and respiration genes and the detoxification genes sodA and the catalase and alkyl hydroperoxide reductase genes. The chaperone thioredoxin catalyzes or removes disulfide bonds in implementing protein folding. The highest predicted expression level for thioredoxin occurs in *B.* subtilis (E = 1.35), followed by those in the other fast-growing bacteria in the order *D.* radiodurans, (1.23), *V.* cholerae (1.21), *H.* influenzae (1.11), and *E.* coli (1.06). Thioredoxin (trxA) and thioredoxin reductase (trxB) ordinarily carry multiple copies, with at least one of these PHX in most eubacterial genomes.

In contrast to *E. coli*, four flagellin genes (flagellin [*hag*], flagellar hook protein [*flgE*], flagellar hook-basal body [*fliE*], and flagellin homolog [*yvzB*]) of *B. subtilis* are PHX, whereas a lone flagellin gene of *E. coli* is PHX. Why this difference? Based on the assumption that soil is the major *B. subtilis* habitat and the human gut is primary for *E. coli*, the habitat localization may be relevant. The movements of *B. subtilis* mediated by PHX flagellar proteins may facilitate its acquisition of food from soil sources. By contrast, nutrition (many sugars) flows easily to *E. coli* in the human lower intestine. Moreover, flagellar genes in *E. coli* are strictly regulated and inducible, but they are constitutive in *B. subtilis* (58).

(iii) D. radiodurans. A mesophilic bacterium, D. radiodurans can survive intense ionizing radiation at a dose of 5,000 Gy(4), which is lethal to virtually all other microorganisms. Such radiation causes DNA single- and double-strand breaks, generates DNA cross-links, and invokes a myriad of other types of DNA, RNA, and protein damage. It was hypothesized (69; but earlier in references 3 and 64) that although D. radiodurans possesses only the traditional prokaryotic repair repertoire, there probably are special mechanisms available that enhance repair. However, it seems paradoxical that recognized repair proteins show predominantly low expression levels, except for RecA (Table 4). RecA promotes and participates in many functions, including homologous recombination, DNA strand exchange, DNA repair and coprotease activity (reacting to DNA damage resulting in the SOS response), prophage induction, and/or mutagenesis subsequent to LexA cleavage (28, 36). Interestingly, RecA has the highest expression level in D. ra*diodurans* of all the genomes shown in Table 1.

Our view of the viability of *D. radiodurans* suggested by the predictions of gene expression levels emphasizes three processes: (i) the many degradation and export vehicles for removing damaged DNA, RNA, and proteins; (ii) the surfeit of chaperonins, which putatively enhance the operations of the repair proteins; and (iii) the manifold detoxification facilities that neutralize and remove free oxygen radicals and other toxic substances.

Strikingly, Table 5 shows that, compared to the other prokaryotic genomes, *D. radiodurans* contains the greatest number of PHX detoxification genes. The major PHX CH genes in *D. radiodurans* include those encoding GroEL (E = 2.35), DnaK (2.24), the general stress protein Ctc (1.89), Lon (two copies: 1.69 and 1.41), FtsZ (1.67), trigger factor (1.48), FtsH (three copies, but only FtsH-3 is PHX) (1.47), DNA binding stress response protein (Dps family) (1.51), cyclophilin (1.46), PPIase (four copies: 1.23 to 1.51), grpE (1.24), fimbrial assembly (pili) (1.23), septum cell division protein MinD (1.14), HSP20 (1.39), ribonuclease PH (1.06), ribonuclease H (1.02), thioredoxin (1.23), and GroES (1.14).

D. radiodurans is distinctive among prokaryotic genomes in having genes for PHX proteases of many kinds, including three serine protease (E = 1.29 to 1.41), protease I (PfpI) (1.46), zinc metalloendopeptidase (1.28), carboxyl-terminal protease (1.48), ClpX (1.32), and signal peptidase (1.08). FtsH, an integral inner membrane protein, is a cell division metalloprotease that facilitates degradation and protein folding (52). FtsH is PHX in *E. coli*, *D. radiodurans*, *Synechocystis*, *M. tuberculosis*, and *T. maritima*. The archaeal genomes commonly feature their own PHX cell division control protein. FtsZ, generally PHX in eubacteria, also contributes to bacterial cell division; it is tubulin-like and very abundant.

D. radiodurans PHX detoxification genes include two catalase genes (1.92 and 1.55), *sodA* (1.75), two *sodC* (1.44 and 1.05), Dps (1.51), chloride peroxidase (1.14), singlet oxygen resistance protein (1.11), MutT-nudix (1.29), organic hydroperoxide resistance protein (1.12), and tellurium resistance protein TerD (two copies) (1.47 and 1.05). SodA and catalase protect cells against toxin components induced by oxygen radicals. Alkyl hydroperoxide reductase serves to protect the cell against DNA damage caused by alkyl hydroperoxides. It reduces organic hydroperoxide to its dithiol form.

Virtually all of the DNA repair genes identified in D. radiodurans have functional analogs in other bacterial species. It is surmised that D. radiodurans possesses high redundancy in repair, but except for the nudix family (69) of nucleoside triphosphate pyrophosphorylases, this has not been verified. Actually, the degree of similarity claimed for the nudix genes is tenuous. Growing D. radiodurans cells contain 4 to 10 genome equivalents, which putatively provide facile opportunities for recombination and repair. However, most rapidly growing bacteria, e.g., E. coli and V. cholerae, contain at least three genome equivalents but fail to be resistant to ionizing radiation. Does D. radiodurans possess novel repair proteins more effective than those in other species or use the common repair machinery in new ways? None of this is established. On the contrary, Harsojo et al. (22) reported that with varied genome numbers in D. radiodurans there is no difference in radiation resistance.

We propose that the proliferation of PHX chaperones and degradation and detoxification proteins helps intrinsically in maintaining the survival and stability of the *D. radiodurans* cell subject to severe conditions of ionizing and/or UV radiation. We speculate that chaperones affect D. radiodurans cells when needed to expedite repair. Along these lines, there are precedents for chaperone influences that enhance the UvrA function in removing DNA damage during the process of nucleotide excision repair (75). Following massive UV radiation damage, D. radiodurans rapidly degrades at least 40% of its DNA and expels it from the cell. Faulty chromosomes could also be expelled, probably harmlessly. The chromosomes are fractured into many subgenomic fragments, but in less than 3 h, they can be accurately reassembled without loss of viability (3). The elaborate CH protein ensemble allows D. radiodurans to maintain the integrity of its essential macromolecules. Along these lines, D. radiodurans contains a preponderance of cell division, degrading, and recycling proteins. The proliferation of antioxidative stress proteins can putatively also mitigate desiccation and negative thermal effects. Most PHX genes of the small chromosome of D. radiodurans are duplicated in the large chromosome.

There are substantial parallels between the principal PHX genes of *E. coli* and *D. radiodurans* for the RP, TF, and CH

TABLE 4. Repair genes of D. radiodurans

0.52 1015 $1801990-$ Excinucless A ($urd+2$) 0.55 730 $1227809+$ Excinucless A ($urd+2$) 0.57 616 $1130069+$ Excinucless C ($urcC$) 0.52 744 $11844529+$ DNA helicase C ($urcC$) 0.52 744 $11844356+$ Uraci-DNA N_2 bycosplase (urg) 0.56 246 $170515+$ Uraci-DNA N_2 bycosplase (urg) 0.57 290 $493975+$ Formamidopyrimiline-DNA glycosplase ($uudy$) 0.65 362 $12281259 AG$ -specific admine glycosplase ($uudy$) 0.66 283 $135670-$ Exdocosyribonuclease III ($uth-1$) 0.76 $1999836-$ DNA mismath repair protein MuL ($uud.$) 0.45 765 $1999936-$ DNA mismath repair protein MuL ($uud.$) 0.46 283 $1282795-$ RecR protein ($recN$) 0.84 2910 $1202795-$ RecR protein ($recN$) 0.84 $1989030-$ recF protein ($recN$) 0.40 0.81 $1262795-$ RecR	E(g)	Length (aa)	Position ^a	Gene
0.52 $^{9}21$ $219705-$ Excinucless B ($urrA-2$) 0.55 730 $1227809+$ Excinucless E ($urrC$) 0.52 744 $180069+$ Excinucless E ($urrC$) 0.52 325 $11849685-$ Transcription-repair coupling factor (nfd) 0.52 325 $11844356+$ UV damage endonuclesse; putative (DR1819) 0.56 246 $1705155+$ $Urait-DNA$ hybrosylase (urg) 0.57 290 $1433975+$ Formanidopyrimidine-DNA glycosylase (urg) 0.56 362 $1223229 AG$ -specific denine glycosylase (urg) 0.86 224 $1291364+$ Endonuclease III ($ufh-1$) 0.79 258 $12349683-$ Endonuclease III ($ufh-2$) 0.54 546 $1137575+$ DNA mismatch repair protein MuL (uuL) 0.55 538 $119990850-$ DNA repair protein MuL (uuL) 0.54 563 $11990295-$ Reck protein ($vecN$) 0.55 415 $11995908-$ DNA repair protein MuL (muL) 0.55 415 194	0.76	1.015	1 1801990-	Excinuclease A (uvrA-1)
0.55 700 $1227809+$ Excinucless C ($urcC$) 0.57 616 $1360069+$ Excinucless C ($urcC$) 0.52 744 $1804529+$ DNA helicase C ($urcC$) 0.52 325 $11844356+$ Urascription-repair coupling factor (mfd) 0.55 226 $1795155+$ Urascription-repair coupling factor (mfd) 0.57 290 $1493975+$ Formanicopyrimidine-DNA glycosylase (mug) 0.57 290 $1493975+$ Formanicopyrimidine-DNA glycosylase (mud) 0.56 362 $1223329 A'G$ ($Sepectific adenic glycosylase (mud) 0.56 362 1223329 A'G (Sepectific adenic glycosylase (mud) 0.56 362 1223329 A'G (Sepectific adenic glycosylase (mud) 0.56 362 123354 DA (Nainsmatch repair protein mud) 0.57 290 123354 DNA mismatch repair protein mud) 0.56 199886 DNA mismatch repair protein Mut; (mud) 0.58 358 119902795- RecR protein (recA)$	0.52	921	2 197055-	Excinuclease A $(uvrA-2)$
0.52 744 $1560069+$ Excinucless C ($urcl$) 0.52 744 $1540685-$ Transcription-repair coupling factor ($mlcl$) 0.52 225 $11549685-$ Transcription-repair coupling factor ($mlcl$) 0.56 246 $1705155+$ $Urail-DNA$ helicase; putative (DR1819) 0.56 246 $1705155+$ $Urail-DNA$ helicase; putative (DR1819) 0.56 362 $1232329 AG$ -specific adening elycosylase (mug) 0.56 362 $1232829 AG$ -specific adening elycosylase (mud) 0.86 224 $129164+$ Endonuclease III (mhc) 0.79 258 $12349683-$ Endonuclease III (mhc) 0.54 546 $117875+$ DNA mismatch repair protein MutL (muL) 0.54 563 $1199985-$ DNA repair protein (mcN) 0.58 583 $1199998-$ DNA repair protein (mcN) 0.51 563 $1192959-$ Ref protein (mcA) 0.54 245 $123795+$ reck protein (mcA) 0.5	0.55	730	12272809 +	Excinuclease B $(uvrB)$
0.52 744 1894829^+ DNA helices U (arD) 0.58 1.053 11844356^+ UV damage endonuclease; putative (DR19) 0.56 246 1705155^+ Ur caci-DNA glycosylase (mug) 0.64 198 173545^- GU minitch-specific DNA glycosylase (mug) 0.57 290 1493975^+ Formanidopyrindine-DNA glycosylase (mug) 0.57 290 $1233259^ AG$ -specific adenine glycosylase (mud) 0.65 362 $1233259^ AG$ -specific adenine glycosylase (mud) 0.86 224 1291364^+ Endonuclease III (mL^-1) $0.79^ 228$ 1239083^- Endonuclease III (mL^+2) 0.54 136700^- Exodeoxyribonuclease III (mL^+2) 0.54 119992856^- DNA mismath repair protein MuL (mud) 0.51 563 11490126^+ DNA respective (eC) 0.40 823 11292060^- Exodeoxyribonuclease V: subunit RecD; putative (RP) 0.49 714 11922060^- Exodeoxyribonuclease V: subunit RecD; putative (RP) 0.55 415 1944494^+ <td< td=""><td>0.57</td><td>616</td><td>11360069+</td><td>Excinuclease C $(uvrC)$</td></td<>	0.57	616	11360069+	Excinuclease C $(uvrC)$
0.58 1.033 $1.549685 Transcription-repair coupling factor (mfd)$ 0.52 325 $1.844356 +$ UV damage endouclease; putative (DR1819) 0.56 246 $1.705155 +$ $Uracil-DNA$ Arglycoylase (mag) 0.64 198 $1.73545 G/U$ mismath-specific DNA glycoylase (mag) 0.57 200 $1.493975 +$ Formamidopyrinding-DNA glycoylase (mag) 0.65 362 $1.228325 AG-specific adenine glycoylase (mag)$ 0.86 224 $1.213545 -$ Endonuclease III (ml-1) 0.79 258 $1.2439683 -$ Endonuclease III (ml-1) 0.76 $1.78755 +$ DNA mismatch repair protein MuL (muL) 0.45 765 $1.999856 -$ DNA repair protein (mcR) 0.45 765 $1.099856 -$ DNA neismatch repair protein MuL (muL) 0.45 765 $1.999856 -$ DNA neismatch repair protein MuL (muL) 0.46 $2.337955 -$ RecR protein (mcR) 0.40 0.51 563 $1.19224060 -$ Exonuclease ShcD; putative (DR1921)	0.52	744	1 1804529+	DNA helicase II $(uvrD)$
322 325 $1844336+$ UV damBy endowices; purative (DR(819)) 0.56 246 $1705155+$ $UrachDyA$ regionalizes; purative (DR(819)) 0.64 198 $1735345 G(U$ mismatch-specific DNA glycosylase (mug) 0.57 200 $1493975+$ Formanidopyrinidine-DNA glycosylase (mug) 0.56 362 $12233259-$ AGC-specific adening glycosylase (mut) 0.86 224 $1291364+$ Endonuclease III (mh-1) 0.79^{-2} 258 $12439683-$ Endonuclease III (mh-1) 0.66 233 $1356790-$ Exodeoxyribonuclease III (mh-1) 0.54 546 $1178755+$ DNA mismatch repair protein MuL (muL) 0.58 358 $10990856-$ DNA mismatch repair protein MuL (muL) 0.80 219 $120795-$ RecR protein (recR) 0.40 823 $11292060-$ Exodeoxyribonuclease V: subunit RecD; putative 0.71^{-1} 908 $11942746+$ Exonuclease SbcC (shcC) 0.55 415 $11941499+$ Exonuclease SbcC (shcC) 0.69^{-1} 200 $11281494-$ Ho	0.58	1.053	1 1549685-	Transcription-repair coupling factor (<i>mfd</i>)
0.56 246 170:155+ Uracli DNA N_{el} prospiase (mg) 0.64 198 173:545- Gradi mismatch-specific DNA glycosylase (mut) 0.57 290 149:3975+ Formanidopyrinidine-DNA glycosylase (mut) 0.65 362 1223:259- A/G-specific adenine glycosylase (mut) 0.86 224 121:364+ Endonuclease III $(mt-2)$ 0.66 283 13:56790- Exodeoxyribonuclease III $(mt-2)$ 0.45 765 11999856- DNA mismatch repair protein MutS; putative $(mutS)$ 0.51 563 11990856- DNA repair protein (meN) 0.51 563 11990856- DNA repair protein (mutS; putative $(mutS)$ 0.40 823 11295908- DNA helicase Rec0 (rec0) 0.40 823 11292906- Exodeoxyribonuclease V; subunit Rec1; putative $(mutS)$ 0.71 908 1194:1499+ Econuclease Sbc1; putative $(DR1921)$ 0.44 123:1494+ Holiday junction binding protein (meA) 0.55 1178 143:818+ Holiday junction DNA helicase (meC) 0.56	0.52	325	1 1844356+	UV damage endonuclease: putative (DR1819)
0.64 198 1733847 GU mismatch-specific DNA glycosylase (mug) 0.57 290 1493975 + Formanidopyrinidine-DNA glycosylase (mud) 0.65 362 1283259 - Λ/G -specific adenine glycosylase (mud) 0.86 224 1213644 + Endonuclease III (ml-1) 0.79 258 12439683 - Endonuclease III (ml-2) 0.66 283 1356700 - Exodeoxyribonuclease III (ml-2) 0.54 546 11718755 + DNA mismatch repair protein Mut1 (mud1) 0.54 563 11999856 - DNA mismatch repair protein Mut1 (mud2) 0.58 358 11098030 - recf protein (rec P) 0.40 823 11292090 - Exodeoxyribonuclease III (ml-2) 0.40 823 11922060 - Exodeoxyribonuclease Rec0 (rec) 0.40 823 11922060 - Exodeoxyribonuclease Rec) (rec) 0.49 714 11922060 - Exodeoxyribonuclease Rec) (rec) 0.49 $212746+$ Exonuclease Shc) (putative (DR1921) 0.40	0.56	246	$1.705155 \pm$	Uracil-DNA <i>N</i> -glycosylase (<i>ung</i>)
0.57 290 $1493975+$ Formanidopyrinidine-DNA glycosylase (matM) 0.65 362 $1223325-$ A/G-specific adenine glycosylase (matM) 0.86 224 $121364+$ Endonuclease III (mh-2) 0.79 258 $12439683-$ Endonuclease III (mh-2) 0.66 283 $1556790-$ Exodeoxyribonuclease III (mh-2) 0.45 765 $11999856-$ DNA mismatch repair protein MutS; putative (mutS) 0.51 563 $11999856-$ DNA repair protein (mcN) 0.80 219 $1202795-$ RecR protein (mcN) 0.81 $1922060-$ Exodeoxyribonuclease V; subunit RecD; putative 0.71 908 $11942746+$ Exonuclease Stoc); putative (DR1921) 2.04 ⁴⁰ 362 $1233795+$ recA protein (mcA) 0.69 200 $123494-$ Holiday junction binding protein (mcA) 0.69 200 $123494-$ Holiday junction binding protein (mcA) 0.69 200 $123494-$ Holiday junction binding protein (mcA) 0.60 332 $1609449-$	0.64	198	1 733545-	G/U mismatch-specific DNA glycosylase (<i>mug</i>)
0.65 362 1283259- A/G-specific adenine glycospiae (muY) 0.86 224 1291364+ Endonuclease III $(mh-1)$ 0.79 238 12439683- Endonuclease III $(mh-2)$ 0.66 283 1356700- Exodeoxyribonuclease III $(mh-2)$ 0.54 546 11718755+ DNA mismatch repair protein MuL((muL) 0.45 765 11999856- DNA mismatch repair protein MuL((muL) 0.51 563 11490126+ DNA repair protein $(mecN)$ 0.80 219 1202795- ReCR protein $(mecN)$ 0.40 823 11295908- DNA hielicase RecQ $(recQ)$ 0.41 1922060- Exonuclease SbcC (bcC) 2019 0.71 908 11942746+ Exonuclease SbcC (bcC) 2020 0.55 415 11941499- Houlday junction binding protein (mvA) 106 0.54 15 1941494- Holliday junction POA helicase (mvB) 0.88 0.78 178 1436818+ Holliday junction DNA helicase (mvB) 0.85 0.57 955 11732431- DNA-diffected DNA polymerase (moL) <td< td=""><td>0.57</td><td>290</td><td>1493975 +</td><td>Formamidopyrimidine-DNA glycosylase (<i>mutM</i>)</td></td<>	0.57	290	1493975 +	Formamidopyrimidine-DNA glycosylase (<i>mutM</i>)
0.86 224 121364+ Endonuclease III (nh^2) 0.79 258 12439683- Endonuclease III (nh^2) 0.66 283 1356790- Exodeoxyribonuclease III (nh^2) 0.54 546 11718755+ DNA mismatch repair protein MuL $(nud.)$ 0.45 765 11999856- DNA mismatch repair protein MuL; putative $(nud.)$ 0.51 563 11490126+ DNA repair protein $(necN)$ 0.80 219 1202795- RcR protein $(necR)$ 0.40 823 11225008- DNA helicase RecQ $(necQ)$ 0.44 1922060- Exonuclease SbcC $(shcC)$ 0.80 0.71 908 11942746+ Exonuclease SbcC $(shcC)$ 0.80 0.69 200 11281494- Holiday junction DNA helicase $(nw.8)$ 0.88 0.73 1894189+ Holiday junction DNA helicase $(nw.8)$ 0.87 0.87 985 11732431- DNA delicase RecG $(necG)$ 0.57 955 11732431- DNA delicase RecG $(necG)$ 0.66 1991985- MutT/nudix f	0.65	362	1 2283259-	A/G-specific adenine glycosylase $(mutY)$
0.79 258 $12439683-$ Endomclease III (<i>wh-2</i>) 0.66 283 $1356790-$ Exodeoxyribonuclease III (<i>wh-2</i>) 0.54 546 $11718755+$ DNA mismatch repair protein Mut1 (<i>mul1</i>) 0.45 765 $19999856-$ DNA mismatch repair protein Mut1 (<i>mul1</i>) 0.51 563 $11490126+$ DNA repair protein (<i>recN</i>) 0.80 219 $1202795-$ RccR protein (<i>recN</i>) 0.40 823 $1129508-$ DNA helicase RecQ (<i>recQ</i>) 0.49 714 $11922060-$ Exodeoxyribonuclease V; subunit RecD; putative 0.71 908 $11942746+$ Exonuclease SbcC (<i>sleCC</i>) 0.55 415 $19441499+$ Exonuclease SbcC (<i>sleCC</i>) 0.69 200 $1281494-$ Holliday junction brach repair protein (<i>recN</i>) 0.69 200 $1281494-$ Holliday junction protein (<i>recN</i>) 0.57 955 $11732431-$ DNA helicase (<i>rucO</i>) 0.57 955 $11732431-$ DNA ilgase (<i>dnH)</i> 0.76	0.86	224	1 291364+	Endonuclease III (<i>nth-1</i>)
0.66 283 1 356790 – Exodeoxyribonuclease III (xhA) 0.54 546 11718755 + DNA mismatch repair protein MutL (mutL) 0.45 765 1999856 – DNA mismatch repair protein MutS; putative (muS) 0.58 358 1090800 – recF protein (recF) 0.51 563 11490126 + DNA repair protein (meR) 0.40 823 1295908 – DNA helicase RecQ (recQ) 0.49 714 11922060 – Exodeoxyribonuclease V; subunit RecD; putative 0.71 908 1942746 + Exonuclease SbcC; putative (DR1921) 2.04 ^b 362 1233795 + recA protein (recA) 0.69 200 11281494 - Holiday junction DNA helicase (racV) 0.69 200 1281494 - Holiday junction DNA helicase (racQ) 0.55 11732431 - DNA-dricease BccC (recC) 0.56 783 1936653 + DNA helicase (racQ) 0.61 699 12084444 - DNA ligase (dnI) 0.76 170 180637 - MutT/nudix family protein (DR079)	0.79	258	1 2439683-	Endonuclease III (<i>nth-2</i>)
0.54 546 1 1718755 + DNA mismatch repair protein MutL ($nutL$) 0.45 765 11999856 - DNA mismatch repair protein MutS; putative ($nutS$) 0.58 358 11998030 - recF protein ($recR$) 0.51 563 11490126 + DNA repair protein ($recR$) 0.40 823 11295908 - DNA helicase RecQ ($recQ$) 0.49 714 11922060 - Exocovyribonuclease SbcC ($sbcC$) 0.55 415 11941499 + Exonuclease SbcC ($sbcC$) 0.40 362 1233795 + recA protein ($recA$) 0.69 200 1281494 - Holiday junction bnA helicase ($reuB$) 0.55 415 1944499 + Holiday junction bnA helicase ($reuB$) 0.60 332 1609449 - Holiday junction bnA helicase ($reuB$) 0.57 955 11732431 - DNA helicase ($reuC$) 0.56 783 11936853 + DNA helicase ($reuB$) 0.61 699 2084844 - DNA ligase (dul) 0.76 177 1276905 + MutT/nudix family protein	0.66	283	1 356790-	Exodeoxyribonuclease III (<i>xthA</i>)
0.45 765 $1999856-$ DNA mismatch repair protein MutS; putative (mutS) 0.58 358 $1109010-$ rccF protein (recF) 0.51 563 $11490126+$ DNA repair protein (recR) 0.40 823 $1129508-$ DNA helicase RecQ (recQ) 0.49 714 $11922060-$ Exodeoxyribonuclease V; subunit RecD; putative 0.71 908 $11942746+$ Exonclease SbcC (sbcC) 0.55 415 $11941499+$ Exonuclease SbcC (sbcC) 0.69 200 $1281494-$ Holliday junction IDNA helicase (nurB) 0.69 200 $1281494-$ Holliday junction TNA helicase (nurB) 0.60 332 $1609449-$ Holliday junction IDNA helicase (nurB) 0.50 783 $1395653+$ DNA helicase RecG (recG) 0.57 955 $11732431-$ DNA directed DNA polymerase (polA) 0.61 699 $1284844-$ DNA ligase (dnU) 0.76 170 $189637-$ MutT/nudix family protein (DR079) 0.76 <t< td=""><td>0.54</td><td>546</td><td>1 1718755+</td><td>DNA mismatch repair protein MutL (<i>mutL</i>)</td></t<>	0.54	546	1 1718755+	DNA mismatch repair protein MutL (<i>mutL</i>)
0.58 358 1 1098030- recF protein (recF) protein (precN) 0.51 563 1 1490126+ DNA repair protein (recN) 0.80 219 1 202795- RecR protein (recR) 0.40 823 1 129508- DNA helicase RecQ (recQ) 0.49 714 1 1922060- Exodeoxyribonuclease V: subunit RecD; putative 0.71 908 1 1942746+ Exonuclease SbcC (sbcC) 0.55 415 1 1941499+ Exonuclease SbcD; putative (DR1921) 0.69 200 1 1281494- Holliday junction binding protein (nvA) 1.06 332 1 609449- Holliday junction resolvase (nvC) 0.59 783 1 936853+ DNA helicase RecG (recG) 0.57 955 1 1732431- DNA-chirected DNA polymerase (pol/4) 0.61 699 1 2084844- DNA ligase (dnU) 0.76 170 1 80637- MutT/nudix family protein (DR079) 0.76 177 1 276905+ MutT/nudix family protein (DR079) 0.73 166 1 991985- MutT/nudix family protein (DR079) 0.74 155 1 1807313- <td>0.45</td> <td>765</td> <td>1 1999856-</td> <td>DNA mismatch repair protein MutS: putative (<i>mutS</i>)</td>	0.45	765	1 1999856-	DNA mismatch repair protein MutS: putative (<i>mutS</i>)
0.51 563 1 490126+ DNA repair protein (recN) 0.80 219 1 202795 – RecR protein (recN) 0.40 823 1 1295988 – DNA helicase RecQ (recQ) 0.49 714 1 1922060 – Exodeoxyribonuclease V; subunit RecD; putative 0.71 908 1 1942746 + Exonuclease SbcC; putative (DR1921) 2.04 ^b 362 1 2337795 + recA protein (recA) 0.69 200 1 1281494 - Holliday junction binding protein (nvA) 1.06 332 1 609449 - Holliday junction box helicase (nvA) 0.50 783 1 356853 + DNA helicase RecG (recG) 0.57 955 1 1732431 - DNA helicase RecG (recG) 0.76 177 1 276905 + MutT/nudix family protein (DR0079) 0.76 177 1 276905 + MutT/nudix family protein (DR079) 0.73 166 1 991985 - MutT/nudix family protein (DR075) 0.73 166 1 991985 - MutT/nudix family protein (DR077) 0.78 175 1 1807313 - MutT/nudix family protein (DR1025) 0.73 166 1 991985	0.58	358	1 1098030-	recF protein (recF)
0.80 219 1 202795 – RecR protein (recR) 0.40 823 1 1295908 – DNA helicase RecQ (recQ) 0.49 714 1 922060 – Exodeoxyribouclease V; subunit RecD; putative 0.71 908 1 1942746 + Exonuclease SbcC (sbcC) 0.55 415 1 941499 + Exonuclease SbcC) putative (DR1921) 0.69 200 1 1281494 - Holliday junction binding protein (nvA) 1.06 332 1 609449 - Holliday junction DNA helicase (nvB) 0.88 178 1 436818 + Holliday junction resolvase (nvC) 0.57 955 1 1732431 - DNA directed DNA polymerase (polA) 0.61 699 1 2084844 - DNA liagse (dnU) 0.76 177 1 276905 + MutT/nudix family protein (DR079) 0.76 177 1 276905 + MutT/nudix family protein (DR079) 0.73 166 1 991985 - MutT/nudix family protein (DR077) 0.83 1 3038101 + MutT/nudix family protein (DR077) 0.84 1 019945 + MutT/nudix family protein (DR1077)	0.51	563	11490126 +	DNA repair protein (recN)
0.40 823 1 1205908- DNA helicase RecQ (recQ) 0.49 714 1 1922060- Exodeoxyribonuclease V; subunit RecD; putative 0.55 415 1 1941499+ Exonuclease SbcC (sbcC) 0.69 200 1 1281494- Holliday junction binding protein (nwA) 1.06 332 1 609449- Holliday junction DNA helicase (nwB) 0.88 178 1 436818+ Holliday junction DNA helicase (nwC) 0.50 783 1 1930853+ DNA helicase RecG (recG) 0.57 955 1 1732431- DNA direcase RecG (net C) 0.61 699 1 2084844- DNA ligase (nuC) 0.62 322 1 32437- MutT/nudix family protein (DR0079) 0.76 170 1 80637- MutT/nudix family protein (DR0079) 0.76 177 1 276905+ MutT/nudix family protein (DR079) 0.73 166 1 991985- MutT/nudix family protein (DR075) 0.73 165 1 1038101+ MutT/nudix family protein (DR107) 0.83 158 1 1038101+ MutT/nudix	0.80	219	1 202795-	RecR protein (recR)
0.49 714 $1922060-$ Exodeoxyribonuclease V; subunit RecD; putative 0.71 908 $11942746+$ Exonuclease SbcC ($sbcC$) 0.55 415 $11941499+$ Exonuclease SbcD; putative (DR1921) 2.04^{b} 362 $12337795+$ recA protein ($recA$) 0.69 200 $11281494-$ Holliday junction binding protein (nwA) 1.06 332 $1609449-$ Holliday junction NA helicase (nwB) 0.88 178 $1436818+$ Holliday junction resolvase (nwC) 0.50 783 $11936853+$ DNA helicase RecG ($recG$) 0.57 955 $11732431-$ DNA-directed DNA polymerase ($polA$) 0.61 699 $12084844-$ DNA ligase (dnU) 0.76 170 $180637-$ MutT/nudix family protein (DR0079) 0.76 177 $127695+$ MutT/nudix family protein (DR0079) 0.76 177 $127695+$ MutT/nudix family protein (DR0329) 0.60 249 $155381+$ MutT/nudix family protein (DR107) 0.83 158 $1103945+$ MutT/nudix family protein (DR107) 0.78 175 $1807313-$ MutT/nudix family protein (DR107) 0.79 711 $12270912+$ MutT/nudix family protein (DR2272) 1.29^{b} 143 $12354268+$ MutT/nudix family protein (DR2272) 0.83 354 $312890+$ Ribonucleoside-diphosphate reductase; hota ($nrdE$) 0.87 502 $11112934+$ DNA repair protein ($ndA1$) 0.52 940	0.40	823	1 1295908-	DNA helicase RecO (recO)
0.71 908 $11942746+$ Exonuclease SbcC (sbcC) 0.55 415 $11941499+$ Exonuclease SbcD; putative (DR1921) 0.69 200 $11281494-$ Holliday junction binding protein (nvA) 1.06 332 $1609449-$ Holliday junction DNA helicase (nvB) 0.88 178 $436818+$ Holliday junction colvase (nvC) 0.50 783 $11936853+$ DNA helicase (creC) 0.57 955 $11732431-$ DNA helicase (nvB) 0.61 699 $20084844-$ DNA ligase (dnI) 0.76 170 $180637-$ MutT/nudix family protein (DR079) 0.76 177 $1276905+$ MutT/nudix family protein (DR079) 0.62 322 $1224367-$ MutT/nudix family protein (DR079) 0.76 177 $1276905+$ MutT/nudix family protein (DR079) 0.73 166 $199185-$ MutT/nudix family protein (DR075) 0.94 165 $1019945+$ MutT/nudix family protein (DR107) 0.83 158 $1038101+$ MutT/nudix family protein (DR107) 0.84 $12270912+$ MutT/nudix family protein (DR2272) 1.29^{b} 143 $122599+$ Ribonucleoside-diphosphate reductase; beta (nrdF) 0.89 140 $312650+$	0.49	714	1 1922060-	Exodeoxyribonuclease V: subunit RecD: putative
0.554151 941499+Exonuclease SbcD; putative (DR1921) 2.04^{b} 3621 2337795+recA protein (recA)0.692001 1281494-Holliday junction binding protein (ravA)1.063321 609449-Holliday junction DNA helicase (ravB)0.881781 436818+Holliday junction resolvase (ravC)0.507831 1936853+DNA helicase EceG (recG)0.579551 1732431-DNA-directed DNA polymerase (polA)0.616991 2084844-DNA ligase (dnII)0.761701 80637-MutT/nudix family protein (DR0079)0.761771 276905+MutT/nudix family protein (DR0274)0.623221 324367-MutT/nudix family protein (DR0550)0.731661 991985-MutT/nudix family protein (DR075)0.941651 1019945+MutT/nudix family protein (DR1025)0.781751 1807313-MutT/nudix family protein (DR1025)0.781711 2270912+MutT/nudix family protein (DR2272)1.29 ^b 1431 2354268+MutT/nudix family protein (DR2356)0.597013 126894+Ribonucleoside-diphosphate reductase; alpha (nrdE)0.583543 128900+Ribonucleoside-diphosphate reductase; beta (nrdF)0.875021 1112934+DNA repair protein (RadA)0.529403 162296+ATP-dependent helicase HepA (hepA)0.554031 233583+CinA protein (RadA)0.597013 126894+ <td< td=""><td>0.71</td><td>908</td><td>1 1942746+</td><td>Exonuclease SbcC (<i>sbcC</i>)</td></td<>	0.71	908	1 1942746+	Exonuclease SbcC (<i>sbcC</i>)
2.04b3621 2337795 +recA protein (recA)0.692001 1281494 -Holliday junction binding protein (nvA)1.063321 609449 -Holliday junction binding protein (nvA)0.881781 436818 +Holliday junction resolvase (nvC)0.507831 1936853 +DNA helicase RecG (recC)0.579551 1732431 -DNA-directed DNA polymerase (polA)0.616992 2084844 -DNA ligase (dnU)0.761701 80637 -MutT/nudix family protein (DR079)0.761771 276905 +MutT/nudix family protein (DR0274)0.623221 324367 -MutT/nudix family protein (DR0329)0.602491 555381 +MutT/nudix family protein (DR0350)0.731661 991985 -MutT/nudix family protein (DR1075)0.831581 1038101 +MutT/nudix family protein (DR1075)0.591711 2270912 +MutT/nudix family protein (DR2272)1.29b1431 2354268 +MutT/nudix family protein (DR2275)0.583543 128900 +Ribonucleoside-diphosphate reductase; alpha (nrdE)0.575021 1112934 +DNA repair protein (ncdA)0.529403 162296 +AT repersor (lexA)0.534333 124398 +Integrase-recombinase XerD; putative (xerD)	0.55	415	1 1941499+	Exonuclease SbcD: putative (DR1921)
0.69 200 1 1281494 – Holliday junction binding protein (nvA) 1.06 332 1 609449 – Holliday junction DNA helicase (nvB) 0.88 178 1 436818 + Holliday junction resolvase (nvC) 0.50 783 1 936853 + DNA helicase RecG ($veCG$) 0.57 955 1 1732431 - DNA-directed DNA polymerase ($polA$) 0.61 699 1 2084844 - DNA higase (dnU) 0.76 170 1 80637 - MutT/nudix family protein (DR0079) 0.76 177 1 276905 + MutT/nudix family protein (DR0274) 0.60 249 1 555381 + MutT/nudix family protein (DR0329) 0.60 249 1 555381 + MutT/nudix family protein (DR0750) 0.73 166 1 991985 - MutT/nudix family protein (DR1070) 0.83 158 1 1038101 + MutT/nudix family protein (DR1071) 0.84 12257012 + MutT/nudix family protein (DR2756) 0.78 175 1 1807313 - MutT/nudix family protein (DR272) 0.59 701 3 126894 +	2.04^{b}	362	1 2337795+	recA protein (<i>recA</i>)
1.063321 609449Holliday junction DNA helicase (nvB) 0.881781 436818 +Holliday junction resolvase (nvC) 0.507831 1936853 +DNA helicase RecG $(recG)$ 0.579551 1732431 -DNA-directed DNA polymerase $(polA)$ 0.616991 2084844 -DNA digase (dnU) 0.761701 80637 -MutT/nudix family protein (DR079)0.761771 276905 +MutT/nudix family protein (DR079)0.623221 324367 -MutT/nudix family protein (DR0550)0.731661991985 -MutT/nudix family protein (DR0755)0.941651 1019945 +MutT/nudix family protein (DR107)0.831581 1038101 +MutT/nudix family protein (DR107)0.831581 1038101 +MutT/nudix family protein (DR272)1.29b1711 2270912 +MutT/nudix family protein (DR2272)1.29b1731 26894 +Ribonucleoside-diphosphate reductase; alpha (nrdE)0.522192 379822 -LexA repressor (lexA)0.529403 126505 +Ribonucleoside-diphosphate reductase; beta (nrdF)0.611,0663 78759 +Extracellular nuclease; putative (DRB0107)0.529403 162505 +ATP-dependent helicase HepA (hepA)0.529403 162206 +ATP-dependent helicase HepA (hepA)0.529403 162505 +Extracellular nuclease; putative (DRB0067)1.021431 100371 +Single-stranded DNA binding protein (ssb) <td>0.69</td> <td>200</td> <td>1 1281494-</td> <td>Holliday junction binding protein ($ruvA$)</td>	0.69	200	1 1281494-	Holliday junction binding protein ($ruvA$)
0.881781 436818+Holliday junction resolvase $(nuC)^{+}$ 0.507831 1936853+DNA helicase RecG $(recG)^{-}$ 0.579551 1732431-DNA-directed DNA polymerase $(polA)^{-}$ 0.616991 2084844-DNA ligase $(dnl)^{-}$ 0.761701 80637-MutT/nudix family protein (DR079)0.761772 76905+MutT/nudix family protein (DR0329)0.602491 555381+MutT/nudix family protein (DR0550)0.731661 991985-MutT/nudix family protein (DR1007)0.831581 1038101+MutT/nudix family protein (DR1007)0.781751 1807313-MutT/nudix family protein (DR1025)0.791711 2270912+MutT/nudix family protein (DR2272)1.29 ^b 1431 2354268+MutT/nudix family protein (DR2356)0.597013 126894+Ribonucleoside-diphosphate reductase; alpha (nrdE)0.891403 12505+Ribonucleoside-diphosphate reductase; beta (nrdF)0.8750211112934+DNA repair protein (radA)0.529403 162296+ATP-dependent helicase HepA (hepA)0.611,0663 78759+Extracellular nuclease; putative (DRB0067)1.021431 100371+Single-stranded DNA binding protein (pR238)0.833433 124398+Integrase-recombinase XerD; putative (xerD)	1.06	332	1 609449-	Holliday junction DNA helicase $(ruvB)$
0.507831 1936853+DNA helicase RecG (recG)0.579551 1732431-DNA-directed DNA polymerase (polA)0.616991 208484-DNA ligase (dnlJ)0.761701 80637-MutT/nudix family protein (DR079)0.761771 276905+MutT/nudix family protein (DR0329)0.602491 555381+MutT/nudix family protein (DR0550)0.731661 991985-MutT/nudix family protein (DR107)0.831581 1019945+MutT/nudix family protein (DR107)0.831581 1038101+MutT/nudix family protein (DR107)0.591711 2270912+MutT/nudix family protein (DR2272)1.29 ⁶ 1431 2354268+MutT/nudix family protein (DR2356)0.597013 126894+Ribonucleoside-diphosphate reductase; alpha (nrdE)0.583543 128900+Ribonucleoside-diphosphate reductase; alpha (nrdE)0.622192 379822-LexA repressor (lexA)0.875021112934+DNA repair protein (madA)0.529403 162206+ATP-dependent helicase HepA (hepA)0.611,0663 78759+Extracellular nuclease; putative (DRB0067)1.021431 100371+Single-stranded DNA binding protein (lossb)0.654031 233583+CinA protein (DRA238)0.833433 124398+Integrase-recombinase XerD; putative (xerD)	0.88	178	1 436818+	Holliday junction resolvase $(ruvC)$
0.57 955 $11732431-$ DNA-directed DNA polymerase (pol4) 0.61 699 $12084844-$ DNA ligase (dnll) 0.76 170 $180637-$ MutT/nudix family protein (DR0079) 0.76 177 $1276905+$ MutT/nudix family protein (DR0274) 0.62 322 $1324367-$ MutT/nudix family protein (DR0550) 0.60 249 $1555381+$ MutT/nudix family protein (DR0570) 0.73 166 $1991985-$ MutT/nudix family protein (DR1007) 0.83 158 $10031045+$ MutT/nudix family protein (DR1007) 0.83 158 $1038101+$ MutT/nudix family protein (DR1025) 0.78 175 $1807313-$ MutT/nudix family protein (DR2356) 0.59 171 $12270912+$ MutT/nudix family protein (DR2356) 0.59 701 $3126894+$ Ribonucleoside-diphosphate reductase; alpha (nrdE) 0.58 354 $3128900+$ Ribonucleoside-diphosphate reductase; beta (nrdF) 0.62 219 $279822-$ LexA repressor (lexA) 0.52 940 $3162296+$ ATP-dependent helicase HepA (hepA) 0.52 940 $3162296+$ ATP-dependent helicase HepA (hepA) 0.61 $1,066$ $378759+$ Extracellular nuclease; putative (DR80067) 1.02 143 $100371+$ Single-stranded DNA binding protein (ssb) 0.65 403 $12335883+$ CinA protein (DR2338) 0.83 343 $3124398+$ Integrase-recombinase XerD; putative (xerD) </td <td>0.50</td> <td>783</td> <td>1 1936853+</td> <td>DNA helicase RecG ($recG$)</td>	0.50	783	1 1936853+	DNA helicase RecG ($recG$)
0.616991 2084844 -DNA ligase $(dnll)$ 0.761701 80637 -MutT/nudix family protein (DR0079)0.761771 276905 +MutT/nudix family protein (DR0274)0.623221 324367 -MutT/nudix family protein (DR0329)0.602491 555381 +MutT/nudix family protein (DR0550)0.731661 991985 -MutT/nudix family protein (DR1007)0.831581 1019945 +MutT/nudix family protein (DR1025)0.781751 1807313 -MutT/nudix family protein (DR1025)0.781751 1807313 -MutT/nudix family protein (DR2356)0.597013 126894 +Ribonucleoside-diphosphate reductase; alpha (nrdE)0.622192 379822 -Lex A repressor (lex 4)0.875021 1112934 +DNA repair protein (nrdA)0.529403 162296 +ATP-dependent helicase HepA (hepA)0.611,0663 78759 +Extracellular nuclease; putative (DRB0067)1.021431 100371 +Single-stranded DNA binding protein (ssb)0.654031 2335883 +CinA protein (DR2338)0.833433 124398 +Integrase-recombinase XerD; putative (xerD)	0.57	955	1 1732431-	DNA-directed DNA polymerase (<i>polA</i>)
0.76 170 $180637-$ MutT/nudix family protein (DR0079) 0.76 177 $1276905+$ MutT/nudix family protein (DR0274) 0.62 322 $1324367-$ MutT/nudix family protein (DR0329) 0.60 249 $1555381+$ MutT/nudix family protein (DR0975) 0.73 166 $199185-$ MutT/nudix family protein (DR0975) 0.94 165 $11019945+$ MutT/nudix family protein (DR1007) 0.83 158 $11038101+$ MutT/nudix family protein (DR176) 0.78 175 $11807313-$ MutT/nudix family protein (DR272) 1.29^b 143 $12354268+$ MutT/nudix family protein (DR2272) 0.59 701 $3126894+$ Ribonucleoside-diphosphate reductase; alpha (nrdE) 0.58 354 $312890+$ Ribonucleoside-diphosphate reductase; beta (nrdF) 0.62 219 $2379822-$ LexA repressor (lexA) 0.87 502 $11112934+$ DNA repair protein (nrdA) 0.52 940 $3162296+$ ATP-dependent helicase HepA (hepA) 0.61 $1,066$ $378759+$ Extracellular nuclease; putative (DRB0067) 1.02 143 $100371+$ Single-stranded DNA binding protein (nsb) 0.65 403 $1233883+$ CinA protein (DR238) 0.83 343 $3124398+$ Integrase-recombinase XerD; putative (xerD)	0.61	699	1 2084844-	DNA ligase (dnlJ)
0.76 177 $1\ 276905 +$ MutT/nudix family protein (DR0274) 0.62 322 $1\ 324367 -$ MutT/nudix family protein (DR0329) 0.60 249 $1\ 555381 +$ MutT/nudix family protein (DR0550) 0.73 166 $1\ 991985 -$ MutT/nudix family protein (DR1007) 0.83 158 $1\ 019945 +$ MutT/nudix family protein (DR1007) 0.83 158 $1\ 038101 +$ MutT/nudix family protein (DR1025) 0.78 175 $1\ 1807313 -$ MutT/nudix family protein (DR2272) 0.78 171 $2\ 270912 +$ MutT/nudix family protein (DR2356) 0.59 701 $3\ 126894 +$ Ribonucleoside-diphosphate reductase; alpha (nrdE) 0.58 354 $3\ 128990 +$ Ribonucleoside-diphosphate reductase; alpha (nrdF) 0.62 219 $2\ 379822 -$ LexA repressor (lexA) 0.87 502 $1\ 1112934 +$ DNA repair protein (nrdA) 0.52 940 $3\ 122396 +$ ATP-dependent helicase HepA (hepA) 0.65 403 $1\ 233583 +$ CinA protein (DR2338) 0.83 343 $3\ 124398 +$ Integrase-recombinase XerD; putative (xerD)	0.76	170	1 80637-	MutT/nudix family protein (DR0079)
0.62 322 $1\ 324367-$ MutT/nudix family protein (DR0329) 0.60 249 $1\ 555381+$ MutT/nudix family protein (DR0550) 0.73 166 $1\ 991985-$ MutT/nudix family protein (DR075) 0.94 165 $1\ 101995+$ MutT/nudix family protein (DR1007) 0.83 158 $1\ 1038101+$ MutT/nudix family protein (DR1025) 0.78 175 $1\ 1807313-$ MutT/nudix family protein (DR2272) 0.59 171 $1\ 2270912+$ MutT/nudix family protein (DR2272) 1.29^b 143 $1\ 2354268+$ MutT/nudix family protein (DR2356) 0.59 701 $3\ 126894+$ Ribonucleoside-diphosphate reductase; alpha (nrdE) 0.58 354 $3\ 128990+$ Ribonucleoside-diphosphate reductase; beta (nrdF) 0.62 219 $2\ 379822-$ LexA repressor (lexA) 0.87 502 $1\ 1112934+$ DNA repair protein (radA) 0.52 940 $3\ 162296+$ ATP-dependent helicase HepA (hepA) 0.65 403 $1\ 2335883+$ CinA protein (DR2338) 0.83 343 $3\ 124398+$ Integrase-recombinase XerD; putative (xerD)	0.76	177	1.276905 +	MutT/nudix family protein (DR0274)
0.60 249 $1555381+$ MutT/nudix family protein (DR0550) 0.73 166 $1991985-$ MutT/nudix family protein (DR0975) 0.94 165 $11019945+$ MutT/nudix family protein (DR1007) 0.83 158 $11038101+$ MutT/nudix family protein (DR1025) 0.78 175 $11807313-$ MutT/nudix family protein (DR1776) 0.59 171 $12270912+$ MutT/nudix family protein (DR2356) 0.59 701 $3126894+$ Ribonucleoside-diphosphate reductase; alpha (nrdE) 0.58 354 $3128990+$ Ribonucleoside-diphosphate reductase; beta (nrdF) 0.89 140 $3126505+$ Ribonucleotide reductase; NrdI family (DRB0107) 0.62 219 $2379822-$ LexA repressor (lexA) 0.87 502 $11112934+$ DNA repair protein (nrdA) 0.52 940 $3162206+$ ATP-dependent helicase HepA (hepA) 0.51 $1,066$ $378759+$ Extracellular nuclease; putative (DRB0067) 1.02 143 $1100371+$ Single-stranded DNA binding protein (ssb) 0.65 403 $12335883+$ CinA protein (DR2338) 0.83 343 $3124398+$ Integrase-recombinase XerD; putative (xerD)	0.62	322	1 324367-	MutT/nudix family protein (DR0329)
0.73 166 $1991985-$ MutT/nudix family protein (DR0975) 0.94 165 $1019945+$ MutT/nudix family protein (DR1007) 0.83 158 $1038101+$ MutT/nudix family protein (DR1025) 0.78 175 $11807313-$ MutT/nudix family protein (DR1776) 0.59 171 $12270912+$ MutT/nudix family protein (DR2272) 1.29^b 143 $12354268+$ MutT/nudix family protein (DR2356) 0.59 701 $3126894+$ Ribonucleoside-diphosphate reductase; alpha (nrdE) 0.58 354 $3128990+$ Ribonucleoside-diphosphate reductase; beta (nrdF) 0.89 140 $3126505+$ Ribonucleoside diphosphate reductase; beta (nrdF) 0.62 219 $2379822-$ LexA repressor (lexA) 0.52 940 $3162296+$ ATP-dependent helicase HepA (hepA) 0.61 $1,066$ $378759+$ Extracellular nuclease; putative (DRB0067) 1.02 143 $100371+$ Single-stranded DNA binding protein (ssb) 0.65 403 $1233583+$ CinA protein (DR2338) 0.83 343 $3124398+$ Integrase-recombinase XerD; putative (xerD)	0.60	249	1555381 +	MutT/nudix family protein (DR0550)
0.94 165 $1019945 +$ MutT/nudix family protein (DR1007) 0.83 158 $1038101 +$ MutT/nudix family protein (DR1025) 0.78 175 $11807313 -$ MutT/nudix family protein (DR1276) 0.59 171 $12270912 +$ MutT/nudix family protein (DR2272) 1.29^b 143 $12354268 +$ MutT/nudix family protein (DR2356) 0.59 701 $3126894 +$ Ribonucleoside-diphosphate reductase; alpha (nrdE) 0.58 354 $3128990 +$ Ribonucleoside-diphosphate reductase; beta (nrdF) 0.89 140 $3126505 +$ Ribonucleotide reductase; NrdI family (DRB0107) 0.62 219 $2379822 -$ LexA repressor (lexA) 0.52 940 $3162296 +$ ATP-dependent helicase HepA (hepA) 0.61 $1,066$ $378759 +$ Extracellular nuclease; putative (DRB0067) 1.02 143 $1100371 +$ Single-stranded DNA binding protein (ssb) 0.65 403 $12335883 +$ CinA protein (DR2338) 0.83 343 $3124398 +$ Integrase-recombinase XerD; putative (xerD)	0.73	166	1 991985-	MutT/nudix family protein (DR0975)
0.83 158 $1038101 +$ MutT/nudix family protein (DR1025) 0.78 175 $11807313 -$ MutT/nudix family protein (DR1776) 0.59 171 $12270912 +$ MutT/nudix family protein (DR2272) 1.29^b 143 $12354268 +$ MutT/nudix family protein (DR2356) 0.59 701 $3126894 +$ Ribonucleoside-diphosphate reductase; alpha (nrdE) 0.58 354 $3128990 +$ Ribonucleoside-diphosphate reductase; beta (nrdF) 0.89 140 $3126505 +$ Ribonucleotide reductase; NrdI family (DRB0107) 0.62 219 $2379822 -$ LexA repressor (lexA) 0.52 940 $3162296 +$ ATP-dependent helicase HepA (hepA) 0.61 $1,066$ $378759 +$ Extracellular nuclease; putative (DRB0067) 1.02 143 $1100371 +$ Single-stranded DNA binding protein (ssb) 0.65 403 $12335883 +$ CinA protein (DR2338) 0.83 343 $3124398 +$ Integrase-recombinase XerD; putative (xerD)	0.94	165	$1\ 1019945 +$	MutT/nudix family protein (DR1007)
0.78 175 $11807313 -$ MutT/nudix family protein (DR1776) 0.59 171 $12270912 +$ MutT/nudix family protein (DR2272) 1.29^b 143 $12354268 +$ MutT/nudix family protein (DR2356) 0.59 701 $3126894 +$ Ribonucleoside-diphosphate reductase; alpha (nrdE) 0.58 354 $3128990 +$ Ribonucleoside-diphosphate reductase; beta (nrdF) 0.89 140 $3126505 +$ Ribonucleoside reductase; NrdI family (DRB0107) 0.62 219 $2379822 -$ LexA repressor (lexA) 0.52 940 $3162296 +$ ATP-dependent helicase HepA (hepA) 0.61 $1,066$ $378759 +$ Extracellular nuclease; putative (DRB0067) 1.02 143 $1100371 +$ Single-stranded DNA binding protein (ssb) 0.65 403 $12335883 +$ CinA protein (DR2338) 0.83 343 $3124398 +$ Integrase-recombinase XerD; putative (xerD)	0.83	158	$1\ 1038101 +$	MutT/nudix family protein (DR1025)
0.59 171 $12270912+$ MutT/nudix family protein (DR2272) 1.29^b 143 $12354268+$ MutT/nudix family protein (DR2356) 0.59 701 $3126894+$ Ribonucleoside-diphosphate reductase; alpha (nrdE) 0.58 354 $3128990+$ Ribonucleoside-diphosphate reductase; beta (nrdF) 0.89 140 $3126505+$ Ribonucleotide reductase; NrdI family (DRB0107) 0.62 219 $2379822-$ LexA repressor (lexA) 0.87 502 $1112934+$ DNA repair protein (radA) 0.61 $1,066$ $378759+$ Extracellular nuclease; putative (DRB0067) 1.02 143 $100371+$ Single-stranded DNA binding protein (ssb) 0.65 403 $12335883+$ CinA protein (DR2338) 0.83 343 $3124398+$ Integrase-recombinase XerD; putative (xerD)	0.78	175	1 1807313-	MutT/nudix family protein (DR1776)
1.29^b 143 $12354268+$ MutT/nudix family protein (DR2356) 0.59 701 $3126894+$ Ribonucleoside-diphosphate reductase; alpha (nrdE) 0.58 354 $3128990+$ Ribonucleoside-diphosphate reductase; beta (nrdF) 0.89 140 $3126505+$ Ribonucleoside diphosphate reductase; beta (nrdF) 0.62 219 $2379822-$ LexA repressor (lexA) 0.87 502 $1112934+$ DNA repair protein (radA) 0.52 940 $3162296+$ ATP-dependent helicase HepA (hepA) 0.61 $1,066$ $378759+$ Extracellular nuclease; putative (DRB0067) 1.02 143 $100371+$ Single-stranded DNA binding protein (ssb) 0.65 403 $1233583+$ CinA protein (DR2338) 0.83 343 $3124398+$ Integrase-recombinase XerD; putative (xerD)	0.59	171	1 2270912+	MutT/nudix family protein (DR2272)
0.59 701 $3126894+$ Ribonucleoside-diphosphate reductase; alpha (nrdE) 0.58 354 $3128990+$ Ribonucleoside-diphosphate reductase; beta (nrdF) 0.89 140 $3126505+$ Ribonucleoside reductase; NrdI family (DRB0107) 0.62 219 $2379822-$ LexA repressor (lexA) 0.87 502 $1112934+$ DNA repair protein (radA) 0.52 940 $3162296+$ ATP-dependent helicase HepA (hepA) 0.61 $1,066$ $378759+$ Extracellular nuclease; putative (DRB0067) 1.02 143 $1100371+$ Single-stranded DNA binding protein (ssb) 0.65 403 $12335883+$ CinA protein (DR2338) 0.83 343 $3124398+$ Integrase-recombinase XerD; putative (xerD)	1.29 ^b	143	1 2354268+	MutT/nudix family protein (DR2356)
0.58 354 $3128990+$ Ribonucleoside-diphosphate reductase; beta (nrdF) 0.89 140 $3126505+$ Ribonucleotide reductase; NrdI family (DRB0107) 0.62 219 $2379822-$ LexA repressor (lexA) 0.87 502 $1112934+$ DNA repair protein (radA) 0.52 940 $3162296+$ ATP-dependent helicase HepA (hepA) 0.61 $1,066$ $37879+$ Extracellular nuclease; putative (DRB0067) 1.02 143 $1100371+$ Single-stranded DNA binding protein (ssb) 0.65 403 $12335883+$ CinA protein (DR2338) 0.83 343 $3124398+$ Integrase-recombinase XerD; putative (xerD)	0.59	701	3 126894+	Ribonucleoside-diphosphate reductase; alpha (<i>nrdE</i>)
0.89 140 $3 \ 126505 +$ Ribonucleotide reductase; NrdI family (DRB0107) 0.62 219 $2 \ 379822 -$ LexA repressor ($lexA$) 0.87 502 $1 \ 112934 +$ DNA repair protein ($radA$) 0.52 940 $3 \ 162296 +$ ATP-dependent helicase HepA ($hepA$) 0.61 $1,066$ $3 \ 78759 +$ Extracellular nuclease; putative (DRB0067) 1.02 143 $1 \ 100371 +$ Single-stranded DNA binding protein (ssb) 0.65 403 $1 \ 2335883 +$ CinA protein (DR2338) 0.83 343 $3 \ 124398 +$ Integrase-recombinase XerD; putative ($xerD$)	0.58	354	3 128990+	Ribonucleoside-diphosphate reductase; beta (<i>nrdF</i>)
0.62 219 $2379822-$ LexA repressor ($lexA$) 0.87 502 $1112934+$ DNA repair protein ($radA$) 0.52 940 $3162296+$ ATP-dependent helicase HepA ($hepA$) 0.61 $1,066$ $378759+$ Extracellular nuclease; putative (DRB0067) 1.02 143 $1100371+$ Single-stranded DNA binding protein (ssb) 0.65 403 $12335883+$ CinA protein (DR2338) 0.83 343 $3124398+$ Integrase-recombinase XerD; putative ($xerD$)	0.89	140	3 126505+	Ribonucleotide reductase; NrdI family (DRB0107)
0.87 502 $1112934+$ DNA repair protein (radA) 0.52 940 $3162296+$ ATP-dependent helicase HepA (hepA) 0.61 $1,066$ $378759+$ Extracellular nuclease; putative (DRB0067) 1.02 143 $1100371+$ Single-stranded DNA binding protein (ssb) 0.65 403 $12335883+$ CinA protein (DR2338) 0.83 343 $3124398+$ Integrase-recombinase XerD; putative (xerD)	0.62	219	2 379822-	LexA repressor (<i>lexA</i>)
0.52 940 3 162296+ ATP-dependent helicase HepA (hepA) 0.61 1,066 3 78759+ Extracellular nuclease; putative (DRB0067) 1.02 143 1 100371+ Single-stranded DNA binding protein (ssb) 0.65 403 1 2335883+ CinA protein (DR2338) 0.83 343 3 124398+ Integrase-recombinase XerD; putative (xerD)	0.87	502	1 1112934+	DNA repair protein (radA)
0.61 1,066 3 78759+ Extracellular nuclease; putative (DRB0067) 1.02 143 1 100371+ Single-stranded DNA binding protein (ssb) 0.65 403 1 2335883+ CinA protein (DR2338) 0.83 343 3 124398+ Integrase-recombinase XerD; putative (xerD)	0.52	940	3 162296+	ATP-dependent helicase HepA (hepA)
1.02 143 1 100371+ Single-stranded DNA binding protein (ssb) 0.65 403 1 2335883+ CinA protein (DR2338) 0.83 343 3 124398+ Integrase-recombinase XerD; putative (xerD)	0.61	1,066	3 78759+	Extracellular nuclease; putative (DRB0067)
0.65 403 1 2335883+ CinA protein (DR2338) 0.83 343 3 124398+ Integrase-recombinase XerD; putative (xerD)	1.02	143	$1\ 100371 +$	Single-stranded DNA binding protein (ssb)
0.833433 124398+Integrase-recombinase XerD; putative (xerD)	0.65	403	1 2335883+	CinA protein (DR2338)
	0.83	343	3 124398+	Integrase-recombinase XerD; putative (xerD)

^{*a*} Position in the genome is indicated by a chromosome number (1, large chromosome; 2, small chromosome; 3, megaplasmid MP1) followed by position of the translation initiation site and orientation of the gene (+, direct strand; -, complementary strand).

^b Predicted highly expressed.

gene classes. However, *D. radiodurans* possesses a collection of PHX S-layer (surface structure) proteins which may provide environmental protection (e.g., against desiccation). *D. radiodurans* also has an abundance of PHX ATP-binding cassette transporters for various peptides, branched-chain amino acids (LivK), phosphates, and maltose (periplasmic maltose-binding protein). Intriguingly, the top PHX gene is the multifunctional tricarboxylic acid cycle gene aconitate hydratase (aconitase).

(iv) *M. tuberculosis*. The PHX isocitrate lyase enzyme (AceA) is very abundant when *M. tuberculosis* inhabits macrophages (5). Concomitant to infection of activated macrophages, *M. tuberculosis* exhibits an increased expression of iso-

citrate lyase, preferred over isocitrate dehydrogenase, the first enzyme of the glyoxylate shunt pathway that yields a net carbon gain in metabolism of fatty acids (5). This is consistent with the elevated expression levels of this gene. When *M. tuberculosis* enters macrophages, induction of stress proteins also results. Many fatty acid biosynthesis genes (e.g., *fas; fadA; fadB; fadE4, -E5*, and *-E7; fadD3* and others) are PHX. Also, mycolic acid synthases 2 and 3 (both PHX) are abundant on the bacterial outer cell wall. Apart from *M. tuberculosis*, of all the genomes in Table 1, the AceA gene appears only in *D. radiodurans* (PHX) and *E. coli* (not PHX). Thirty-six genes labeled PE-PGRS (9) and distinguished by a preponderance of glycine-glycine doublets and anomalous repetitive structures are

TABLE 5. Selected classes of PHX genes

	No. of genes					
Genome	Flagellar	Histidine kinase ^a	Electron transfer ^b	Detoxifi- cation ^c	Aminoacyl tRNA synthetase ^d	
E. coli	1	0	20	8	20	
H. influenzae	0	0	5	3	7	
B. subtilis	4	0	5	4	4	
V. cholerae	1	0	8	1	8	
D. radiodurans	0	0	9	12	1	
R. prowazekii	0	0	1	0	0	
C. trachomatis	0	0	0	0	0	
C. pneumoniae	0	0	0	0	1	
M. genitalium	0	0	0	0	1	
M. pneumoniae	0	0	0	0	1	
B. burgdorferi	5	0	0	0	1	
T. pallidum	7	1	0	1	1	
H. pylori	2	1	5	2	0	
M. tuberculosis	0	5	8	3	6	
Synechocystis	0	7	16	4	1	
A. aeolicus	4	0	37	5	9	
T. maritima	3	0	11	2	5	
M. jannaschii	1	0	5	0	0	
M. thermoauto- trophicum	0	2	6	1	2	
A. fulgidus	1	0	24	4	7	
P. horikoshii	3	0	10	1	6	
P. abyssi	3	0	20	2	10	

^{*a*} Also includes sensors and other two-component system proteins.

^b Includes cytochrome- and NADH-related proteins, ferredoxins, and flavodoxins.

^c Includes Sod, glutathione peroxidase, thiol-specific antioxidant-peroxidase, thiosulfate sulfurtransferase, cytochrome *c* peroxidase, glutathione-*S*-transferase, alkyl hydroperoxide reductase, catalase, LexA repressor, Dps, and rubrerythrin (peroxide oxidase).

^d Also includes amidotransferase.

PHX. These may obstruct the host immune system (cf. references 31 and 46).

The bacterial cell has developed complex mechanisms to deal with membrane translocation, secretion of polypeptides, and subsequent folding. SecA, essential and unique to eubacteria (i.e., not found in archaea), is fundamental for protein translocation to the periplasm. Secretion-specific chaperones include SecB and the signal recognition particle. In these activities, the major chaperones GroEL, DnaK, and the trigger factor are also involved (13). In addition to structural subunits, such as SecY, SecE, and SecG, the translocase has a mechanical motor device, the SecA ATPase, that binds to SecYEG to establish the functional translocase core. M. tuberculosis possesses two SecA paralogs with distinct substrate specificities. The SecA gene is also PHX in V. cholerae, E. coli, Synechocystis, Mycoplasma pneumoniae, T. pallidum, B. burgdorferi, and A. aeolicus. The secretion pathway is used by many protein substrates. The cellular destination of all secretory polypeptides is governed by a 20- to 30-residue aminoterminal sequence, the leader peptide, which also helps guide SecA binding to the substrate. SecA, SecB, and SecG are all involved in protein export and chaperonin activity. Gramnegative bacteria also secrete a variety of proteins into the extracellular milieu mediated by secretion apparatus types I to IV (13). These proteins can influence bacterium-host interactions.

(v) Synechocystis (Cyanobacterium). In the Synechocystis genome, the chaperonin GroEL-2 attains the highest expression level (E = 1.51) and the duplicate GroEL-1 (E = 1.47) and DnaK (E = 1.40) also have high expression levels. Apart from GroEL and DnaK, several CH proteins are among the most

highly expressed, including ClpC (E = 1.46) and three copies of FtsH (1.49, 1.30, and 1.17). A fourth FtsH has an expression level of 0.82. In many genomes, duplicated genes have only a single copy that is PHX (S. Karlin, A. D. Kaiser, A. M. Campbell, and J. Mrázek, unpublished data).

The majority of primary photosynthesis genes attain very high expression levels, e.g., the phycobilisome LCM core-membrane linker polypeptide (ApcE) gene records the highest predicted expression level ($\vec{E} = 1.51$) in Synechocystis. The PHX genes include more than 30 genes contributing to photosynthesis. The large and small subunits of rubisco are both highly expressed. Interestingly, rubisco is also PHX in the archaeal genome of Archaeoglobus fulgidus. There are several glycolysis and gluconeogenesis genes in Synechocystis that satisfy our criteria as PHX, including those for phosphoglycerate kinase, fructose-1,6-bisphosphate aldolase, GAPDH, enolase, fructose-1,6-bisphosphatase, phosphofructokinase, and pyruvate kinase. These genes also act in photosynthesis. The PHX "giant" ribosomal protein S1, weakly homologous to S1 of E. coli, is only 327 aa, much reduced from the usual size exceeding 500 aa. There are many PHX genes for aerobic respiration and many contributing to electron transport. This is consistent with the proposition that respiration and photosynthesis are linked in Synechocystis.

(vi) *H. pylori. H. pylori* lives in the thick mucus lining that protects the stomach from its own digestive juices. Among the most PHX genes are those for urease α (UreA), urease β (UreB), and the accessory *ureI* (all occurring as a cluster, or operon), which convert urea from gastric juices into bicarbonate and ammonia (NH₃), which help to neutralize the highly acidic stomach environment and allow *H. pylori* to safely traverse the mucus layer to the epithelium surface. Ammonia could also serve as a nitrogen source for amino acids (19, 45). Other accessory proteins, UreE, UreF, UreG, and UreH, that are not part of the urease enzyme and are not PHX help to incorporate Ni²⁺ ions required for urease enzyme assembly and activity. UreI pumps urea from the outside to the inside of the cell.

The *H. pylori* genome is rife with genes encoding a family of outer membrane proteins encompassing more than 32 members (many duplicated). The genes encoding outer membrane proteins 2, 9, 11, 14, 21, and 28 of this family are PHX. Some porins may be involved importantly in the antibiotic susceptibility of *H. pylori*. The PHX chaperone genes of *H. pylori* include those for GroEL and the two DnaK cochaperones GrpE and DnaJ (the expression level of DnaK is 1.00, at the boundary of PHX genes); the general stress gene *ctc* and the thioredoxin (*trxA*) and thioredoxin reductase (*trxB*) genes are in a cluster. The two chemotaxis genes *cheW* and *cheA* as an operon are PHX, as are the two flagellar genes *motB* and *flaE*. The *cheA* gene controls flagellar motor on-off changes. Urease plays a key role in chemotactic motility (48).

The mycoplasma Ureaplasma urealyticum genome has just been sequenced. It is distinguished by an operon of three urease complex components, ureG, ureE, and ureC, among the most PHX genes in this genome (data not shown). Several fatty acid biosynthesis genes (those for FadA, biotin carboxylase, and cyclopropane fatty acid synthase), a spectrum of cytochrome genes, and the genes for two antioxidants (catalase and alkylhydroperoxide reductase) carry high expression values.

(vii) *C. trachomatis* and *C. pneumoniae* (mammalian obligate intracellular parasites). Chlamydia live in vacuoles which burst to spread. Their PHX genes include one of two ATP-ADP exchange translocase genes. This antiporter takes ATP from host cytoplasmic sources and releases ADP from the bacterial cell;

the standard mitochondrial exchange is reversed. The ATP-ADP translocase is very uncommon among bacteria and has been found only in *Chlamydia* and *Rickettsia* bacteria and in a spectrum of plant plastids. The two *C. trachomatis* ATP-ADP translocases function differently (71). One exchanges ATP and ADP as described above, and the other is a nucleotide transporter (62). NusA and NusG contribute to termination and antitermination as components of the transcription process (41). The highest expression level of NusA (E = 1.28) is applicable to *C. pneumoniae*. The Nus proteins are PHX in *E. coli*, *V. cholerae*, *D. radiodurans*, *Synechocystis*, *M. pneumoniae*, *T. pallidum*, *B. burgdorferi*, *C. pneumoniae*, *M. jannaschii*, and *P. abyssi*.

The PHX OmpA receptor of *C. pneumoniae* makes up more than 60% of all its membrane proteins (R. S. Stephens, personal communication). There are three GroEL chaperonins each in *C. trachomatis* and *C. pneumoniae*, of which only GroEL-1 is PHX. The chaperonin trigger factor and another PPIase are significantly PHX in *C. pneumoniae*.

(viii) *R. prowazekii*. Unlike *C. trachomatis*, the human obligate intracellular parasite *R. prowazekii* is not able to metabolize glucose (70). A distinctive PHX gene encodes the cell division protein FtsZ. In contrast to *C. trachomatis*, *R. prowazekii* contains five ATP-ADP exchange translocase genes, but individually none is PHX. Perhaps the redundancy of five suffices. There are no PHX glycolysis genes in *R. prowazekii* and *C. trachomatis*. *R. prowazekii* does engage in some respiration, but these microbes apparently extract substantial energy straight from the host.

(ix) T. pallidum. Spirochetes in general, but not T. pallidum in particular, are mostly free living and are found in soil and freshwater, but they are also commensal with clams and other animals. T. pallidum is restricted to human hosts. Its genome stands out, with the greatest number of PHX flagellar genes among the genomes shown in Table 5 (see also reference 8). These include the genes for the flagellar filament outer layer proteins FlaA-1 and FlaA-2, flagellar motor switch protein, flagellar filament 33-kDa core protein, and flagellar basal-body rod protein FlgG-2. Flagellar proteins in spirochetes operate in the periplasm. This is different from most other genomes, where flagella are extracellular at the surface. Why is T. pallidum so mobile? The bacterium (which causes syphilis) invades all parts of the human body, including the brain. The abundance of highly expressed flagellar genes in T. pallidum could facilitate its movement and enhance its survival by spreading. The genes encoding the response regulator CheY and the purine binding CheW are also PHX genes. The recombinationrepair proteins RecA, RecX, and Ssb (single-stranded binding) are PHX. T. pallidum features the longest S1 RP (862 aa) among all complete genomes (Table 6).

(x) A. aeolicus. On the basis of the 16S RNA sequence, the A. aeolicus genome is classified as that of a deeply branching gram-negative hyperthermophile. However, with respect to PHX genes, there is much resemblance to E. coli. A. aeolicus, like a classical gram-negative bacterium, contains an S1 RP of 534 aa which is very highly expressed in this genome. Many electron transport proteins stand out as PHX, including cytochrome c oxidase, cytochrome b, cytochrome c_{552} , several NADH dehydrogenase subunits, and most subunits (α , β , and γ) of the iron sulfur ferredoxin oxidoreductase. The detoxification genes sodA, the alkyl hydroperoxide reductase gene, ahpC1, and ahpC2 and four flagellar genes are PHX. Biotin carboxylase is a PHX protein important in fatty acid biosynthesis. Biotin acts as a coenzyme covalently linked to carboxylase (67). Its highest expression level occurs in A. aeolicus.

TABLE 6. RPs

Genome	No. of RP genes ^a	S1 length (aa)	S2 isolated ^j	L7-L12 ^g	\mathbf{P}_0^f	
E. coli	55	556	+	+	_	
H. influenzae	55	548	+	+	_	
B. subtilis	57	381^{b}	+	+	_	
V. cholerae	49 ^c	555	+	+	_	
D. radiodurans	52	628	+	+	_	
R. prowazekii	54	567	+	+	_	
C. trachomatis	52	568	+	+	_	
C. pneumoniae	52	579	+	+	_	
M. genitalium	50	Missing	+	+	_	
M. pneumoniae	50	Missing	+	+	_	
B. burgdorferi	53	552	+	+	_	
T. pallidum	52	862	+	+	_	
H. pylori	52	555	+	+	_	
M. tuberculosis	55	480	+	+	_	
Synechocystis	53	327, 304 ^h	+	+	_	
A. aeolicus	53	534	+	+	_	
T. maritima	55	542^{i}	+	+	_	
M. jannaschii	60	Missing	_	d	+	
<i>M. thermoautotrophicum</i>	61	Missing	_e	d	+	
A. fulgidus	61	Missing	_ ^e	d	+	
P. horikoshii	54	Missing	_e	d	+	
P. abyssi	61	Missing	_e	d	_	

^a Including duplicated genes.

^b S1 homolog.

^c More RP genes may be present in *V. cholerae* that failed to be identified by similarity searches.

^d The archaeal genomes have distinct RP L7 and L12, but none is similar to *E. coli* L7-L12.

^e M. thermoautotrophicum RP Sa (homolog of E. coli S2) gene is 2 kb downstream from S16 (E. coli S9) separated by genes for DNA-dependent RNA polymerase subunit N, DNA-dependent RNA polymerase subunit K, and enolase. Exactly the same sequence of genes occurs in A. fulgidus, P. horikoshii, and P. Abyssi (with the only exception that in P. horikoshii, two ORFs replace the subunit K and enolase genes).

 ${}^{f}P_{0}$ is functionally equivalent to *E. coli* L10 (SwissProt) and is sometimes called L10E. P₀ has a hyperacidic charge run at the C terminus. Eukaryotes also have acidic RPs P₁ and P₂ (not present in the archaeal genomes at hand). +, present; -, absent.

^g Unlike most bacterial RPs, which are basic (mostly >20% basic residues), L7-L12 is acidic. For example, *E. coli* L7-L12 contains 17% acidic residues. +, present; -, absent.

^h Synechocystis has two copies of S1, and both are significantly smaller than the typical S1 from most other eubacterial genomes.

^{*i*} T. maritima S1 contains a frameshift.

^{*j*} +, isolated; -, proximal to other RP genes.

Biotin carboxylase is also PHX in *E. coli*, *H. influenzae*, *V. cholerae*, *H. pylori*, *Synechocystis*, *C. trachomatis*, and *A. fulgidus*.

(xi) *M. jannaschii. M. jannaschii* (strictly anaerobic) carries out no fermentation. Energy conservation proceeds exclusively by the conversion of H_2 plus CO₂ to CH₄. Special PHX genes are those for thermosome (*ths*) (this applies to all archaea) and flagellin (*flaB1*). As expected, the PHX genes include more than 20 participating in methanogenesis. Actually, the three genes with the highest expression levels participate in methanogenesis. The thermosome "homolog" of GroEL is very highly expressed (E = 1.56). The absence of the giant S1 RP applies to all archaeal genomes. The top PHX genes correlate with the greatest protein abundances verified by 2-D-gel analysis (C. Giometti, Argonne National Laboratory, personal communication).

(xii) *M. thermoautotrophicum*. The predominant PHX genes in *M. thermoautotrophicum* are the thermosome subunits *thsA* (E = 1.33) and *thsB* (1.38). Again, most PHX genes of *M. thermoautotrophicum* are involved in methanogenesis. DnaK, missing from *M. jannaschii*, is PHX in *M. thermoautotrophicum*.

(xiii) A. fulgidus. In A. fulgidus, both thermosome units α and β are PHX. The elaborate proteasome complex is PHX in

A. fulgidus and P. abyssi. Intriguingly, A. fulgidus contains two PHX copies of rubisco (rbcL-1 and rbcL-2). A. fulgidus has more than 300 PHX genes, compared to about 150 in M. jannaschii and M. thermoautotrophicum. Many NADH dehydrogenases and general anaerobic respiration proteins (electron acceptors) are based on nitrate and sulfate. A. fulgidus grows using sulfate or thiosulfate as an electron acceptor and H₂ as an electron donor. Although A. fulgidus is not a methanogen, there are several methanogenesis homologs among the PHX genes. Cells are regular to irregular spheres and have flagella at one end for motility; flaB1 is PHX. A. fulgidus seems to have much more metabolic flexibility with organic and inorganic sources than the methanogens. The polyamine spermidineputrescine transporter in the periplasm is PHX in H. influenzae, V. cholerae, M. pneumoniae, T. maritima, and A. fulgidus. These apparently help to maintain charge homeostasis. The polyamines are small organic molecules generally present in all living organisms. They are synthesized by a highly regulated pathway from arginine or ornithine and can also be transported in and out of cells. Polyamines influence the transcriptional and translational stages of protein synthesis, stabilize membranes, and, in mammalian systems, modulate neurophysiological functions and may act as intracellular messengers. The five archaeal genomes are rife with PHX genes which conduct electron transfer as needed with anaerobic respiration.

Selected classes of highly expressed genes. Three gene groups are prominently PHX: RP, CH, and TF. This finding is consistent with protein abundance assessments deduced from 2-D-gel assays for *E. coli* (65) (see below). These results support the choice of the RP, CH, and TF gene classes as representative standards for PHX genes in prokaryotes. Five specialized classes of genes were examined in Table 5 for PHX genes.

(i) Flagellar genes. Assembly of a flagellum, the motive organelle produced by many bacteria, requires export of protein subunits from the cytoplasm to the outer surface of the cell by a mechanism resembling type III secretion (74). Flagella generally consist of three main components: the basal body, the hook, and the filament. Flagellum biogenesis and chemotaxis occur in coordination with flagellum assembly and in response to environmental signals. In this context, class I flagellar genes, consisting of *flhD* and *flhC*, are first produced. Class II genes encode structural and accessory proteins needed for assembly of the basal body and hook components. Class III proteins are required for maturation of the flagellum and the chemosensory system. This and recent evidence indicate that the flagellum regulon can influence bacterium-host interactions independent of motility (74). There is also an established selective connection of flagellar motion and chemotaxis responses. The flagellum secretion apparatus may be viewed as part of the chaperone family essential for bacterial viability. Flagella are generally absent in nonmotile prokaryotes.

Why do flagellar PHX genes proliferate among the spirochete genomes of *B. burgdorferi* and *T. pallidum*? It is known that the flagella of spirochetes are enclosed in a compartment inside the periplasm, whereas in most other bacteria they are attached to a cell surface receptor outside the cell. Moreover, the flagellar genes of the spirochetes respond to a specialized sigma factor, σ^{28} , whereas the flagellar genes of *E. coli* are commonly activated by the standard σ^{70} . Several flagellar PHX genes of *H. pylori* have mixed controls, e.g., σ^{28} and σ^{54} . The flagellar export apparatus in *E. coli* also functions as a protein secretion system (74).

(ii) Detoxification genes. PHX genes acting in detoxification are preponderant in *D. radiodurans* and significant in *E. coli*,

TABLE 7. Expression levels of two-component system genes of *E. coli* and *B. subtilis*

E(g)	Length (kb)	Position ^a	Gene and comments
E. coli			
0.39	430	417113+	Phosphate regulon sensor protein (<i>phoR</i>)
0.37	893	723637-	Sensor protein (kdpD)
0.40	485	1188999-	Sensor protein $(phoQ)$
0.40	597	1276841-	Nitrate-nitrite sensor protein (<i>narX</i>)
0.39	653	1973348-	Chemotaxis protein (<i>cheA</i>)
0.36	565	2583751 +	Nitrate-nitrite sensor protein (<i>narQ</i>)
0.47	449	3533506-	Osmolarity sensor protein $(envZ)$
0.39	500	3847766-	Sensor protein (<i>uhpB</i>)
0.43	348	4053919-	Negative regulation of glnA (glnL or ntrB)
0.70	456	4102556-	Chemosensory transducer (<i>cpxA</i>)
0.38	473	4634265+	Sensor protein (creC)
B. subtilis			
0.42	605	1469428 +	Histidine kinase (kinA)
0.55	428	3229144+	Histidine kinase (kinB)
0.65	145	2443877-	Anti-sigma factor and serine kinase (<i>spoIIAB</i>)

^{*a*} Position in the genome is shown as position of the translation initiation site and orientation of the gene (+, direct strand; –, complementary strand).

A. aeolicus, and *Synechocystis*. We suggested that the high levels of CH and detoxification proteins in *D. radiodurans* contribute to its capacities for prevention of damage to DNA and RNA and for repair of DNA, RNA, and protein damage caused by severe ionizing radiation.

(iii) Electron transfer genes. Many electron transfer PHX genes are prominent in fast-growing bacteria, in the archaeal genomes, in the deeply branching thermophilic eubacteria *A. aeolicus* and *T. maritima*, and in *Synechocystis*, *M. tuberculosis*, and *H. pylori*. By contrast, parasitic eubacteria apparently do not possess electron transfer PHX genes. This is to be expected, since they derive much of their energy from the host.

(iv) Histidine kinase genes. None of the rapidly dividing bacteria possess PHX genes for sensors, histidine kinases, regulatory protein kinases, or chemotaxis proteins (see also Table 7). Of course, there are many metabolic kinases which are highly expressed. By contrast, several PHX histidine kinase genes are contained in *Synechocystis* and *M. tuberculosis*.

Highly expressed RP genes in prokaryotic genomes. In our original predictions of PHX genes for each genome, the RP genes served as a representative group. Following the analysis based on definition I and equations 2 and 3, we observed that practically all RP genes of all sizes qualify as highly expressed. Those with the highest expression levels (arranged by decreasing predicted levels in *E. coli*) are the genes encoding L2, S2, L4, S3, S1, L1, L3, S9, L20, L5, L13, S4, L14, and S13. Among the prokaryotic genomes at hand, distinct RPs number from 50 to 60 (Table 6), and in eukaryotes they number 79 (in yeast, 78) (68, 72, 73). Special cases and distributional properties of RPs stand out, as described below.

The eubacterial RP genes generally feature a large cluster (operon) encompassing 20 to 40% of all RP genes. Some of the main TF including Tuf, Fus, RpoA, RpoB, and RpoC are often encoded within or proximal to the large RP gene operon. Other operons usually consist of two to five RP genes. For example, the cluster of L7-L12, L10, L1, L11, rpoB, and rpoC stands out. *B. subtilis* unites in its genome the equivalents of the two largest *E. coli* clusters. In many genomes (e.g., *Synechocystis* and *M. tuberculosis*), several major CH proteins are proximal to the major RP operons. It is tempting to speculate that these chaperones may contribute to ribosomal assembly. In the presence of a unique *oriC*, the bulk of eubacterial RP clusters are positioned near the origin of replication.

A giant RP (labeled S1, RpsA, or Rps1 and generally exceeding 500 aa) is recognized in most eubacteria. The S1 gene is essential in *E. coli* and putatively contributes to the initiation of protein synthesis. In *Synechocystis*, S1 (327 aa) occurs as a drastically reduced version of the typical S1. The major RP cluster in *Synechocystis* has the genes for RpoB, RpoC, and GroEL-1 nearby. In *B. subtilis*, there is a putative S1 homolog of 380 aa, and S1 is definitely missing from the mycoplasma genomes *Mycobacterium genitalium*, *M. pneumoniae*, and *U. urealyticum*. S1, when extant, is isolated (not part of an RP operon) and tends to score among the highest expression levels. The deeply branching gram-negative *A. aeolicus* encodes an S1 homolog. None of the archaeal genomes possesses an S1 homolog, and eukaryote genomes also lack an S1 homolog.

Unlike the giant eubacterial S1, *Saccharomyces cerevisiae* RP genes are all less than 350 aa in length (mostly between 50 and 250 aa) and are randomly distributed over the 16 yeast chromosomes. This is consistent with the general absence of operons from yeast. Most yeast RPs are duplicated and achieve impressively high expression levels (68).

The S2 RP gene in eubacterial genomes is separated from other RPs. However, S2 in the archaeal genomes (those of *M. thermoautotrophicum, A. fulgidus, P. abyssi*, and *P. horikoshii*) is proximal to RP clusters.

RPs are generally very cationic and tend to bind nucleic acids, particularly RNA. The acidic RPs (containing a carboxyl hyperacidic residue run) P_0 , P_1 , and P_2 are found in eukaryotes; P_0 is generally part of the RP repertoire in the archaeal genomes. *P. horikoshii* contains a ribosomal P_0 -like acidic protein of 341 aa. Acidic RPs have not been detected in eubacterial genomes, except for L7-L12.

Comparison of predicted expression levels with 2-D gel patterns. For some E. coli proteins, 2-D gel electrophoresis data on protein abundances under different growth conditions are available (65, 66). We compared the molar abundances (protein abundance divided by protein molecular weight) of 96 proteins of ≥ 100 aa with the set of PHX genes. The genes for the 20 most molar abundant of the 96 proteins include (in decreasing abundances) tufA, metE, rplL, ompA, fabB, rpsA, rpsF, groEL, eno, fusA, hns, purC, glyA, ilvE, tsf, folA, dnaK, tig, atpA, and *glnA*. Seventeen of these genes were identified as PHX by our method. The three that were not identified as PHX are metE (methionine synthase), ilvE (branched-chain amino acid aminotransferase), and folA (dihydrofolate reductase). Interestingly, all three are involved in amino acid or nucleotide biosynthesis. At the other extreme, among the 20 least abundant proteins of the 96, only five qualify as highly expressed. These include the aminoacyl tRNA synthetases LeuS and ValS, the RP RpII, N utilization substance protein B (NusB), and phosphoenolpyruvate carboxykinase (PckA). The results for the remaining 56 proteins of intermediate molar abundance include 28 identified as highly expressed.

PHX genes and SD signals. Initiation of gene translation in *E. coli* and in many eubacteria involves interactions between a conserved SD sequence immediately upstream of the initiation codon in the mRNA leader and an equally conserved anti-SD sequence at the 3' end of the 16S rRNA. Not all mRNAs possess a recognizable SD sequence. The consensus SD sequence features at its core the purine run GGAGG, generally traversing positions -9 to -5 relative to the initiation codon and the 16S rRNA gene which persistently carries the anti-SD sequence CACCTCCTTTC at its 3' end. The bulk of genomes,

TABLE 8. Shine-Dalgarno (SD) sequence in prokaryotic genomes^a

Genome	Туре	Expression [E(g)] value	No. of genes examined	No. (%) with SD	No. (%) of strong SD sequences with GAGG
		100 0 11		a (a (a)	01 00A0
E. coli	PHX	1.00 - 2.66	306	248 (81)	161 (53)
	PMX	0.50-0.99	258^{b}	156 (66)	82 (34)
	PLX	0.27-0.37	113	64 (57)	35 (31)
V. cholerae	PHX	1.00 - 2.25	238	156 (66)	52 (45)
	PLX	0.27-0.37	113 ^b	65 (57)	48 (25)
H. pylori	PHX	1.00 - 1.21	73	67 (92)	30 (41)
1.	PMX	0.74-0.99	154 ^b	128 (83)	32 (21)
T. maritima	PHX	1.00 - 1.40	181	177 (97)	146 (81)
	PMX	0.74-0.99	209^{b}	192 (92)	148 (71)
A. fulgidus	PHX	1.00-1.39	176	106 (66)	79 (45)
, ,	PMX	0.58-0.99	188^{b}	105 (56)	53 (28)

^{*a*} Putative SD sequences are defined as purine runs \geq 5 bp long within 20 bp upstream of the translation start. A strong SD sequence includes GAGG or GGAG.

A random sample of genes in this category was examined.

including those of all five archaea, have at least one copy of 16S rRNA that has the CCTCCT terminal motif. Two bacterial genomes, those of B. burgdorferi and R. prowazekii, do not have rRNA genes with this motif, and two bacterial genomes (those of Synechocystis and D. radiodurans) have an additional copy of a 16S rRNA gene with a different 3' end. In several genomes, we investigated the proportion of genes in possession of a strong SD sequence among three groups of genes: PHX genes, genes with predicted moderate expression levels (PMX), and genes with predicted low expression levels (PLX). The statistics are displayed in Table 8. The collection of PHX genes examined is complete. A random sample of the PMX and PLX genes was investigated. The data show that more PHX genes than genes with an average or low expression level tend to possess a strong SD sequence, indicating a significant positive correlation between predicted expression levels of genes and the existence of a strong SD sequence.

Gene classes not highly expressed. Proteins required in few copies per cell cycle are not expected to be highly expressed. In fact, the following gene groups are seldom highly expressed: (i) specific regulatory proteins, (ii) specific transcription factors, and (iii) strict replication proteins. We display in Table 7 the expression levels for several two-component sensor genes (histidine kinases) in E. coli and B. subtilis. In all the examples, the expression levels are emphatically low, ranging from 0.30 to 0.70, with most values about 0.40. A second gene group with prevalent low expression levels are those for the repair proteins of D. radiodurans (Table 4). Only the paramount recombination protein, RecA, is significantly highly expressed (E =2.04). However, the bulk of repair proteins of D. radiodurans score in the interval E = 0.4 to 0.8 (Table 4). The repair protein repertoire of E. coli (again with the exception of RecA and RuvB) and those in almost all prokaryotic genomes are not PHX.

Pathways for the synthesis of vitamins, of which only small amounts are generally needed to achieve adequate function, also record low *E* values, about 0.4 to 0.8 (Karlin et al., unpublished). Exceptionally, RibH (riboflavin synthase β subunit) of *E. coli*, in a pathway of vitamin synthesis, is PHX. RibE (riboflavin synthase α subunit; not PHX) forms a complex with RibH composed of three units of RibE joined with 60 units of RibH (52a). This stoichiometric anomaly on RibH makes it likely that RibH furnishes structural support and in this purview may be used in multiple capacities.

Perspectives. Why are CH proteins outstandingly PHX? Their functional attributes are far ranging. Chaperones are vitally needed during both rapid growth and stationary phase. In normal cell physiology, these proteins contribute decisively to ensuring proper protein folding, to correcting misfolded structures, to coordinating protein transport, and to directing protein secretion. Chaperones also contribute to conformational changes and to minimizing protein damage during stress. Accordingly, during starvation, molecular chaperones reduce protein denaturation. Starvation is accompanied by toxic metabolites and oxidative stress. Dps also controls proteins concerned with oxidative protection. A large number of starvation proteins are involved in protein and DNA repair (43). Overall, the three protein families-RPs, TFs, and CHs—are needed in large quantities at many stages of the life cycle, and putatively the organism has evolved codon usages to promote, as needed, growth, stability, and plasticity. From this perspective, codon usage has evolved to accommodate most situations of the cell's existence.

Protein synthesis can be divided into four essential steps: initiation, elongation, termination, and ribosome disassembly. The major highly expressed proteins involved in these processes rely on initiation factors (InfB and InfC), elongation factors (EF-Tu, EF-G, and EF-Ts), and the ribosome release factor (Rrf). The principal genes of the transcriptome also feature the prime components of RNA polymerase, RpoA, RpoB, and RpoC, sometimes supplemented by the sigma factors RpoD, RpoE, and RpoH and terminators-antiterminators (NusA, NusB, and NusG) (Table 2). However, not all transcription factors are PHX. For example, RpoA is selectively highly expressed, and most σ factors are not PHX. EF-Tu and EF-G genes often localize in eubacterial genomes among RP clusters, but these are not found near to archaeal RP operons. For eubacteria with a unique bidirectional origin of replication, PHX genes are predominantly encoded from the leading strand.

More than 80% of PHX genes possess an unambiguous SD sequence compared to genes of average or lower expression levels, with percentages indicating a positive association of E(g) values and an extant strong SD sequence. Generally, PHX genes exploit favorable codon usages, tend to possess a strong SD sequence, and are probably endowed with a strong promoter sequence.

Questions for future study include the following. Are the prime prokaryotic PHX genes "ancient," meaning significantly conserved across many genomes? How does proteome content (in terms of protein abundances) correlate with transcriptome data? This also concerns correlations of 2-D-gel assessments with mRNA levels. Several reports depict these correlations as weak (21, 60). What are the core (essential) numbers and types of genes that genomes require for fast growth? The rapidly growing bacteria E. coli, H. influenzae, V. cholerae, B. subtilis, and D. radiodurans attain the highest expression levels [E(g) values] of genes among bacterial genomes (Table 1). What is the relation of induced versus constitutive protein expression to PHX genes? What is the influence of stoichiometry of subunits or the half-life of a protein on expression levels? Do operons and complexes entail components concordant or discordant with respect to their PHX status? In these contexts, PHX ORFs are attractive targets for knockout studies.

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