Whole-Genome Random Sequencing and Assembly of Haemophilus influenzae Rd

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An approach for genome analysis based on sequencing and assembly of unselected pieces of DNA from the whole chromosome has been applied to obtain the complete nucleotide sequence (1,830,137 base pairs) of the genome from the bacterium *Haemophilus influenzae* Rd. This approach eliminates the need for initial mapping efforts and is therefore applicable to the vast array of microbial species for which genome maps are unavailable. The *H. influenzae* Rd genome sequence (Genome Sequence DataBase accession number L42023) represents the only complete genome sequence from a free-living organism.

A prerequisite to understanding the complete biology of an organism is the determination of its entire genome sequence. Several viral and organellar genomes have been completely sequenced. Bacteriophage $\phi X174$ [5386 base pairs (bp)] was the first to be sequenced, by Fred Sanger and colleagues in 1977 (1). Sanger et al. were also the first to use strategy based on random (unselected) pieces of DNA, completing the genome sequence of bacteriophage λ (48,502 bp) with cloned restriction enzyme fragments (1). Subsequently, the 229-kb genome of cytomegalovirus (CMV) (2), the 192-kb genome of vaccinia (3), and the 187-kb mitochondrial and 121-kb chloroplast genomes of Marchantia polymorpha (4) have been sequenced. The 186-kb genome of variola (smallpox) was the first to be completely sequenced with automated technology (5).

At the present time, there are active genome projects for many organisms, including Drosophila melanogaster (6), Escherichia coli (7), Saccharomyces cerevisiae (8), Bacillus subtilis (9), Caenorhabditis elegans (10), and

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Homo sapiens (11). These projects, as well as viral genome sequencing, have been based primarily on the sequencing of clones usually derived from extensively mapped restriction fragments, or λ or cosmid clones. Despite advances in DNA sequencing technology (12) the sequencing of genomes has not progressed beyond clones on the order of the size of λ (~40 kb). This has been primarily because of the lack of sufficient computational approaches that would enable the efficient assembly of a large number (tens of thousands) of independent, random sequences into a single assembly.

The computational methods developed to create assemblies from hundreds of thousands of 300- to 500-bp complementary DNA (cDNA) sequences (13) led us to test the hypothesis that segments of DNA several megabases in size, including entire microbial chromosomes, could be sequenced rapidly, accurately, and cost-effectively by applying a shotgun sequencing strategy to whole genomes. With this strategy, a single random DNA fragment library may be prepared, and the ends of a sufficient number of randomly selected fragments may be sequenced and assembled to produce the complete genome. We chose the free-living organism Haemophilus influenzae Rd as a pilot project because its genome size (1.8 Mb) is typical among bacteria, its G+C base composition (38 percent) is close to that of human, and a physical clone map did not exist.

Haemophilus influenzae is a small, nonmotile, Gram-negative bacterium whose only

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natural host is human. Six H. influenzae serotype strains (a through f) have been identified on the basis of immunologically distinct capsular polysaccharide antigens. Non-typeable strains also exist and are distinguished by their lack of detectable capsular polysaccharide. They are commensal residents of the upper respiratory mucosa of children and adults and cause otitis media and respiratory tract infections, mostly in children. More serious invasive infection b caused almost exclusively by type b strains, with meningitis producing neurological sequelae in up to 50 percent of affected children. A vaccine based on the type b capsulat antigen is now available and has dramatically reduced the incidence of the disease in Europe and North America.

Genome sequencing. The strategy for a shotgun approach to whole genome sequencing is outlined in Table 1. The theory follows from the Lander and Waterman (14) application of the equation for the Poisson distribution. The probability that a base is not sequenced is $P_0 = e^{-m}$, where m is the sequence coverage. Thus after 1.83 Mb of sequence has been randomly generated for the *H. influenzae* genome (m = 1, 1× coverage), $P_o = e^{-1} = 0.37$ and approximately 37 percent of the genome is unsequenced. Fivefold coverage (approximately 9500 clones sequenced from both insert ends and an average sequence read length of 460 bp) yields $P_0 = e^{-5} = 0.0067$, or 0.67 percent unsequenced. If L is genome length and n is the number of random sequence segments done, the total gap length is Le^{-m} . and the average gap size is L/n. Fivefold coverage would leave about 128 gaps averaging about 100 bp in size.

To approximate the random model during actual sequencing, procedures for library construction (15) and cloning (16) were developed. Genomic DNA from H. influenzae Rd strain KW20 (17) was mechanically sheared, digested with BAL 31 nuclease to produce blunt ends, and size-fractionated by agarose gel electrophoresis. Mechanical shearing maximizes the randomness of the DNA fragments' Fragments between 1.6 and 2.0 kb in size were excised and recovered. This narrow range was chosen to minimize variation in growth of clones. In addition, we chose this maximum size to minimize the number of complete genes that might be present in a single fragment, and thus might be lost as a result of expression of deleterious gene products. These fragments were ligated to Sma I-cut, phosphatase-treated pUC18 vector, and the ligated products were fractionated on an agarose gel. The linear vector plus insert band was excised and recovered. The ends of the linear recombinant molecules were repaired with T4 polymerase, and the molecules were then ligated into circles. This two-

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Plasmid I ble-stranded were prepare collaboration Technology were prepare stages of DN growth throu Template co with Hoechst 2350. DNA co ed, but low-yi were identifie quenced. Ter from two H. i (20). An amp in vector λ (library was cor libraries conta of 15 to 20 kb prepared from plates were pre tesin (Qiagen) carried out on of a Catalyst L Ready Reactic mencing Kits M13-21) and

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stage procedure resulted in a collection of ingle-insert plasmid recombinants with ninimal contamination from double-insert himeras (<1 percent) or free vector (<3) Fircent). Because deviation from randomless is most likely to occur during cloning, 3. coli host cells deficient in all recombinaion and restriction functions (18) were sed to prevent rearrangements, deletions, and loss of clones by restriction. Transmed cells were plated directly on antibitic diffusion plates (16) to avoid the usual noth recovery phase that would have alwed multiplication and selection of the 10st rapidly growing cells and could lead to eviation from randomness. All colonies ere used for template preparation regardss of size. Only clones lost because of pression of deleterious gene products -ould be deleted from the library, resulting a slight increase in gap number over that spected.

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To evaluate the quality of the H. influnzae library, sequence data were obtained ∞ ~4000 templates by means of the 組3-21 primer. Sequence fragments were sembled with the AUTOASSEMBLER stware [Applied Biosystems division of erkin-Elmer (AB)] after obtaining 1300, 800, 2500, 3200, and 3800 sequence fragients, and the number of unique assembled 1se pairs was determined. The data obined from the assembly of up to 3800 squence fragments were consistent with a bisson distribution of fragments with an verage "read" length of 460 bp for a geme of 1.9×10^6 bp, indicating that the ld brary was essentially random.

Plasmid DNA templates that were dourk-stranded and of high quality (19,687) ırere prepared by a method developed in rv llaboration with Advanced Genetic echnology Corporation (19). Plasmids re ere prepared in a 96-well format for all nlly ges of DNA preparation from bacterial to with through final DNA purification. bv Amplate concentration was determined al th Hoechst dve and a Millipore Cytofluor he 50. DNA concentrations were not adjust-.6 but low-yielding templates ($<30 \text{ ng/}\mu$ l) vre identified where possible and not senenced. Templates were also prepared ıdm two H. influenzae λ genomic libraries n-1). An amplified library was constructed nat vector λ GEM-12 and an unamplified nd rary was constructed in λ DASH II. Both on raries contained inserts in the size range ig-15 to 20 kb. Liquid lysates (10 ml) were OSpared from selected plaques and temliites were prepared on an anion-exchange gain (Qiagen). Sequencing reactions were nd ried out on plasmid templates by means he a Catalyst LabStation (AB) and PRISM red ady Reaction Dye Primer Cycle Seles encing Kits (AB) for the M13 forward vo- 113-21) and the M13 reverse (M13RP1)

primers (21). Dye terminator sequencing reactions were carried out on the λ templates on a Perkin-Elmer 9600 Thermocycler with the Applied Biosystems Prism Ready Reaction Dye Terminator Cycle Sequencing Kits. We used T7 and SP6 primers to sequence the ends of the inserts from the λ GEM-12 library and T7 and T3 primers to sequence the ends of the inserts from the λ DASH II library. Sequencing reactions (28,643) were performed by eight individuals using an average of 14 AB 373 DNA Sequencers per day over a 3-month period. All sequencing reactions were analyzed with the Stretch modification of the AB 373 sequencer. These sequencers were modified to include a heat plate and the height of the laser was reduced. With standard gel plates the "well-to-read" length was increased to 34 cm when standard sequencing plates were used and to 48 cm when 60-cm plates were used. The sequencing reactions in this project were analyzed primarily with a 34-cm well-to-read distance. The overall sequencing success rate was 84 percent for M13-21 sequences, 83 percent for M13RP1 sequences, and 65 percent for dye-terminator reactions. The average usable read length was 485 bp for M13-21 sequences, 444 bp for M13RP1 sequences, and 375 bp for dye-terminator reactions. The highthroughput sequencing phase of the project is summarized in Table 2.

We balanced the desirability of sequencing templates from both ends, in terms of ordering of contigs and reducing the cost of lower total number of templates, against shorter read lengths for sequencing reactions performed with the M13RP1 primer compared to the M13-21 primer. Approximately one-half of the templates were sequenced from both ends. Altogether, 9297 M13RP1 sequencing reactions were done. Random reverse sequencing reactions were done on the basis of successful forward se-

quencing reactions. Some M13RP1 sequences were obtained in a semidirected fashion; for example, M13-21 sequences pointing outward at the ends of contigs were chosen for M13RP1 sequencing in an effort to specifically order contigs. The semidirected strategy was effective, and clone-based ordering formed an integral part of assembly and gap closure.

In the course of our research on expressed sequence tags (ESTs), we developed a laboratory information monagement system for a large-scale sequencing laboratory (22). The system was designed to automate data flow wherever possible and to reduce user error. It has at its core a series of databases developed with the Sybase relational data management system. The databases store and correlate all information collected during the entire operation from template preparation to final analysis. Although the system was originally designed for EST projects, many of its features were applicable or easily modified for a genomic sequencing project. Because the raw output of the AB 373 sequencers is collected on a Macintosh system and our data management system is based on a Unix system, it was necessary to design and implement multiuser, client-server applications that allow the raw data as well as analysis results to flow seamlessly into the database with a minimum of user effort. To process data collected by the AB 3735, sequence files were first analyzed with FACTURA, an AB program that runs on the Macintosh and is designed for automatic vector sequence removal and end-trimming of sequence files. The Macintosh program ESP, written at The Institute for Genomic Research (TIGR), loaded the feature data extracted from sequence files by FAC-TURA to the Unix-based H. influenzae relational database. Assembly was accom-

Table 1. Whole-genome sequencing strategy.

Stage	Description		
Random small insert and large insert library construction	Shear genomic DNA randomly to ~2 kb and 15 to 20 kb, respectively		
Library plating	Verify random nature of library and maximize random selection of small insert and large insert clones for template production		
High-throughput DNA sequencing	Sequence sufficient number of sequence fragments from both ends for 6× coverage		
Assembly	Assemble random sequence fragments and identify repeat regions		
Gap closure			
Physical gaps	Order all contigs (fingerprints, peptide links, λ clones, PCR) and provide templates for closure		
Sequence gaps	Complete the genome sequence by primer walking		
Editing	Inspect the sequence visually and resolve sequence ambiguities, including frameshifts		
Annotation	Identify and describe all predicted coding regions (putative identifications, starts and stops, role assignments, operons, regulatory regions)		

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plished by first retrieving a specified set of sequence files and their associated features by means of STP, another TIGR program, which is an X-windows graphical interface that retrieves sequences from the database with user-defined queries.

TIGR ASSEMBLER is the software component that enabled us to assemble the H. influenzae genome. It simultaneously clusters and assembles fragments of the genome. In order to obtain the speed necessary to assemble more than 10⁴ fragments, the algorithm builds a table of all 10-bp oligonucleotide subsequences to generate a. list of potential sequence fragment overlaps. When TIGR ASSEMBLER is used, a single fragment begins the initial contig; to extend the contig, a candidate fragment is chosen with the best overlap based on oligonucleotide content. The current contig and candidate fragment are aligned by a modified version of the Smith-Waterman (23) algorithm, which provides for optimal gapped alignments. The contig is extended by the fragment only if strict criteria for the quality of the match are met. The match criteria include the minimum length of overlap, the maximum length of an unmatched end, and the minimum percentage match. The algorithm automatically lowers these criteria in regions of minimal coverage and raises them in regions with a possible repetitive element. The number of potential overlaps for each fragment determines which fragments are likely to fall into repetitive elements. Fragments representing the boundaries of repetitive elements and potentially chimeric fragments are often rejected on the basis of partial mismatches at the needs of alignments and excluded from the contig.

TIGR ASSEMBLER was designed to take advantage of clone size information coupled with sequence information from both ends of each template. It enforces the

Table 2. Summary of features of whole-genome sequencing of H. influenzae Rd.

Description	Number
	19,687
Double-stranded templates	19,346
Forward-sequencing reactions (M13-21 primer)	16,240 (84)
Successful (%)	485
Average edited read length (bp)	9,297
Reverse sequencing reactions (M13RP1 primer)	7,744 (83)
Successful (%)	444
Average edited read length (bp)	24,304
Sequence fragments in random assembly	11,631,485
Total base pairs	140
Contigs	42
Physical gap closure	37
PCR	15
Southern analysis	23
λ clones	2
Peptide links	3,530
Terminator sequencing reactions*	2,404 (68)
Successful (%)	375
Average edited read length (bp)	1,830,137
Genome size (bp)	
G+C content (%)	6
rRNA operons	723
rrnA, rrnC, rrnD (spacer region) (bp)	478
rmB, rmE, rmF (spacer region) (bp)	54
tRNA genes identified	1,743
Number of predicted coding regions	736 (42)
Unassigned role (%)	389
No database match	347
Match hypothetical proteins	1,007 (58)
Assigned role (%)	68 (6.8
Amino acid metabolism	
Biosynthesis of cofactors, prosthetic groups, and carriers	54 (5.4)
Cell envelope	84 (8.3)
Cellular processes	53 (5.3)
Central intermediary metabolism	30 (3.0)
Energy metabolism	105 (10.4
Fatty acid and phospholipid metabolism	25 (2.5
Purines, pyrimidines, nucleosides and nucleotides	53 (5.3
Regulatory functions	. 64 (6.3
Replication	87 (8.6
Transcription	27 (2.7
Translation	141 (14.0
Transport and binding proteins	123 (12.2)
Other	93 (9.2)

*Includes gap closure, walks on rRNA repeats, random end-sequencing of λ clones for assembly confirmation, and alternative reactions for ambiguity resolution.

constraint that sequence fragments from two ends of the same template point to ward one another in the contig and are located within a certain range of base pairs (definable for each clone on the basis of the insert length or the clone size range for a given library). In order for the assembly process to be successful it was essential that the sequence data be of the highest quality and that sequence fragment lengths be sufficient to span most small repeats. Less than 13 percent of our random sequence fragments were smaller than 400 bp after vector removal and end trimming. Assembly of 24,304 sequence fragments of H. influenzae required 30 hours of central processing unit time with the use of one processor on a SPARCenter 200 containing 512 Mb of RAM. This process resulted in approximately 210 contigs. Because of the high stringency of the TIGR ASSEMBLER, all contigs were searched against each other with GRASTA, which is a modified version of the program FASTA (24). In this way, additional overlaps that enabled compression of the data set into 140 contigs were detected. The location of each fragment in the contigs and extensive information about the consensus sequence itself were loaded into the H. influenzae relational database. After assembly, the relative positions of

After assembly, the relative positions of the 140 contigs were unknown. The program ASM_ALIGN, developed at TIGR, identified clones whose forward and reverse sequencing reactions indicated that the were in different contigs and ordered and displayed these relationships. With this program, the 140 contigs were placed into 4. groups totaling 42 physical gaps (no template DNA for the region) and 98 sequence gaps (template available for gap closure).

Four integrated strategies were developed to order contigs separated by physical gaps. Oligonucleotide primers were designed and synthesized from the end of each contig group. These primers were then available for use in one or more of the strategies outlined below:

1) DNA hybridization (Southern) analysis was done to develop a "fingerprint" for a subset of 72 of the above oligonucleotides This procedure was based on the supposttion that labeled oligonucleotides homologous to the ends of adjacent contigs should hybridize to common DNA restriction fragments, and thus share a similar or identical hybridization pattern or fingerprint (25). Adjacent contigs identified in this manner were targeted for specific PCR reactions.

2) Peptide links were made by searching each contig end with BLASTX (26) against a peptide database. If the ends of two contigs matched the same database sequence appropriately, then the two contigs were tentatively considered to be adjacent. Figure region shown role ca bers co nation

Table 3

bers are its role (Sim) o shown. is BLOS informa http://ww notransf zyme A: ethyl sul phate (acetylglu methyltr tein: PR adenosy transfera other spa identifica variabilis 0404(47) 0687(49) 1648(65) aphidicol vinosum: kluyveri: (1590(74) 0003(52) 0044(80) 0065(75) 0107(54) 0147(61) 0182(60) 0227(51) 0257(76) 0282(59) 0329(79), 0344(85), 0371(84),

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The Genome of Haemophilus influenzae Rd

Figure 2. Gene map of the *H. influenzae* Rd genome. Predicted coding regions are shown on each strand. The rRNA and tRNA genes are shown as lines and triangles, respectively. Genes are color-coded by role category as described in the Figure key. Gene identification numbers correspond to those in Table 3. Where possible, three-letter designations are also provided. In the region containing ribosomal proteins

HI0782-HI0796 some identification numbers have been omitted because of space limitations. Predicted coding regions with similarity to database sequences designated as hypothetical coding regions are represented as white, cross-hatched rectangles. Predicted coding regions that have no database match are represented as white, unfilled rectangles.

overe data Table 3. Identification of H. influenzae genes. Gene identification numbers are listed with the prefix HI in Fig. 3. Each identified gene is listed in . The its role category [adapted from Riley (36)]. The percentage of similarity ontigs (Sim) of the best match to the NRBP (as described in the text) is also e conshown. The amino acid substitution matrix used in the BLAZE analysis to the is BLOSUM60. An expanded version of this table with additional match information, including species, is available via World Wide Web (URL: ions of http://www.tigr.org/). Abbreviations used: Ac, acetyl; ATase, amie pro- notransferase; BP, binding protein; biosyn, biosynthesis; CoA, coenzyme A; DCase, decarboxylase; DHase, dehydrogenase; DMSO, dim-TIGR, ethyl sulfoxide; f-Met, formylmethionine; G3PD, glyceraldehyde-3-phosreverse phate dehydrogenase; GABA, γ-aminobutyric acid; GlcNAc, Nit they acetylglucosamine; LOS, Lipooligosaccharide; lpp, lipoprotein; MTase, ed and methyltransferase; MurNAc, N-acetylmuramyl; P, phosphate; prt, pronis protein; PRTase, phosphoribosyltransferase; RDase, reductase; SAM, Sinto 42 adenosylmethionine; Sase, synthase-synthetase; sub, subunit; Tase, o tem- transferase. The following hypothetical proteins were matched from the other species as indicated (percent similarity in parentheses after gene quence identification number): Alcaligenes eutrophus: 1053(52); Anabaena sure). variabilis: 1349(54); Bacillus subtilis: 0115(53), 0259(54), 0355(61), devel- 0404(47), 0415(69), 0416(63), 0417(66), 0454(64), 0456(56), 0522(54), physical 0687(49), 0775(54), 0959(50), 1083(53), 1203(63), 1627(59), 1647(81), ere de- 1648(65), 1654(64); Bacteriophage P22: 1412(54); Buchnera of each aphidicola: 1199(65); Campylobacter jejuni: 0560(71); Chromatium e then vinosum: 0105(75); Clostridium acetobutylicum: 0773(72); Clostridium kluyveri: 0976(48); Clostridium perfringens: 0†43(58); Coxiella burnetii: of the 1590(74), 1591(50); Erwinia carotovora: 1436(72); Escherichia coli: 0003(52), 0012(67), 0017(91), 0028(68), 0033(90), 0034(84), 0035(79), n) anal-0044(80), 0045(67), 0050(70), 0051(50), 0052(56), 0053(56), 0059(72), rint" for 0065(75), 0072(65), 0081(71), 0091(72), 0092(49), 0093(59), 0103(71), eotides. 0107(54), 0108(65), 0125(88), 0126(87), 0135(68), 0145(69), 0146(58), supposi-147(61), 0148(62), 0162(47), 0172(67), 0174(84), 0175(70), 0176(87), 0182(60), 0183(66), 0184(73), 0187(58), 0188(81), 0198(75), 0203(86), nomolo-0227(51), 0230(71), 0232(69), 0235(80), 0241(82), 0242(50), 0258(95), s should 2257(76), 0265(77), 0266(83), 0270(80), 0271(73), 0276(70), 0281(76), ion frag 0282(59), 0293(61), 0303(81), 0306(70), 0308(58), 0315(87), 0316(68), identical0329(79), 0336(91), 0338(68), 0340(72), 0341(84), 0342(60), 0343(67), nt (25),0344(85), 0345(82), 0346(77), 0347(67), 0364(55), 0365(86), 0367(48), manner³371(84), 0374(64), 0375(62), 0376(75), 0379(57), 0380(58), 0386(76);

0393(93), 0396(54), 0398(72), 0400(65), 0409(69), 0412(85), 0418(68). 0423(67), 0424(66), 0431(76), 0432(68), 0442(93), 0452(73), 0464(78), 0467(80), 0493(64), 0494(69), 0500(63), 0508(82), 0509(69), 0510(74), 0519(71), 0520(59), 0521(58), 0562(83), 0565(63), 0568(71), 0570(80), 0572(70), 0574(63), 0575(80), 0576(65), 0597(57), 0617(54), 0624(72), 0626(81), 0634(78), 0638(68), 0647(64), 0656(74), 0658(56), 0668(76), 0670(83), 0671(87), 0696(54), 0697(64), 0700(77), 0702(71), 0719(86), 0721(78), 0723(73), 0724(64), 0730(65), 0733(55), 0744(70), 0755(61), 0756(60), 0766(87), 0767(72), 0810(74), 0817(68), 0826(70), 0827(86), 0831(77), 0837(74), 0839(69), 0840(72), 0841(66), 0849(75), 0851(71), 0852(66), 0855(75), 0858(68), 0860(86), 0862(81), 0864(92), 0878(71), 0881(81), 0890(69), 0891(79), 0906(71), 0918(81), 0929(58), 0933(71), 0934(52), 0935(63), 0936(64), 0943(83), 0948(67), 0955(72), 0956(73), 0963(67), 0965(81), 0979(79), 0984(79), 0986(81), 0988(85), 1000(80), 1001(75), 1005(61), 1007(86), 1010(53), 1019(65), 1020(65), 1021(71), 1024(67), 1026(85), 1027(72), 1028(77), 1029(83), 1030(62), 1031(87), 1032(79), 1064(57), 1072(57), 1073(62), 1082(67), 1084(61), 1085(76), 1086(89), 1089(70), 1090(82), 1091(76), 1092(73), 1093(72), 1094(81), 1095(79), 1096(64), 1104(53), 1118(84), 1125(87), 1129(77), 1130(80), 1146(80), 1147(68), 1148(88), 1149(73), 1150(59), 1151(81), 1153(84), 1155(79), 1165(87), 1181(68), 1195(76), 1198(85), 1216(73), 1234(80), 1240(77), 1243(74), 1252(93), 1262(61), 1280(71), 1282(74), 1288(84), 1289(74), 1297(67), 1298(69), 1300(58), 1301(82), 1309(67), 1314(70), 1315(66), 1333(79), 1337(84), 1342(57), 1364(56), 1368(53), 1369(44), 1437(72), 1463(84), 1542(61), 1545(80), 1558(62), 1598(58), 1608(76), 1612(72), 1628(61), 1643(70), 1652(68), 1653(88), 1655(56), 1656(69), 1657(65), 1664(50), 1677(72), 1679(69), 1703(74), 1704(73), 1714(78), 1715(86), 1721(71), 1723(92); Klebsiella pneumoniae: 0021(63); Lactobacillus johnsonii: 0112(54), 1720(55); Lactococcus lactis: 0555(69); Mycobacterium leprae: 0004(62), 0019(62), 0136(58), 0260(56), 0694(54), 0740(56), 0920(57), 1663(55); Mycoplasma hyopneumoniae: 1281(71); Pasteurella haemolytica: 0219(92); Pseudomonas aeruginosa: 0090(68), 0177(56); Rhodobacter capsulatus: 0170(62), 0672(59), 1439(65), 1683(75), 1684(60), 1688(58); Salmonella typhimurium: 0405(51), 0964(67), 1434(76), 1607(51); Shigella flexneri: 0277(52); Streptococcus parasanguis: 0359(65); Synechococcus sp.: 0961(70); Vibrio parahaemolyticus: 0323(87), 0325(75); Vibrio sp.: 0333(70); Yersinia enterocolitica: 0753(69)

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Identification %Sim <u>HI#</u> **Identification** <u>H</u>I# Amino acid biosynthesis 0457 aminodeoxychorismate lyase (pabC) 67 55 Aromatic amino acid family 0970 3-dehydroquinase (aroQ) 0208 3-dehydroquinase (aroB) 0472 amidotransferase (hisH) 1387 anthranilate Sase component II (trpE) 1388 anthranilate Sase component II (trpD) 1399 anthranilate Sase component II (trpD) 1629 dedA 1629 dedA 0699 dehydrofolate RDase, type I (folA) 1366 dihydropteroate Sase (folP) 1464 dihydropteroate Sase (folP) 1261 folylpolyglutamate Sase (folC) 1447 GTP cyclohydrolase I (folE) 1170 p-aminobenzoate Sase (pabB) 83 68 71 71 68 79 54 70 73 74 anthranilate isomerase (trpC) anthranilate Sase Gln amidotransferase 1389 75 59 Heme and porphyrin 1160 ferrochelatase (visA) 0113 heme utilization prt (hxuC) 0263 heme-hemopexin utilization (hxuB) (trpG) 0468 ATP PRTase (hisG) 69 82) ATP PHTase (INSG)) chorismate mutase (IyrA) i chorismate mutase-prephenate dehydratase (pheA) i chorismate Sase (aroC) DAHP Sase (aroC) dehydroquinase shikimate DHase orobyrru udet bikimaterbasphateS) 46 77 75 99 52 Vess and the second seco 0196 88 64 57 84
 UcU/
 denydroquinase shikimate DHase
 48

 1589 enolpyruvylshikimatephosphateSyn (aroA)
 98

 1166 Gln amidotransferase (hisH)
 61

 0499 histidinol dehydrogenase (hisD)
 78

 0474 hisF cyclase (hisF)
 91

 0470 histidinol-P ATase (hisC)
 77

 0471 imidazoleglycerol-P dehydratase (hisB)
 81

 0475 phosphoribosyl-AMP cyclohydrolase
 77

 (hisIE)
 77
 0607 73 60 Lipoate 0026 lipoate biosyn prt A (lipA) 0027 lipoate biosyn prt B (lipB) 84 Menaguinone and ubiquinone (hislE) (hisIE)
 0473 phosphoribosylformimino-5-aminoimidazole caarboximde ribotide isomerase (hisA)
 055 shikimate 5-DHase (aroE)
 0207 shikimic acid kinase I (aroK) 0283 2-succinyl-6-hydroxy-2,4-cyclohexadiene-1-carboxylate Sase (menD) 0969 4-(2'-carboxyphenyl)-4-oxybutyric acid 64 77 Sase (menC) 189 coenzyme PQQ synthesis prt III (pqdIII) 49 0968 dihydroxynaphthoic acid Sase (menB) 1438 famesyldiphosphate Sase (ispA) 71 71 88 1432 Trp Sase α chain (trpA) 73 71 67 1431 Trp Sase β chain (trpB) 90 0194 O-succinylbenzoate-CoA Sase (menE) Aspartate family 0564 Asn Sase A (asnA) 0286 Asp ATase (aspC) 1617 Asp ATase (aspC) Molybdopterin 77 54 79 85 1675 molybdenum biosyn prt A (moaA) 1675 molybdenum biosyn prt C (moaC) 1370 molybdenum-pterin-BP (mopl) 78 89 74 73 53 1617 ASD Class (43)CU 0646 Asp-semialdehyde DHase (asd) 1632 aspartokinase III (lysC) 0089 aspartokinase-homoserine DHase (thrA) 1042 B12-dependent homocysteine-N5-1448 molybdopterin biosyn prt (chlE 73 77 70 118 molybdopterin biosyn prt (chlN) 1449 molybdopterin biosyn prt (chlN) 1674 molybdopterin converting factor, sub 1 78 79 methyltetrahydrofolate transmethylase (moaD) metH) 1673 molybdopterin converting factor, sub 2 (moaE) 0844 molybdopterin-dinucleotide biosyn prt 76 0122 β-cystathionase (metC) 84 0086 cystathionine γ-Sase (metB) 1308 dehydrodipicolinate RDase (dapB) R 62 83 79 (mob) 1308 oenydrodpicolintale HDase (ddpb)
 0727 diaminopimelate DCase (lysA)
 0750 diaminopimelate epimerase (ddpF)
 0255 dihydrodipicolinate Sase (ddpA)
 1263 homoserine acetyltransferase (met2)
 088 homoserine kinase (thrB)
 0102 succinyl-diaminopimelate desuccinylase
 (ddpC) 8 Pantothenate õ 0953 pantothenate metabolism flavoprotein 77 (dfp) 0631 pantothenate kinase (coaA) 81 78 80 (dapE) Pvridoxine (dapb) 1634 tetrahydrodipicolinate *N*-succinyltransferase (dapD) 1702 tetrahydropteroyltrigiutamate MTase (metb) 0087 Thr Sase (thrC) 99 0863 pyridoxamine phosphate oxidase (pdxH) 65 68 Riboflavin 7rt4 3:4-dihydroxy-2-butanone 4-P Sase (ribB) 83 (212 GTP cyclohydrolase II (ribA) 0944 riboffavin biosyn prt (ribG) 76 1613 riboflavin Sase a chain (ribC) 82 81 Branched chain family 0989 3-isopropylmalate dehydratase (leuD) 0987 3-isopropylmalate DHase (leuB) 0737 acetohydroxy acid Sase II (livG) 1585 acetolactate Sase III large chain (ilvH) 1584 acetolactate Sase III small chain (ilvH) 86 1303 riboflavin Sase β chain (ribE) 90 80 Thioredoxin, glutaredoxin, and glutathione 0161 glutathione RDase (gor) 1115 thioredoxin (trxA) 1159 thioredoxin (trxA) 85 193 branched-chain amino acid transat 0738 dihydroxyacid dehydrase (ilvD) 0983 α isopropylmalate Sase (leuA) 59 49 62 90 100 0084 thioredoxin m (trxM) 79 0682 ketol acid reductoisomerase (ilvC) 90 Cell envelope Membranes, lipoproteins, and porins 1579 15 kD peptidoglycan-assoc lop (lpp) 0620 28 kD membrane pt (lipA) 0302 apolipoprotein Avacylitansferase (cute) 0407 hydrophobic membrane pt Glutamate family 0811 argininosuccinate lyase (argH) 1727 argininosuccinate Sase (argG) 0900 γ-glutamyl kinase (proB) 84 100 87 80 64 61 1239 y-glutamyl-P RDase (proA) 79 (ioA)
 (ioA)
 (ioA) 67 51 1865 Glin Sase (glnA) 0189 Glu DHase (gdhA) 0596 omithine carbamoyltransferase (arcB) 1719 uridylyl Tase (glnD) 86 (iroA) 6933 lpp (hel) 6933 lpp (hpD) 703 lpp (hpD) 703 lpp B (hpB) 6894 membrane fusion prt (mtrC) 6401 outer membrane prt P2 (ompP1) 1194 outer membrane prt P5 (ompA) 904 prolipoprotein diacylglyceryl Tase (lgt) 0300 rare lpp A (rlpA) 0322 rare lpp B (rlpB) 100 68 65 90 Pyruvate family 1575 Ala racemase, biosynthetic (alr) 54 97 98 75 Serine family 1102 Cys Sase (cysK) 103 Cys Sase (cysK) 045 phosphodycerate DHase (serA) 1167 phosphoserine ATase (serC) 1033 phosphoserine phosphatase (serB) 0606 Ser acetyltransferase (cysE) 0889 Ser hydroxymethyltransferase (glyA) 96 30 80 58 62 76 84 84 72 70 88

 Murein sacculus and peptidoglycan

 1140 D-Ala-D-Ala igase (ddlB)

 7630 D-alanyl-D-Ala carboxypeptidase (dacB)

 7133 GlcNAc transferase (murG)

 764

 770440 penicillin-BP (ponA)

 7725 penicillin-BP 1B (ponB)

 766 penicillin-BP 2 (bpp2)

 746

 768 penicillin-BP 5 (dacA)

 769 penicillin-BP 5 (dacA)

 760 penicillin-BP 5 (dacA)

 Biosynthesis of cofactors, prosthetic groups, and carriers Biotin 1554 7,8-diamino-pelargonic acid ATase (bioA) 1553 7-keto-8-aminopelargonic acid Sase (bioF) 74 56 (bioF) 1551 biotin synthesis prt (bioC) 0643 biotin sulfoxide RDase (bisC) 1022 biotin Sase (bioB) 1550 dethiobiotin Sase (bioD) 1445 dethiobiotin Sase (bioD) 47 72 (mepA) poptidoglycan-assoc outer membrane lpp 100 (pal) phospho-N-acetylmuramoyl-pentapeptide-89 0381 R 1135 Tase E (mrat) 0031 rod shape-determining pt (mreB) 0037 rod shape-determining pt (mreB) 0038 rod shape-determining pt (mreC) 0039 rod shape-determining pt (mreC) Folic acid 1444 5,10-methylenetetrahydrofolate RDase 81 83 90 74 72 59 (file) (folD) 0069 5,10-methylenetetrahydrofolate DHase (folD) 0064 7,8-dihydro-6-hydroxymethylpterin-pyrophosphokinase (folK) 82 0240 0829 soluble lytic murein transglycosylase (slt) 1081 UDP-GICNAc enolpyruvyl Tase (murZ) 082 78 85

%Sim <u>Hi#</u> **Identification** %Sim 1139 UDP-MurNAc-Ala ligase (murC) 1136 UDP-MurNAc-Ala-DGlu ligase (murD) 1134 UDP-MurNAc-pentapeptide Sase (murF) 1133 UDP-MurNAc-tripeptide Sase (murE) 82 68 73 76 UDP-NAc-enolpyruvoylglucosamine RDase (murB) 0268 Surface polysaccharides, lipopolysaccharides and Sufface pory-account of the second se 92 70 (kdtA) ADP-heptose-lps heptosyltransferase II 1105 79 (rfaF) 1114 ADP-L-glycero-D-mannoheptose-6-88 ADP-L-glycero-D-mannoheptose-6-epimerase (rfaD)
O68 CTP-CMP-3-deoxy-D-manno-octulosonate-cytidylyt-transferase (kdsB)
0868 glycosyl Tase (lgtD)
1578 glycosyl Tase (lgtD)
1578 kpsF prt (kpsF)
1537 lic-1 operon pt (licA)
1538 lic-1 operon pt (licB)
1539 lic-1 operon pt (licC)
1540 lic-1 operon pt (licD) 82 55 100 99 399 94 77 1540 lic-1 operon prt (licD) 1060 lipid A disaccharide Sase (lpxB) 0765 LOS biosyn prt 0550 LOS biosyn prt ÷. lipopolysacharide core biosyn prt (kdtB) Isg locus prt 1 Isg locus prt 1 0651 76 1700 100 83 99 97 0867 0867 Isg locus prt 1 1699 Isg locus prt 2 1698 lsg locus prt 3 1697 lsg locus prt 3 1697 lsg locus prt 4 1696 lsg locus prt 5 98 98 98 99 1695 Isg locus prt 6 1694 1693 lsg locus prt 7 lsg locus prt 8 98 399 57 77 lipopolysaccharide biosyn prt (opsX) rfe prt UDP-3-O-acyl GIcNAc deacetylase 0261 1716 88 (envA) 0915 UDP-3-O-(R-3-hydroxymyristoyl) 91 glucosamine N-acetyltransferase (firA) 1061 UDP-GlcNAc acetyltransferase (ipA) 0873 UDP-GlcNAc epimerase (rffE) 0872 undecaprenyl-P Gal-P Tase (rfbP) 79 79 75 Surface structures 0119 adhesin B precursor (fimA) 0362 adhesin B precursor (fimA) 0330 cell envelope pt (oapA) 0331 opacity assoc pt (oapB) 48 62 100 99 59 91 1174 opacity prt (opa66) 0414 opacity prt (opa66) 1457 opacity prt (opa0) 1460 outer membrane adhesin (yopA) 5823658 0299 pilin biogenesis prt (pilA) 0298 pilin biogenesis prt (pilB) 0297 pilin biogenesis prt (pilC) 0917 protective surface antigen D15 Cellular processes Cell division 0769 cell division ATP-BP (ftsE) 1208 cell division inhibitor (sulA) 1142 cell division prt (ftsA) 78 56 74 88 89 60 1335 cell division prt (ftsH) 1465 cell division prt (ftsH) 1334 cell division prt (ftsH) 1131 cell division prt (ftsL 1141 cell division prt (ftsQ) 1137 cell division prt (ftsQ) 0768 cell division prt (ftsY) 587588877870971 1143 cell division prt (ftsZ 1143 ceil division pr (tts2) 1374 ceil division pr (tmkB) 1353 cytoplasmic axial filament prt (cafA) 0770 ceil division membrane prt (ttsX) 1065 mukB suppressor prt (smbA) 1132 penicillin-BP 3 (ttsl) Cell killing 0301 hemolysin (tlyC) 1658 hemolysin, 21 kD (hly) 1373 killing prt (kicA) 58 72 84 83 55 1372 killing prt suppressor (kicB) 1051 leukotoxin secretion ATP-BP (lktB) Chaperones Cnaperones 0373 heat shock cognate prt 66 (hsc66) 1238 heat shock prt (dnaJ) 1237 heat shock prt 70 (dnaK) 0104 heat shock prt G62.5 (htpG) 0543 heat shock prt groEL (mopA) 0540 heat shock prt groES (mopP) 82 83 88 88 95 95 0542 heat shock prt groES (mopB) Detoxification 0928 catalase (hktE) 1088 superoxide dismutase (sodA) 1002 thiophene and furan oxidation prt (thdF) 100 85 Protein and peptide secretion 1467 colicin V secretion ATP-BP (cvaB) 0016 GTP-binding membrane prt (lepA) 1006 lpp signal peptidase (lspA) 56 91 72 71 1642 peptide transport system ATP-BP (sapF) 0716 preprotein translocase (secE) 62 87 87 73 0798 preprotein translocase (secY) protein-export membrane prt (secD) protein-export membrane prt (secF)

Energy metabolism Aerobic 1163 D-lactate DHase (dld) 1649 D-lactate DHase (dld) 0656 glycerol-3-P DHase (gpsA) 0747 NADH DHase (ndh) Amino acids and amines Armino actos and annines 0534 aspartase (aspA) 0595 carbamate kinase (arcC) 0745 *L*-asparaginase II (ansB) 0288 *L*-Ser deaminase (sdaA) Anaerobic 1047 anaerobic DMSO RDase A (dmsA) 1046 anaerobic DMSO RDase B (dmsB) 1045 anaerobic DMSO RDase C (dmsC) 0644 cytochrome C-type prt (lorC) 0348 denitrification system component (nirT) 0009 formate DHase pathway prt (dhE) 0006 formate DHase (tdnG) 0006 formate DHase-N affector (fdhD) 0006 formate DHase-O, β sub (fdoH) 0007 formate DHase-O, β sub (fdoH) 0007 formate JHase-O, β sub (fdoH) Anaerobic (007 formate DHase-O, β sub (fdoH)
 (1089 formate-dependent nitrite RDase (nrfA)
 (1086 formate-dependent nitrite RDase (nrfB)
 (1087 formate-dependent nitrite RDase (nrfB)
 (1086 formate-dependent nitrite RDase transmembrane prt (nrfD)
 (1083 fumarate RDase (frdC)
 (1083 fumarate RDase (17dC)
 (1083 fumarate RDase (17dC) (frdD) (frdD) 0835 fumarate RDase, flavoprotein sub (frdA) 0834 fumarate RDase, iron-sulfur prt (frdB) 0885 G3PD, sub A (glpA) 0884 G3PD, sub B (glpB) 0883 G3PD, sub C (glpC) 0679 glpE 0618 glpG 1390 hydrogenase isoenzymes formation prt (hvpC)

H#

0601

Other

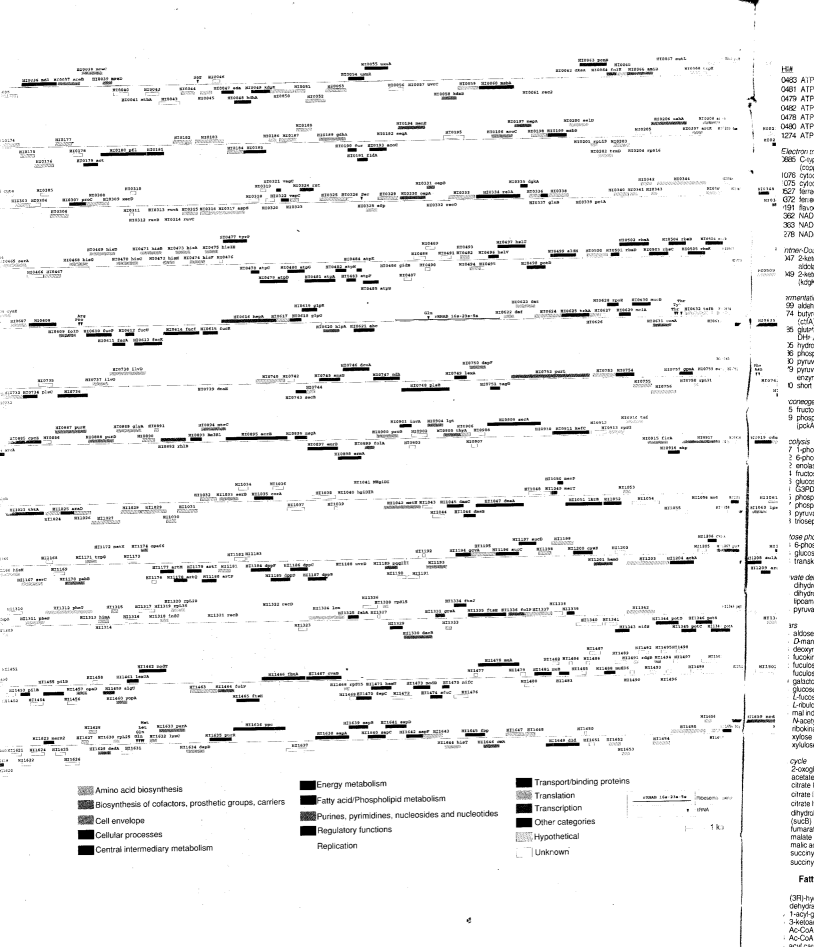
Transformation

- ATP-proton motive force interconversion 0484 ATP Sase C chain (atpE) 0485 ATP Sase F0 α sub (atpB)

%Sim Identification HI# 0445 protein-export membrane prt (secG) 81 10445 protein-export memorane pri (secCa)
10745 protein-export pri (secCl)
10909 preprotein translocase sub (secA)
1015 signal peptidase (lepB)
1016 signal recognition particle prt 54 (ffh)
0713 trigger factor (tig)
(226 type 4 prepiln-like prt specific leader peptidase (hopD) 0483 81 Ř 048 66 91 80 49 0479 0482 0478 0480 нго 1274 нто *Elec* 3885 1008 competence locus E (comE1) 0601 tfoX π 100 0439 transformation prt (comA) 100 1076 1075 1527 1372 191 0439 transformation prt (comB) 0438 transformation prt (comB) 0437 transformation prt (comC) 0436 transformation prt (comD) 100 100 HI034 0435 transformation prt (comE) 0434 transformation prt (comF) 100 RIC 100 362 Central intermediary metabolism 363 Amino sugars 0140 GlcNAc-6-P deacetylase (nagA) 278 0429 Gln amidotransferase (glmS) 0141 glucosamine-6-P deaminase (nagB) 84 intne 88)47 87050 X9 2 Degradation of polysaccharides 1356 amylomaltase (malQ) 62 erme Other 0048 7-o-hydroxysteroid DHase (hdhA) 1204 acetate kinase (ackA) 0949 GABA transaminase (gabT) 0111 glutathione Tase (bphH) 0691 glycerol kinase (glpK) 0594 hippuricase (hipO) 55 99 a 74 b 84 56 57 35 n 89 560 90 90 90 50 0541 urease (ureA) 0539 urease α sub (urea amidohydrolase) 76 82 Phe Apri TT (ureC) 0537 urease accessory prt (UreF) 0538 urease prt (ureG) 0536 urease prt (ureG) 0535 urease prt (ureH) 0540 urease sub B (ureB) 55 57 87 HI07 10 si cone 54 77 5 fn 9 pt (p Phosphorus compounds 0695 exopolyphosphatase (ppx) 0124 inorganic PPase (ppa) 0645 lysophospholipase L2 (pldB) 77 colys 7 1-i 50 53 2 6-j 2 en 1 fru Polyamine biosynthesis 0099 nucleotide-BP (potG) 0591 omithine DCase (speF) 67 80 glu G3 NT10 5 ph Polysaccharides - (cytoplasmic) 1357 1,4-α-glucan branching enzyme (glgB) 1060 1 80 3 pyr 3 trio 1361 α-glucan phosphorylase (glgP) 79 1359 ADP-glucose Sase (glgC) 1358 glycogen operon prt (glgX) 1360 glycogen Sase (glgA) 74 è9 tose 71 6-p gluo trar Sulfur metabolism 0805 arylsulfatase regulatory prt (aslB) 1371 desulfoviridin γ sub (dsvC) 67 871209 58 vate 0559 sulfite synthesis pathway prt (cysQ) 56 dihy dihy lipo pyru 48 78 81 75 NT: ars aldo D-m deox fuco 89 ar15 fucui 88 81 fucul galad 83 ğlucc L-tuc L-ribu 86 mali JI1659 ni 85 N-aci 68537272 riboki xyluk 79 71 72 *cycle* 2-oxo aceta citrate 86 75 67 81 citrate citrate dihvdr (sucB 68 fumari malate 72 77 malic a succin succin 87 85 Fat 83 ŝ (3R)-h dehydr 63 66 1-acyl-3-ketoa õ Ac-CoA acvl car 82 78 acvl-Co

β-ketoa

RT0021 RT0022 citP HI0024 citD HI0026 110A HI0028 HI0023 data HI0021 HI0028 citP HI0028 ci NX 0017 \$7773 HI0019 HI0020 HICOOS HIGOOG ETTTS HIG177 537.37 HID178 HI0147 HI0139 E10154 acpF BI0156 febD HI0157 fabR HI0159 Asp Asp HI0138 rnb W MRN HIG161 gor HIG16: HI0152 HIG143 H10176 H10 012 cute HI0275 HI0280 udp #10284 HI028 HI0281 HI0282 HI0283 m 810271 HI0428 dabB Lo 1443 rock HI0445 se BI0427 nhaB BI0455 hol9 HI0457 pa omD HI0439 Ala Ile VV rRNAA 168-238-54 HI0593 Origin HI0503 cpds H1058 Lindessenations 10588 810589 HI0602 hemY Annalder HI0603 hem Stylinger Styl HI0582 gidA Ala Ile Yy rRNAC 168-238-58 HI0715 clp3 X H10723 10722 pep0 10714 clpp H10716 seck H10708 sel HI0876 nd) HI0875 pepA нтов57 ытов58 пов59 сърв а нтов57 ытов58 сърв нтов55 клов55 рода ШД нтов55 клов55 рода нтов55 клов55 клов55 клов55 клов55 клов55 нтов55 клов55 клов55 клов55 нтов55 клов55 клов55 клов55 нтов55 клов55 клов55 нтов55 клов55 клов55 нтов55 клов55 нтов55 клов55 нтов55 клов55 клов55 нтов55 клов55 нтов55 клов55 нтов55 клов55 нтов55 клов55 нтов55 клов55 нтов55 н H10879 2pL27 HILOOA HI1004 E11003 // H11019 V///////// HI1021 HI1021 bic HT1013 HT1014 HI0990 igal HIL164 cmpA 0(1165 HI1153 pmbA HI1154 gltP ddlB HI1142 ftsk HI1144 00 MILLSS LINE HILLSO VISA HILLS? SI1133 murE HI1135 mray BI1139 BurC HI1166 hiss 811169 HI1146 BI1148 EI1150 HI1155 HI1156 cydC HI1163 did HI1167 serC HI1157 cydD HI1159 trxA B11149 HT1304 mugB HT1307 ml.4 HT1305 HT130 H11289 ZUUUUUUU N1280 H11290 tyrA VII280 H11290 H11292 H11294 H11296 N1291 H11292 H11292 H11293 H1 HI1284 infB BI1280 HI1277 RED HI1279 siab HI1281 HI1282 Manganan Wei CHARGO 773 Y V/7778 HI1283 nush H11274 at HZ1301 HT1302 ABAB HI1285 hedR EI1286 HI1449 chiN HI1451 NI1420 NI1420AIL62 Eau NI1426 NI1419 NI1420AIL623 Lou NI1428 faz NI1427 NI1421 NI1426 rot NI1428 puzh 9 NI1421 NI1424 rot NI1428 puzh RII44 ener MINUN RII44 chir MARKAN HI144 ener HI144 ene HI1431 trpB HI1438 ispA HI1437 HI1439 HI1430 HI1432 trph HI HI1430 HI1432 trph HI HI1433 usg HT1434 877777 HI141: HI1435 HI1436 HI1406 HI1407 traN HI1410 HI1411 HI1413 HI1416 HI1418 HI1429 DURM HI1414 HI1409 HI1415 HI1616 purK Manandifish HI1606 cca RI1608 92.20256-2025 20202020 EI1605 HI1607 HI1609 press 2020202 402020 402020 HI1588 purU BI1613 ribC 1625 HI1589 325369 8 bio HI1621 HI16 H H 1619 HI1622 HI1620 HI1610 tyrs R11612 H11601 ST1584 11vH . HI1526 HI1600 H11602 EI1585 ilvI 311727 arg Ile Ala VV HT1735 p HI1740 1 HI1732 HI1733 HI1736 RI1738 BI173



acyl car acyl-Co. β-ketoa

 Bito077
 Bit0077
 Bit0070
 < Cys Lys Gy En TT math HELOOF ther HELOOF ther HELOOF ther HELOOF the HELOOF T НІЛООО 2007/02 HILOOOS HILOOOS HILOOOS HILOOOS IDO HILOOOS GerC2 2 pecd HI0103 dept H10104 bi cd HI0103 HI 107 HI H10100 HI0103 dept H10104 bi cd HI0107 HI0107 HI H01040 HI0103 dept H10104 bi cd HI0107 HI010 an S(210 HI0222 riba NAMES 10211 pgpB HI0213 HI0214 priC HI0215 hadm HI0216 hadg HI0218 prrD HI-0221 guad HI-0222 guad HI-0224 LTD HI HI-0223 guad HI-0223 HI0219 HI0220 arch HI0229 pmp HI0230 F.0231 deat HI0237 HI6233 HI 2 HIG356 tena HIG355 INSE IL 1722/73 INSE ILOST unti 1722/73 INSE ILOST UNTI HIG355 ILOSE HIG361 funk HIG365 6277/2798 ILOSE HIG361 funk HIG365 HIG361 funk HI0234 HI0236 a HIO365 HIO369 HIO369 HIO365 RIO367 HIO369 POC HIO370 Dunning running r HI0387 ding HI0389 F(0390 rnd -nd # B20391 H10392 N16 H10393 H10393 826394 ; 10530 10503 HIOSII teba HIOSI3 hinoIIM HIOSI4 Fpc: 1073 HIOSI2 HincII HI0515 FDOB HI0516 FDL1 HI0518 480D HIG518 HI0520 HI0515 FDOB HI0516 FDL1 HI0518 480D HIG518 HI0520 HI0520 HI0522 HI0524 FDA HI0517 FDL1 HI0517 FDL1 HI0521 FDL1 HI0522 HI0521 HI0521 HI0521 HI0521 HI0522 FDA HIO543 9 urec HIO541 urea HI HIO540 ureb Bx0531 rp821 HI0527 fdx HI0533 HI0526 BI0528 tyrP HI0530 gcp HI0535 ureHI0537 UreF HI0539 HI0536 ureG H10538 ureE HIG632 glad gar HIG632 glad HIG64 grad HIG64 grad HIG64 grad HIG64 grad HIG650 HIG653 HIG655 HIG657 HIG655 HIGST ETDER HIGST E10668 HI0675 H10671 HI0673 10000. HIG672 HIG674 1000000000 - 10000 RIO810 HIO811 argH 10812 galU HIG //////// RENDERRENT IN AND CETA HIO814 als: RIO813 CETA HIO814 als: Bioliging of Angel also arg HIGSIS USDA ET0809 pcka HU922 - LU92 HU922 - LU9 Setting Set HI0923 holA II 9928 hktE glyg HI 0939 HI 0930 HI 0932 eno HI 0933 USUMU MI USU 10931 USU 1000 HI 0931 J HI0935 HI0941 HI0945 httN H10945 httN H10946 dut H10956 HI0940 HI0942 rec HI0944 tibo HI0948 HI0946 tp H10948 HI0957 crp HI0946 tp H10947 vapC H10950 rpl.1) W/W/W HI0957 crp HI0946 tp H10947 vapC H10950 rpl.1) H10958 HI0957 kl/2 HI0946 tp H10947 vapC H10950 rpl.3 18 i cdsA BI0920 #10935 HI0934 HI0936 HI0952 rade HI1065 ambh 9 2:51 lpas H11065 ambh 2 2:51 lpas H11066 nrfD H11066 nrfD H11066 nrfD H11075 H11072 H11075 cydb 2 1ew H11062 fabz H11067 nrfC H11069 nrfh H11071 H11073 H11075 cydb H11075 cydb H11075 cydb H11075 cydb H11075 cydb H11075 tabz H11075 tabz H11075 H110 HI1220 FRI1221 HI1225 H rila Hil211 yeb Hil212 prfs Signature States BI1219 go FII363 pntB HI1365 topA HI367 thrs HI1366 glgA HI365 topA HI365 topA HI367 thrs HI366 H oga HI1353 cafA RI1354 clos HI HI1351 BI1352 putP HI1359 glgC HI1350 cda Millionan T1349 HI13 BI1357 glgB HI1373 kica HI1372 kice HI1504 EUI HI HI1504 EUI HI1 HI502 HZ1503 MUG HX1505 NII595 NII557 NII510 NII512 NII514 NII516 N NII567 NII560 NII511 NII515 MM NII517 NII515 NII569 NII551 NII515 MM NII517 NII515 HI1520 HT1522 HI1520 PART HI1535 HI1537 LICA HI1539 LICC HI1523 HI1524 HI1527 IDDB HI1529 PARC HI1530 QLG HI1532 GTR HI1534 HI1536 HI1537 HICA HI1539 LICD HI1542 HI1542 HI1525 mode HI1529 TO HI1529 TO HI1529 TO HI1529 TO HI1529 TO HI1529 HI1539 LICD HI1542 HI1539 HI1542 HI1539 HI1541 HI1539 HI1549 HI1539 HI1539 HI1549 HI1549 HI1539 HI1549 HI1539 HI1549 HI erid. Nileso nrdl Nileso nrdl Nileso nrdl Nileso nrdl Nileso nrieso United nileso nrieso Nileso nrieso Nileso nrieso Nileso nrdl Nileso nrieso Nileso nrieso Nileso nrdl Nileso nrieso Nileso nrdl Nileso nrd Nil 871544 HI1672 HI1673 De RI2673 most HI1675 HI1672 filo HI1672 filo HI1672 nosc HI1684 HI1686 rnfE H1169 HI1682 sohB HI1683 HI1685 tolA H11687 H11688 H11689 hth Shaddbala UTU23 HI1694 HI1691 modC HI1693 BI1695 HI1692 modC HI1693 BI1695 RI1697 HI16

B:0133 ded [50,000 nt HIOIIS E Modeland Het MIOII7 HIOIIS Chin HIOIIS HIOIIS Chin HIOIIS HIOIIS HIOIISHIMAL H10119 fim2 Leu Gly Lys HI0123 DGSA HI0125 MIGIO7 HI0108 HI0110 serS RI0111 bpbH Glu T Pro V BT0126 B10127 810120 HI0112 TRNAE 58-238-16 HX0114 NZ0131 HI0121 300,000 nt (10267 narQ HI0255 dapa HI0257 Ser Manager III gi0250 sab H10265 H10237 HI0230 HI0231 нт0242 HT0241 HT0244 tgt HT0245 queA 7/72 U2 жылынын балынынын 40 secD HT0243 HT0245 HI0251 t BI0247 igal BI026 810235 HI0240 secD 450,000 nt BI0416 BIG418 RI0425 pask HI0427 nhaB КІСЭЭЭ ВІСЭЭЭ КІСЭЭЭ КІСЭЭЭ КІСЭЭЭ КІСЭЭЭ КІСЭЭЭ КІСЭЭЭ HI0414 op H10423 BI0402 dat1 HI0419 prtC HI0420 HI0415 820417 BI0409 HI0426 fadR BI0421 HI0422 armB HIC390 rnd 20389 HI0406 accA HI0408 RIG411 hfg BIO41: H10404 HI0400 HI0391 HI0407 12 810405 600,000 nt 0///: HI0572 HI0574 HI0575 HI0577 Things Wings 1 HI0552 HIO543 kpga ELO553 kpga HIO543 kpga ELO553 kpga elogospa H10562 H10564 HI0573 slyD HI0580 EpS7 HI0553 gnd HI0554 BI0556 devB HI0559 cysQ HI0560 2/11/1 BI0579 fush HI0558 HI0538 750,000 nt H10704 н10689 hpd HI0691 HI0701 HI0702 BI0706 slpb UC HIG675 PODD HIG676 CPLA ANTHONYSTER HIG671 HIG676 X007 HIG676 CPLA HIG671 HIG676 X007 HIG679 GJAR HIG681 11VX WWW. HIG674 HIG677 HIG660 X007 HI0693 hel HI0694 Sectorial 2011 HIG669 mioC glpT HI0687 HI0688 HI0668 HI0670 HI0703 lppB HI:0700 N10692 BIO695 DOX HIO696 mmmmu BI0697 HI0699 # PD 10672 HI0674 **HI0666** 900,000 nt HT0852 H10843 H10847 H10845 H10846 H10845 H10846 trmA m H10845 H10846 trmA m H10844 mob H8886 H10941 HI0827 HI0829 RIOSS 4ppA HI0825 HI0826 HI0828 kch HI0 HIO832 fraD HIO835 fraA HIO833 fraC HIO834 fraB HI0815 USDA HI0817 HI0618 HIOSIC HIOSII argH HI0812 galU EI0816 pepP HI0813 CSTA HI0814 alaS L050.00 nt H10968 mebB H10970 arcQ H10972 accC <u>ысыклум</u> H10965 H10967 H10969 manC H10971 accB H10973 Z) _______ HI0986 HI0987 Leuk H10988 HI0989 LeuD H10954 dos H10956 MMR 10055 H10955 ddp H10955 H10957 crp H10953 ddp 10052 1165 H10952 1165 H10958 H10958 H10959 H10951 H10952 1165 H10952 1165 H10952 1165 H10952 H1095 HI0981 ampB HI0983 lauA HI0985 dpr/ H10990 igal HI0956 HI0982 pfkA HI0984 HI0950 rpL33 abr HI0951 rpL28 HI0960 1.200,00 nt L 200,00 ml HI126 M1130 HI132 fter HI137 ceta HI120 HI133 fter HI137 ceta HI120 HI133 fter HI137 ceta HI130 HI133 fter HI133 mut2 HI1096 HI1098 H11125 HI1119 lap BI1124 oppA 806 HI1102 CYSZ HOMMAN HI1120 oppy HI1122 oppC HI1126 RT1117 HT1118 HI1100 Lig HI1101 HI1107 HI1115 #11269 #11270 1.350.00 nt HI1268 HI1273 HI1258 mfd HI1254 HI1256 H11251 vapA H11254 H11256 H11250 H11252 H11255 H11255 H11257 HI1267 HI1272 fepC HI1263 met2 Statestade BI1264 gyrA HI1241 HI1244 HI1259 htra EI1248 HE1266 HI1239 proA RI1240 BI1242 811249 1,500,000 nt HI1387 trof HI1389 trp HII390 hypC BII391 val9 Sigginaticsimit BII HI1398 fumC HI1:99 sgA HI1385 HI1388 trpD NI1400 HI1402 HI1404 als HII393 bindIIIR HII396 MANNANCE HII392 bindIIIM HII394 HII397 hold bhoR HI1380 pstB HI1382 pate HI1401 DYTD HI1403 HI1369 HI1370 mop1 HI1371 davo H11395 HI1574 dnaß BI1576 pgi B11578 lgt 1.650.000 nt H11583 Arg9 pyka Bil575 alr HI1565 tbp1 HI1567 iroA ica HI1539 licC HI1579 1pp HI1546 G HII549 GAVA HII551 DIOC HII553 DIOF HII555 HII557 MAGA HII555 DAMED HII551 DIOFA HII556 MANDALE MANDALE MANAGEMENT HII560 HII552 HII552 HII556 HII556 HII556 HII566 HII666 HI HI1568 muG HI15 HI1568 muG HI157 HI1569 HI1571 HI1570 H1154 811572 538 licB HT1540 licD 143 HI1545 2010/07/00/00 HI1544 BI1547 HI1581 HI1592 HI1541 sppA HI1543 HI1562 HI1564 1,800,000 nt HI1727 argG MONNE BI1726 purC #11725 pon8 DA RIIJ15 GUY MARKAR RIIJ05 BIS HIIJ06 beet RIIJ07 bass HIIJ09 BIJJ11 orr HIV13 ptes RIIJ14 BIJJ56 HIJ37 HIJ19 JAD HIJ706 bass HIIJ09 BIJJ11 orr HIV13 ptes HIJ706 bass RIIJ00 HIJ712 ptes HI1720 HI1721 HI1723 HI1'24 ни1705 рерА данияниян HI1701 HI1722 map HI1702 metE HI1703 H11695 HI1697 HI1698 HI1700 HI1691 modC HI1693 دا EI1704 *سیسیس* HT1699

	Hit Identification 0483 ATP Sase F0 β sub (atpF) 0491 ATP Sase F1 α sub (atpA) 0479 ATP Sase F1 β sub (atpD) 0482 ATP Sase F1 δ sub (atpH) 0478 ATP Sase F1 ε sub (atpC) 0470 ATP Sase F1 ε sub (atpC) 0478 ATP Sase F1 ε sub (atpC) 0479 ATP Sase Sub 3 region ptt (atpC)	% <u>Si</u> 9 9 7 7 8 5
ETO ETO ETO ETO	Electron transport 2685 C-type cytochrome biogenesis prt (copper tolerance) (cyc2) 1076 cytochrome oxidase d sub I (cydA) 1075 cytochrome oxidase d sub II (cydB) 527 ferredoxin (fdx) 1372 ferredoxin (fdx) 1391 flavodoxin (fldA) 362 NAD(P) transhydrogenase sub α (pntA) 363 NAD(P) transhydrogenase sub β (pntB)	6 8 7 8 8 8 8 8 8 8
1:050 332204	 278 NAD(P)H-flavin oxidoreductase <i>intner-Doudoroff</i> 147 2-keto-3-deoxy-6-phosphogluconate aldolase (eda) 149 2-keto-3-deoxy-D-gluconate kinase (kdgK) 	5: 6: 6:
31063) 31063) 41 41 41 41 41 41 41 41 41 41 41 41 41	 #mentation ge aldehyde DHase (aldH) dutyrate-acetoacetate CoA-Tase sub A (ctfA) glutathione-dependent formaldehyde DHase (gd-faldH) hydrogenase gene region (hypE) phosphoenolpyruvate carboxylase (ppc) pyruvate formate-lyase activating enzyme (act) short chain alcohol DHase 	62 75 78 48 99 85 69
i	<i>coneogenesis</i> 5 fructose-1,6-bisphosphatase (fbp) 9 phosphoenolpyruvate carboxykinase (pckA)	84 83
3119 ct 3110 35 2160 1 2263	colysis 7 1-phosphofructokinase (fruK) 2 6-phosphofructokinase (pfkA) 2 enolase (eno) 4 fructose-bisphosphate aldolase (fba) 5 glucose-6-P isomerase (pgi) 1 G3PD (gap) 5 phosphoglycerate kinase (pgk) 7 phosphoglyceromutase (gpmA) 3 ptruvate kinase type II (pykA) 3 trosephosphate isomerase (tpiA)	74 84 79 86 89 99 175 87 81
SH su	tose phosphate pathway 1 6-phosphogluconate DHase (gnd) 1 glucose-6-P 1-DHase (G6PD) 1 transketolase 1 (tktA)	71 65 88
51209 -4 1990	 vate dehydrogenase dihydrolipoamide acetyltransferase (aceF) dihydrolipoamide acetyltransferase (acoC) lipoamide DHase (lpdA) pyruvate DHase (aceE) 	82 49 92 84
211 2 2 4 stil 2 2 59 p	ars aldose 1-epimerase precursor (mro) D-mannonate hydrolase (uxuA) deoxyribose aldolase (deoC) fucclinase (fucK) fuculose-1-P aldolase (fucA) fuculose-1-P aldolase (fucA) galactokinase (galK) glucose kinase (glK) L-fucose isomerase (fucl) L-fucose isomerase (fucl) M-acetylneuraminate lyase (nanA) ribokinase (hosK) xylose isomerase (xylA) xylose kinase	5368633383362367555
	cycle 2-oxoglutarate DHase (sucA) acetate:SH-citrate lyase ligase (AMP) citrate lyase α chain (citF) citrate lyase β chain (citE) citrate lyase γ chain (citD) dhydrolipoamide succinyltransferase (sucB)	81 68 86 81 72 84
The second	fumarate hydratase (fumC) malate DHase (mdh) malic acid enzyme succinyl-CoA Sase α sub (sucD) succinyl-CoA Sase β sub (sucC)	74 85 89 80
	Fatty acid and phospholipid metabolism	
.	(3R)-hydroxymyristol acyl carrier prt dehydrase (fabZ) , 1-acyl-glycerol-3-P acyltransferase (plsC) 3-ketoacyl-acyl carrier prt RDase (fabG) Ac-CoA acetyltransferase (fadA) ; Ac-CoA carboxylase (accA) acyl carrier prt (acpP) ; acyl-CoA thioesterase II (tesB) t R-ketoacyl-ACP Sase I (fabB)	85 78 80 89 91 73 84

%Sim

76

75

80

85

una glim

HI0347

t 10507 NIOSO8 WINE

Y H10761

HI0918 181

1348 pept - 1348 pept

BI154 8

58 hly

on 1

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77

78

otj

3 acyl-CoA thioesterase II (tesB)	
β-ketoacyl-ACP Sase I (fabB)	

in	n HI# Identification	%Sin	<u>а н</u>	#	
Æ	9 0157 β-ketoacyl-acyl carrier prt Sase III (fat) H) 8	03	34 A	
æ æ	5 09/1 biotin carboxyl carrier prt (accB)	8 9'	3		re ai
72	0919 CDP-diglyceride Sase (cdsA)	67	7 08	13 c	a
76	denyoratase (rabA)	92	12	00 ç	yc ys
ю 50		72 68)) €0 f€	cy eri
	0748 glycerol-3-P acyltransferase (plsB)	76	5 14	53 fi	m pill
98		53 se 82		55 fi	m
2	(fabD) 0211 phosphatidylglycerophosphate	60	126	(1 50 fc	oill oly
8 7	phosphatase B (pgpB)	83		e	хŗ
14	(pgsA)			re	эġ
17 14	0160 phosphatidylserine DCase proenzyme (psd)	76	; 082 075		ala luc
8	0425 phosphatidylserine Sase (pssA)	71	119	14 G	ily cti
5	0689 prt D (hpd) 1734 short chain alcohol DHase homolog	99 85	100	19 gl	lyc
з	(envM) 1433 USG-1 prt (usg)	54	061		yc Tl
			087 057	7 G	T
4	Purines, pyrimidines, nucleosid and nucleotides	es,		(Ö	İΧ
	2'-Deoxyribonucleotide metabolism	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	061	9 la	tu cz
2	0075 anaerobic ribonucleoside-triphosphate RDase (nrdD)	88	022 159	4 Le	eu
5	0133 deoxycytidine triphosphate deaminase (dcd)	87	074	9 Le	эx.
3	0954 deoxyuridinetriphosphatase (dut)	91	146 161		
3	1532 glutaredoxin (grx) 1660 ribonucleoside diphosphate RDase 82 su 1659 ribonucleoside-diphosphate RDase 1	80 ab 93	029	4 m	et
3	1659 ribonucleoside-diphosphate RDase 1 α chain (nrdA)	92	019	3 m 9 m	sb
5	1158 thioredoxin RDase (trxB)	86	076 071	3 na 3 ne	
)	0905 thymidylate Sase (thyA)	55	062	Э пе	ga
	Nucleotide and nucleoside interconversions		026	7 nit 3 nit	
Ļ	1077 CTP Sase (pyrG) 1299 dGTP triphosphohydrolase (dgt)	90 58	033	(na	arl
ł	0132 uridine kinase (udk)	85	174	l pe	nt
	Purine ribonucleotide biosynthesis		1378	py 3 ph	
	1616 5'-phosphoribosyl-5-amino-4-imidazole carboxylase II (purK)	72	1379) ph	os
	1429 5'-phosphoribosyl-5-aminoimidazole Sase	87	1635	reg pu	
	(purM) 1743 5-guanylate kinase (gmk)	82	0163	(pu	١rF
	U349 adenviate kinase (adk)	100 88	0506	i rbs	s n
	0639 adenylosuccinate lyase (purB) 1633 adenylosuccinate Sase (purA)	87	0563		jul Sre
	1207 amidóPRTase (purF) 0752 formylglycineamide ribonucleotide Sase	84 82	0269	I RN	IA
	(purL)		0533		
	(purL) 1588 formyltetrahydrofolate hydrolase (purU) 0222 GMP Sase (guaA) 0221 imeanhath Dillose (guaP)	85 88	1707 1440		
	0221 inosine-5'-monophosphate DHase (guaB) 0876 nucleoside diphosphate kinase (ndk)	81 74	1441	stri	'ng
	0888 phosphoribosylamine-Gly ligase (purD)	85	1739	trai (me	
	0887 phosphoribosylaminoimidazõle carboxamide formyltransferase (purH)	87	0358	trar	٦S
	1615 phosphoribosylaminoimidazole	97	0681 1708	trar trar	
	carboxylase catalytic sub (purE) 1428 phosphoribosylglycinamide	71	0410 0830	trar Trp	
	formyltransferase (purN) 1609 phosphoribosylpyrophosphate Sase	91	0054	uxu	10
	(prsA) 1726 SAICAR Sase (purC)		1106	xylo)S
	1726 SAICAR Sase (purc)	55	Dog	adat	i.
	Pyrimidine ribonucleotide biosynthesis	77	<i>Degr</i> 1689	епс	lo
	1401 dihydroorotate DHase (pyrD) 0272 orotate PRTase (pyrE)	84	0249 1247	exc exc	
	1225 orotidine 5'-monophosphate DCase 1224 orotidine-5'-monophosphate DCase (pyrF)	88 79	0057	exc	in
	0459 uracil PRTase (pyrR)	74	1377 1321	exo	de
	Salvage of nucleosides and nucleotides		0942 1322	exo	de
	Salvage of nucleosides and nucleotides 0583 2',3'-cyclic nucleotide 2'- phosphodiesterase (cpdB)	78	0041	exo	nι
	1230 adenine PRTase (apt)	83	0397 1214	exo sino	nı Ile
	0551 adenosine tetraphosphatase (apaH) 1350 cytidine deaminase (cda)	73 63		(rec	
	1646 cýtidylate kinase (cmk) 1219 cytidylate kinase (cmk)	77 79	DNA	repli	ica
	0518 purine-nucleoside phosphorylase (deoD)	90	<i>reco</i> 0759	mbir A/G	na -e
	1277 putative ATPase (mrp) 0529 thymidine kinase (tdk)	79 82	1226	chro	n
	1228 uracil PRTase (upp)	94	0993 0314	cnrc	
	0280 uridine phosphorylase (udp) 0674 xanthine-guanine PRTase	85 88		(ruv	C)
	0692 xanthine-guanine PRTase	88	1264	DN/	٩d
	Sugar-nucleotide biosynthesis and conversions		0567	DN/ DN/	٩ç
	0206 5'-nucleotidase (ushA) 1279 CMP-NeuNAc Sase (siaB)	55 64	1188	DN/	۱
	0820 Gal-1-P uridylyltransferase (galT) 1	100 86	0654	DN/ DN/	۱3
	0812 Glc-P uridylyltransferase (galU) 0351 UDP-Glc 4-epimerase (galE)	99	0403	DN/ DN/	٩r
	0642 UDP-GICNAC pyrophosphorylase (glmU)	83	0707	DN/	٩r
	Regulatory functions	m	0856 0992	DN/	۱ŗ
	0604 adenylate cyclase (cyaA) 1 0884 aerobic respiration control prt (arcA)	00 88	0923		۱ŗ
	0220 aerobic respiration control sensor prt	70		DN/ DN/	
	(arcB) 1052 araC-like transcription regulator	48	0739	DNA	۱ŗ
	1209 Arg repressor prt (argR) 0236 arsC prt (arsC)	81 57	1397		

78 88 88 91 73 84 1209 0236 arsC prt (arsC) 0462 ATP-dependent proteinase (Ion)

Identification <u>%Sin</u> TP:GTP 3'-pyrophosphotransferase iA) arbon starvation prt (cstA) arbon storage regulator (csrA) rclic AMP receptor (crp) sr regulon transcriptional activator ysB) rric uptake regulation prt (fur) 1brial transcription regulation repressor IB) brial transcription regulation repressor 73 IB) ylpolyglutamate-dihydrofolate Sase pression regulator (accD) marate (and nitrate) reduction gulatory prt (fnr) lactose operon repressor (galS) œ cockinase explaint repressor (gais) cockinase regulator y cleavage system transcriptional tivator (gcvA) vcerol-3-P regulon repressor (glpR) rcerol-3-P regulon repressor (glpR) IP-BP (chg) IP-BP (obg) 6€ 77 87 71 86 IP-bP (009) drogen peroxide-inducible activator cyR) ucose operon activator (fucR) Z expression regulator (icc) u responsive regulatory prt (Irp) I responsive regulatory prt (irp) I responsive regulatory prt (irp) Ar repressor (lexA) oligosaccharide prt (lex2A) Itose regulatory prt sts1 (stsA) IF aporepressor (metJ) lybdenum transport system (modD) bB dB dAB transcriptional regulator (nadR) gative regulator of translation (relB) gative rpo regulator (mclA) ate sensor prt (narQ) 88 79 ate, nitrite response regulator prt rP) ogen regulatory prt P-II (glnB) ita-P guanosine-3'-77 ra-r guanosine-3- 77 phosphohydrolase (spoT) sphate regulon sensor prt (phoR) 67 sphate regulon transcriptional 72 ulatory prt (phoB) ne nucleotide synthesis repressor prt 74 R) rFl) ative murein gene regulator (bolA) repressor (rbsR) ulatory prt (asnC) ressor for cytochrome P450 (Bm3R1) A polymerase sigma-32 factor (rpoH) A polymerase sigma-70 factor (rpoE) sor prt for basR (basS) neent starvation prt (sspB) 87 88 56 81 son pri toi bash (bass) igent starvation prt (sspB) igent starvation prt A (sspA) s-activator of metE and metH tR) (n) scriptional activator (tenA) scriptional activator prt (livY) scriptional regulatory prt (basR) scriptional regulatory prt (tyrR) repressor (trpR) operon regulator (uxuR) se operon regluatory prt (xyIR) Replication Hepiication on of DNA onuclease III (nth) nuclease ABC sub A (uvrA) nuclease ABC sub B (uvrB) nuclease ABC sub C (uvrC) on of DNA ponuclease III (nth) 92 nuclease ABC sub A (uvrA) 91 nuclease ABC sub C (uvrC) 80 leoxyribonuclease I (sbcB) 75 leoxyribonuclease V (recC) 61 leoxyribonuclease V (recC) 59 nuclease III (sthA) 84 nuclease VII, large sub (xseA) 74 e-stranded DNA-specific exonuclease 77 n) ation, restriction, modification, auor, restrictori, indonication, ation, and repair specific adenine glycosylase (mutY) 75 nosomal replication initiator (dnaA) 75 nosomal replication initiator (dnaA) 80 sover junction endodeoxyribonuclease 88 20 adenine methylase (dam) gyrase, sub A (gyrA) gyrase, sub B (gyrB) helicase (recQ) helicase II (uvrD) licase (II) 86 98 polymerase III δ sub (holA) polymerase III δ ' sub (holB) polymerase III ɛ sub (dnaQ) polymerase III α chain (dnaE) 1397 DNA polymerase III χ sub (holC) 0011 DNA polymerase III psi sub (holC) 0532 DNA primase (dnaG) 1354 Gin-tHNA Sase (ginS) 0274 Glu-tRNA Sase (gitX) 0927 Gly-tRNA Sase α chain (glyQ) 74

1		<u>6Sim</u>
0 4	1740 DNA recombinase (recG) 0070 DNA repair prt (recN) 0657 DNA topoisomerase I (topA)	80 67 55
1	0566 dod 0062 dosage-dependent dnaK suppressor prt	93
ð	(dksA) 0946 formamidopyrimidine-DNA glycosylase	75
5	(fpg) 0582 glucose-inhibited division prt (gidA)	87
3	0486 glucose-inhibited division prt (gidB) 0980 Hin recombinational enhancer BP (fis)	78 93
3	0512 HincII endonuclease (HincII) 1392 HindIII modification MTase (hindIIIM)	98 99
)	1393 HindIII restriction endonuclease (hindIII 0313 Holliday junction DNA helicase (ruvA)	80
)	0312 Holliday junction DNA helicase (ruvB) 0676 integrase-recombinase prt (xerC)	90 74
; ;	0309 integrase-recombinase prt (xerD) 1313 integration host factor α sub (himA)	88 88
)	1221 integration host factor β sub (IHF-β) (himD)	77
	0402 methylated-DNAprt-Cys MTase (dat1) 0669 mioC	62 72
,	1041 modification methylase HgiDI (MHgiDI) 0513 modification methylase HinclI (hinclIM)	70 99
	0910 mutator mutT 0192 negative modulator of initiation of	72 72
	replication (seqA) 0546 primosomal prt n precursor (priB)	100
	0339 primosomal prt replication factor (priA) 0387 probable ATP-dependent helicase (dinG)	
	0991 DNA, ATP-BP (recF) 0332 DNA repair prt (recO)	76 77
	0600 recombinase (recA) 0061 recombination prt (rec2)	100 100
	0443 recR prt (recR) 0599 regulatory prt (recX)	88 50
	0649 rep helicase (rep) 1229 replication prt (dnaX)	83 70
	1574 replicative DNA helicase (dnaB) 1040 restriction enzyme (hgiDIR)	83 64
	1172 SAM Sase 2 (metX) 1424 shufflon-specific DNA recombinase (rci)	92 56
	0250 single-stranded DNA BP (ssb) 1572 site-specific recombinase (rcb)	98 57
	1365 topoisomerase I (topA) 0444 topoisomerase III (topB)	84 79
	1529 topoisomerase IV sub A (parC) 1528 topoisomerase IV sub B (parE)	85 89
	1258 transcription-repair coupling factor (mfd) 0216 type I restriction enzyme ECOK1	83 59
	specificity prt (hsdS) 1287 type I restriction enzyme ECOR124/31	54
	M (hsdM) 0215 type I restriction enzyme ECOR124/31	89
	M (hsdM) 1285 type I restriction enzyme ECOP124/3 R	53
	(hsdR) 1056 type III restriction-modification ECOP15	56
	enzyme (mod) 0018 uracil DNA glycosylase (ung)	ж 80
	Transcription	00
	Degradation of RNA 0218 anticodon nuclease masking-agent (prrD)	86
	1733 exoribonuclease II	68
	0413 ribonuclease E (rne)	65 72
	0138 ribonuclease H (mh) 1059 ribonuclease HII	76 83
	0014 ribonuclease III (rnc) 0273 ribonuclease PH (rph)	80 88
	0999 RNase P (mpA)	81 81
	RNA synthesis, modification, and DNA	
1	transcription 2616 ATP-dependent helicase (hepA)	74
¢	2231 ATP-dependent RNA helicase (deaD) 2892 ATP-dependent RNA helicase (mB)	79 84
	2422 ATP-dependent RNA helicase (srmB) 2802 DNA-directed RNA polymerase α chain	61 97
((rpoA) 515 DNA-directed RNA polymerase () chain	92
0	(rpoB) 514 DNA-directed RNA polymerase β' chain	91
1	(rpoC) 304 N utilization substance prt B (nusB)	71
Ç	063 plasmid copy number control prt (pcnB) 1229 polynucleotide phosphorylase (pnp)	73 87
1	742 RNA polymerase omega sub (rpoZ) 459 sigma factor (algU)	76 49
C	717 transcription antitermination prt (nusG) 331 transcription elongation factor (greA)	70 84 90
C	569 transcription elongation factor (greB) 283 transcription factor (nusA)	30 79 84
	225 transcription termination factor rho (rho)	95
,	Translation Amino acyl tRNA synthetases and tRNA	
	modification 1814 Ala-tRNA Sase (alaS)	83
1	583 Arg-tRNA Sase (argS)	84 91
C	817 Asp-tRNA Sase (aspS) 708 Cvs-tRNA selenium Tase (selA)	86 76
C	078 Cys-tRNA Sase (cysS)	87 87

0924 Gly-tRNA Sase β chain (glyS)

95

Identification HI#
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 Identification

 0369
 His-tRNA Sase (hisS)

 0962
 IIe-tRNA Sase (lieS)

 0962
 IIe-tRNA Sase (lieS)

 1211
 Lys-tRNA Sase (lieUS)

 1211
 Lys-tRNA Sase (lieUS)

 0836
 Lys-tRNA Sase (lieUS)

 1217
 Met-tRNA formyltransferase (fmt)

 1276
 Met-tRNA Sase (metG)

 0394
 peptidyl-tRNA hydrolase (pth)

 1311
 Phe-tRNA Sase α sub (pheS)

 1322
 Pha-tRNA Sase α sub (pheT)
 (054) peptidyminkni ydiotso (pur)
(0729) Pro-tRNA Sase α sub (pheS)
(0729) Pro-tRNA Sase (proS)
(0742) pro-tRNA Sase (proS)
(0745) queuosine biosyn prt (queA)
(0205 selenium metabolism prt (selD)
(0110) Ser-tRNA Sase (serS)
(0202) tRNA (guanine-N1)-MTase (trmD)
(0204) tRNA (guanine-N1)-MTase (trmD)
(0204) tRNA (guanine-N1)-MTase (trmD)
(0205 tRNA h2(2)-isopentenylpyrophosphate
Tase (trpX)
(0205 tRNA nucleotidyttransferase (cca)
(0244) tRNA sase (tyrS)
(130) Tyr-tRNA Sase (tyrS)
(130) Tyr-tRNA Sase (tyrS)
(131) Tyr-tRNA Sase (tyrS)
(132) Val-tRNA Sase (tyrS)
(133) Dasa detring pertirfes and Degradation of proteins, peptides, and Degradation of proteins, peptides, and givcopeptides 0875 aminopeptidase A (pepA) 1705 aminopeptidase A (pepA) 1614 aminopeptidase N (pepN) 0816 aminopeptidase P (pepP) 0714 ATP-dependent cip protease (cipP) 1597 ATP-dependent protease (sms) 0715 ATP-dependent protease ATPase sub (cipY) (clpX) (clpX) (clpX) (059 ATP-dependent protease ATP-binding sub (clpB) 0419 collagenase (prtC) 0150 HfC 0419 Collagerises (phc) 0150 HifC 0950 IgA1 protease (iga1) 0247 IgA1 protease (iga1) 124 Ion protease (ion) 0675 peptidase D (pepD) 0687 peptidase E (pepE) 1348 peptidase E (pepE) 1348 peptidase T (pepT) 1259 periplasmic Ser protease Do (htrA) 0722 Pro dipeptidase (pepQ) 1682 protease (sohB) 1541 protease IV (sppA) 0151 protease for λ cl1 repressor (hflK) 0530 sigloycoprotease (scp) 0150 0530 sialoglycoprotease (gcp) Nucleoproteins 0186 DNA-BP 1491 DNA-BP (rdgB) 1587 DNA-BP H-NS (hns) 0430 DNA-BP HU-α

 O430
 DNA-BP HU-α

 Protein modification and translation factors

 06430
 DNA-BP HU-α

 0945
 DNA processing chain A (dprA)

 0945
 DNA processing chain A (dprA)

 0946
 DNA processing chain A (dprA)

 0947
 elongation factor EF-Ts (tsf)

 0578
 elongation factor EF-Tu (tufB)

 0579
 elongation factor P (etp)

 0522
 elongation factor P (etp)

 0523
 elongation factor P (etp)

 0523
 elongation factor P (etp)

 0524
 elongation factor P (etp)

 0525
 elongation factor P (etp)

 0526
 Glu-ammonia-ligase adenylyltransferase

 (glnE)
 maintation factor IF-3 (infA)

 1318
 initiation factor IF-3 (infC)

 1318
 initiation factor IF-3 (infC)

 1318
 minopeptidase (map)

 0428
 oxido-RDase (dsbB)

 1517
 peptide chain release factor 1 (prfA)

 1521
 peptide chain release factor 2 (prfB)

 1735
 peptidy-prolyl cis-trans isomerase B (op0B)

 0698
 robsome releasing factor (frr)

 0573
 rotamase, peptidyl p

 1213 thiol:disulfide interchange prt (xprA)

 Ribosomal proteins: sthesis and modification

 0516 ribosomal prt L1 (rpL1)

 0640 ribosomal prt L10 (rpL10)

 0517 ribosomal prt L11 (rpL11)

 0978 ribosomal prt L11 (rpL13)

 0788 ribosomal prt L13 (rpL13)

 0788 ribosomal prt L15 (rpL16)

 0789 ribosomal prt L15 (rpL16)

 0784 ribosomal prt L16 (rpL16)

 0785 ribosomal prt L17 (rplQ)

 0784 ribosomal prt L17 (rplQ)

 0794 ribosomal prt L17 (rplQ)

 0794 ribosomal prt L12 (rpL2)

 0707 ribosomal prt L12 (rpL2)

 0710 ribosomal prt L2 (rpL2)

 0728 ribosomal prt L2 (rpL2)

 0738 ribosomal prt L23 (rpL20)

 0789 ribosomal prt L23 (rpL22)

 0779 ribosomal prt L23 (rpL24)

 0788 ribosomal prt L23 (rpL24)

 0789 ribosomal prt L23 (rpL24)

 0879 ribosomal prt L27 (rpL27)

 0879 ribosomal prt L28 (rpL28)

 0879 ribosomal prt L28 (rpL28)

HI

Arc 091

020 047 136

13£ 117

046 123 114

019 154

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247

¥66

20 43

43

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63 08

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08 30) 72

75 25

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702

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989 987

737 58: 58: 19: 73:

8

682

lut 311

727 300

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365 189

719

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103

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506 389

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550 445

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65 87

86

80 70

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93

85

91

 Hitt
 Identification
 %Sim

 0785
 ribosomal pt L29 (rpL29)
 87

 0777
 ribosomal pt L3 (rpL3)
 92

 0796
 ribosomal pt L3 (rpL3)
 96

 0650
 ribosomal pt L3 (rpL31)
 86

 0650
 ribosomal pt L3 (rpL33)
 91

 0996
 ribosomal pt L3 (rpL3)
 93

 1319
 ribosomal pt L4 (rpL4)
 93

 0793
 ribosomal pt L5 (rpL7/L12)
 92

 0641
 ribosomal pt L9 (rpL9)
 96

 1200
 ribosomal pt S1 (rpS1)
 90

 0601
 ribosomal pt S11 (rpS1)
 96

 0797
 ribosomal pt S16 (rpS16)
 87

 0798
 ribosomal pt S16 (rpS16)
 87

 0791
 ribosomal pt S16 (rpS16)
 87

 0791
 ribosomal pt S16 (rpS16)
 87

 0781
 ribosomal pt S17 (rpS7)
 94

 HI# H# **Identification** %Sim

 USB1 streptomycin resistance prt (strA)
 100

 Transport and binding proteins

 Amino acids, peptides and amines
 1177

 1177
 Arg permease (artM)
 60

 1178
 Arg transport ATP-BP artP (artP)
 83

 1285
 biopolymer transport pt (exbD)
 55

 1286
 biopolymer transport pt (exbD)
 55

 1288
 branched chain AA transport system II
 50

 0283
 D-Ala permease (dagA)
 65

 1186
 dipeptide permease (dpD)
 73

 1186
 dipeptide permease (dpD)
 73

 1186
 dipeptide permease (dpD)
 65

 1186
 dipeptide transport ATP-BP (dpDP)
 84

 1186
 dipeptide permease (glnP)
 59

 1186
 dipeptide transport ATP-BP (dpDF)
 87

 1180
 dipopeptide-BP (oppA)
 60

 1180
 dipopeptide permease (glnP)
 50

 1180
 dipopeptide permease (oppE)
 67

 1180
 dipopeptide permease (popA)
 60

 1180
 dipopeptide permease (oppE)
 67

 1180
 dipopeptide permease (op Other (potA) 1344 spermidine-putrescine-BP (potD) 0498 spermidine-putrescine-BP (potD) 0287 Trp-specific permease (mtr) 0287 tyr-specific transport prt (tyrP) 0477 Tyr-specific transport prt (tyrP) 859798688 72 75 73 65 68 Anions 1691 hydrophilic membrane-bound prt (modC) 75 1692 hydrophobic membrane-bound prt (modB) 85 1381 integral membrane prt (pstA) 78 0354 initrate transporter ATPase component 58 õ 73 (nasD) (nasD) 1380 peripheral membrane prt B (pstB) 1382 peripheral membrane prt C (pstC) 1383 periplasmic phosphate-BP (pstS) 79 87 79 68 60 1604 phosphate permease Carbohydrates, organic alcohols, and acids 0020 2-oxogiutarate/malate translocator 0153 Asp transport prt (dcuA) 0746 Asp transport prt (dcuA) 1110 *D*-xylose transport ATP-BP (xyIG) 1111 *D*-xylose-BP (rbsB) 1712 enzyme I (ptsI) 0181 formate transporter 0484 fructose permease IIA/FPR component (fruB) 93 60 70 70 86 83999 88 73 68 96 92 91 (fruB) 0446 fructose permease IIBC component 72
 0446
 fructose permease IIBC component
 72

 0612
 fucose operon prt (fucU)
 80

 1711
 Glo phosphotransferase enzyme III (crr)
 83

 1717
 glocerol uptake facilitator prt (glpF)
 55

 0600
 glocerol uptake facilitator prt (glpF)
 87

 015
 gluconate permease (gntF)
 56

 0686
 glycerol-3-phosphatase transporter (fbpA)
 85

 0503
 high affinity ribose transport prt (rbsA)
 85

 0501
 high affinity ribose transport prt (rbsD)
 78
 96 93 97 86 97 83 387799

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 Identification

 0610
 L-fucose permease (fucP)

 1218
 L-lactate permease (fucP)

 1228
 Lactate permease (fucP)

 1229
 lactate permease (fucP)

 1229
 Lactate permease (fucP)

 0823
 methylgalactoside permease ATP-BP (mgIA)

 0824
 methylgalactoside -BP (mgIB)

 0824
 nethylgalactoside permease (mgIC)

 1630
 Na+ and CI- dependent GABA
 58 54 85 90 53 1000 Vat+and Cho department of vertice transporter
10736 Na+-dependent noradrenaline transporter
10540 periplasmic ribos-BP (rbsB)
1713 phosphohistidinoprotein-hexose phosphotransferase (ptsH)
1028 potassium channel homolog (kch)
1109 ribose permease (xylH) 87 88 80 84 Cations Cations 0254 bacterioferritin comigratory prt (bcp) 0251 energy transducer (tonB) 1272 ferric enterobactin transport ATP-BP 80 98 energy transducer (tonB)
ferric enterobactin transport ATP-BP (fepC)
ferric enterobactin transport ATP-BP (fepC)
ferric enterobactin transport ATP-BP (fepC)
ferriin like prt (rsgA)
ferriin dita transport ATP-BP (fecE)
magnesium and cobalt transport prt (corA)
magnesium and cobalt transport ptt (corA)
magnesium and cobalt transport ptt (merT)
mover and transport ptt (merT)
mover and transport ptt (merP)
to mercury transport ptt (merP)
to mover antipoter (nhaB)
to part, H+ antipoter (nhaC)
periplasmic-BP-dependent iron transport (stuB)
fard periplasmic-BP-dependent iron transport (stuC) 51 55 49 74 79 61 56 85 82 54 46 67 43 87 62 75 59 (STUB) 1474 periplasmic-BP-dependent iron transport (sfuC) 58 (sfuC) 66 0911 potassium efflux system (kefC) 66 0290 potassium, copper-transporting ATPase 64 A (copA) 79 1352 sodium, Pro symporter (putP) 79 0625 TRK system potassium uptake prt (trkA) 83 Nucleosides, purines and pyrimidines 1087 ribonucleotide transport ATP-BP (mkl) 1227 uracil permease (uraA) 61 62 Other 0621 ATP-BP (abc) 0060 ATP-dependent translocator (msbA) 1619 cystic fibrosis transmembrane conductance regulator 0633 heme-binding ipp (dopA) 0264 heme-hemopexin-BP (hxuA) 1471 hemin permease (hemU) 0262 hemin receptor precursor (hemR) 1706 high-affinity choline transport pt (betT) 0661 lactoterrin-BP (lbpA) 0600 Na+, sulfate cotransporter 0975 pantothenate permease (panF) 09712 transferrin-BP (tbA) 0712 transferrin-BP (tbp1) 1565 transferrin-BP 1 (tbp1) 1276 transferrin-BP 1 (tbp1) 0394 transferrin-BP 1 (tbp1) 0353 transferrin-BP 1 (tbp1) 0353 transferrin-BP 1 (tbp2) 0365 transport ATP-BP (cydD) 1157 transport ATP-BP (cydD) 100 61 89 33 46 62 Δ۶ 48 78 48 458883557 1157 transport ATP-BP (cydD) Other categories Adaptations and atypical conditions 1526 autotrophic growth prt (aut) (071 heat shock pt R253 (grpE) 0720 heat shock pt (htpX) 1527 heat shock pt (htpX) 1527 heat shock pt (htpA) 1537 heat shock pt (htpA) 1544 NAD(P)H:menadione oxidoreductase 0456 survival pt (surA) 0815 universal stress pt (uspA) 1251 virulence assoc pt C (vapC) 0947 virulence assoc pt C (vapC) 0947 virulence assoc pt C (vapC) 0347 virulence plasmid pt (mlgA) 0321 virulence plasmid pt (mlgA) 0321 virulence functions 61682717365585755566558 Colicin-related functions C382 colicin tolerance pt (toIB) 1206 colicin V production pt (cvpA) 0384 inner membrane pt (toIR) C385 inner membrane pt (toIQ) 1685 outer membrane integrity pt (toIA) 0383 outer membrane integrity pt (toIA) 78 79 79 80 48 57 Drug and analog sensitivity 0995 acritlavine resistance prt (acrB) 0300 ampD signalling pt (ampD) 1242 bicyclomycin resistance prt (bcr) 1623 mercury resistance regulatory prt (merR2) 55 75 69 58 (merFI2) 0648 modulator of drug activity (mda66) 0897 multidrug resistance prt (emrB) 0896 multidrug resistance prt (ermA) 0036 multidrug resistance prt (mdl) 75 86 66 51

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 Hit
 Idemucation

 1462
 nodulation prt T (nodT)

 0549
 rRNA (adenosine-N6.N6-)-dimethyltransferase (ksgA)

 0511
 tellurite resistance prt (tehA)

 1275
 tellurite resistance prt (tehB)

 Phage-related functions and prophages

 1488
 E16 pt (muE16)

 1503
 G prt (muG)

 1588
 G pt (muG)

 1483
 gam pt

 C411
 host factor-I (HF-I) (hfq)

 1504
 I prt (muB)

 1516
 N prt (muN)

 1516
 P pt (muP)

 1411
 terminase sub 1

 1478
 transposase A (muA)
 Radiation sensitivity 0952 DNA repair prt (radC) Transposon-related functions 1577 IS1016-V6 1329 IS1016-V6 1018 IS1016-V6 Other 1161 15 kD ptt (P15) 0055 2-hydroxyacid dehydrogenase (ddr.) 0460 β-lactamase regulatory ptt (mazG) 0223 chloramphenicol-sensitive ptt (rarC) 0690 chloramphenicol-sensitive ptt (rarC) 1670 conjugative transfer co-repressor (info) 0307 & 1-pyroline-5-carboxylate RDase (proC 5-00 betworkst maturation ptt (devA) 1670 conjugative transfer co-repressor (inO) 0307 &-1-pyrroline-5-carboxylate RDase (proC) 1549 heterocyst maturation prt (devA) 1339 embryonic abundant prt, group 3 0916 export factor homolog (skp) 0367 extragenic suppressor (suhB) 0467 gip regulon prt (glpX) 013 glyoxylate-induced prt 0496 heat shock prt (hsIU) 0496 heat shock prt (hsIU) 0496 heat shock prt (hsIU) 0496 heat shock prt (hsIV) 1117 ilv-related prt 0285 isochorismate Sase (entC) 1618 membrane assoc ATPase (cbiO) 0461 membrane prt (lapB) 119 membrane prt (lapB) 0530 mucoid status locus prt (mucB) 0538 A/carbamyl-L-amino acid amidohydrolase 1295 nitrogen fixation prt (nifS) 1343 nitrogen fixation prt (nifS) 1343 nitrogen fixation prt (nifS) 1367 nitrogen fixation prt (nifS) 1368 nitrogen fixation prt (nifS) 1379 nitrogen fixation prt (nifS) 1389 nitrogen fixation prt (nifS) 1398 nitrogen fixation prt (nifS) 1399 nitrogen fixation prt (nifS) 1399 nitrogen fixation prt (nifS) 1495 partitioning system prt (parB) 171 phenolhydroxylase 0368 prt E (gpcE) 0556 putative glucose-6-P Dhase isozyme i (devB) Construction of the second secon 1407 traN 0664 transport ATP-BP (cydC) 1156 transport ATP-BP (cydC) 1556 vanamycin-resistance prt (vanH)

the templ between t 4) To c by the oth order of 1 formed am reaction (range (XL was done of physica DNA fing and the proparticularly jacent to ea ber of cor achieve cor strategies to genome pro efficiency c the program ysis data, i forward and clones, and the relative by physical gaps ordered techniques i Lambda c completion assembly. It ments of the nonclonable they would p the E. coli ho provide DNA such genes w duction. Fur tion from the particularly su viding genera sembly. Becau likely to span mately 100 r from the ampl prepared, and tained from

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3) Th

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3) The two λ libraries constructed from H. influenzae genomic DNA were probed with oligonucleotides designed from the \mathbf{q} ends of contig groups (27). The positive plaques were then used to prepare templates, and the sequence was determined from each end of the λ clone insert. These sequence fragments were searched with

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46 81

62 71

535454555753653

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72

61

68 73 73

35526

89

(finO)

vdrolase

GRASTA against a database of all contigs. Two contigs that matched the sequence from the opposite ends of the same λ clone were ordered. The λ clone then provided the template for closure of the sequence gap between the adjacent contigs.

4) To confirm the order of contigs found by the other approaches and establish the order of the remaining contigs, we performed amplifications by polymerase chain reaction (PCR), both standard and long range (XL) (28). Although a PCR reaction was done for essentially every combination of physical gap ends, techniques such as DNA fingerprinting, database matching, and the probing of large insert clones were particularly valuable in ordering contigs adjacent to each other and reducing the num-77 ber of combinatorial PCRs necessary to achieve complete gap closure. Use of these strategies to an even greater extent in future genome projects will increase the overall efficiency of complete genome closure. In the program ASM_ALIGN Southern analysis data, identification of peptide links, 48 forward and reverse sequence data from λ clones, and PCR data are used to establish 68 57 the relative order of the contigs separated by physical gaps. The number of physical 91 gaps ordered and closed by each of these ve growth Stechniques is summarized in Table 2.

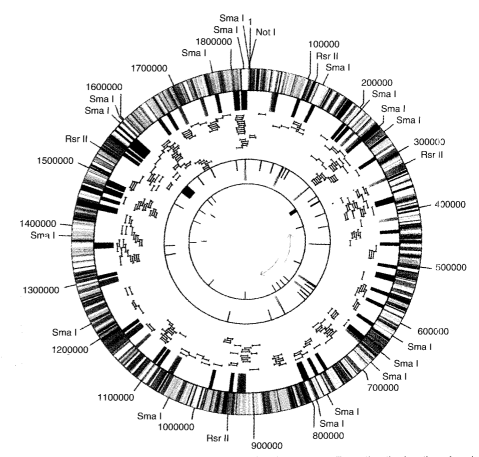
Lambda clones were a central feature for ⁵⁶ ⁷⁶ recompletion of the genome sequence and ⁵⁶ assembly. It was probable that some fraggments of the H. influenzae genome would be nonclonable in a high copy plasmid because they would produce deleterious proteins in the *E*. coli host cells. Lytic λ clones would provide DNA for these segments because such genes would not inhibit plaque production. Furthermore, sequence information from the ends of 15- to 20-kb clones is particularly suitable for gap closure and providing general confirmation of genome assembly. Because of their size, they would be kely to span any physical gap. Approximately 100 random plaques were picked from the amplified λ library, templates were prepared, and sequence information was obained from each end. These sequences were searched (GRASTA) against the conigs and linked in the database to their ppropriate contig, thus providing a scafblding of λ clones that contributed addiional support to the accuracy of the genome assembly (Fig. 1). In addition to conirmation of the contig structure, the λ dones provided closure for 23 physical gaps.

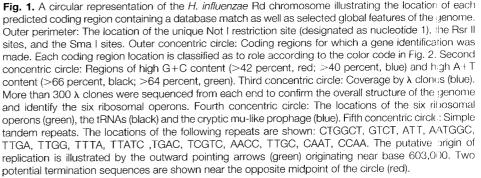
Approximately 78 percent of the genome was covered by λ clones.

The λ clones were particularly useful for solving repeat structures. All repeat structures identified in the genome were small enough to be spanned by a single clone from the random insert library, except for the six ribosomal RNA (rRNA) operons and one repeat (two copies) that was 5340 bp in length. The ability to distinguish and assemble the six rRNA operons of H. influenzae (each containing in order 16S, 23S, and 5S subunit genes) was a test of our overall strategy to sequence and assemble a complex genome that might contain a significant number of repeat regions. The high degree of sequence similarity and the length of the six operons caused the assembly process to cluster all the underlying sequences into a few indistinguishable contigs. To de-

termine the correct placement of the operons in the sequence, unique sequences were identified at the 5S ends. Oligonucleotide primers were designed from these six flanking regions and used to probe the two λ libraries. For five of the six rRNA operons at least one positive plaque was identified that completely spanned the rRNA operon and contained uniquely identifying flanking sequence at the 16S and 5S ends. These plaques provided the templates for obtaining the sequence for these rRNA operons. For rrnA a plaque was identified that contained the particular 58 end and terminated in the 16S end. The 16S end of rrnA was obtained by PCR from H. influenzae Rd genomic DNA.

An additional confirmation of the global structure of the assembled circular genome was obtained by comparing a computer-





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generated restriction map based on the assembled sequence for the endonucleases Apa I, Sma I, and Rsr II with the predicted physical map of Lee et al. (29). The restriction fragments from the sequence-derived map matched those from the physical map in size and relative order (Fig. 1).

At the same time that the final gap filling process occurred, each contig was edited visually by reassembling overlapping 10-kb sections of contigs by means of the AB AUTOASSEMBLER and the Fast Data Finder hardware. AUTOASSEMBLER provides a graphical interface to electropherogram data for editing. The electropherogram data was used to assign the most likely base.at each position. Where a discrepancy could not be resolved or a clear assignment made, the automatic base calls were initially left unchanged. Individual sequence changes were written to the electropherogram files and a program was designed (CRASH) to maintain the synchrony of sequence data between the H. influenzae database and the electropherogram files. After the editing, contigs were reassembled with TIGR AS-SEMBLER prior to annotation.

Potential frameshifts identified in the course of annotating the genome were saved as reports in the database. These frameshifts were used to indicate areas of the sequence that might require further editing or sequencing. Frameshifts were not corrected for cases in which clear electropherogram data disagreed with a frameshift. Frameshift editing was done with TIGR EDITOR. This program was developed as a collaborative effort between TIGR and AB and is a modification of the AB AUTOAS-SEMBLER. TIGR EDITOR can download contigs from the database and thus provides a graphical interface to the electropherogram for the purpose of editing data associated with the aligned sequence file output of TIGR ASSEMBLER. The program maintains synchrony between the electropherogram files on the Macintosh system and the sequence data in the H. influenzae database on the Unix system. TIGR EDITOR is now our primary tool for sequence viewing and editing for the purpose of genome assembly.

The final assembly of the H. influenzae genome with the TIGR ASSEMBLER was precluded by the rRNA and other repeat regions, and was accomplished by means of COMB_ASM (a program written at TIGR) that splices together contigs on the basis of short sequence overlaps.

Throughout the project, we paid particular attention to the accuracy of the sequence generated and included various quality control measures. In particular, we constructed random small and large insert libraries (as described above), used strict criteria for excluding any single sequence in which more than 3 percent of the nucleotides could not be identified with certainty, determined that there was no vector contamination in each sequence, and rejected chimeric sequences from the assembly process. The most important measure of the sequence accuracy is the correct assembly of the 1.8-Mb genome. Any deviation from inclusion of only high-quality sequences would have resulted in an inability to assemble the final genome. In addition, the use of the large insert λ clones confirmed the accuracy of the final assembly. Our finding that the restriction map of the H. influenzae Rd genome based on our sequence data is in complete agreement with that previously published (29) further confirms the accuracy of the assembly.

As a consequence of our shotgun approach, we reached an average of more than sixfold redundancy across the genome, although there are some regions in which the coverage is lower. The criteria that we used to define overall sequence quality and completion were as follows: (i) The sequence should have less than 1 percent single sequence coverage. Because H. influenzae is a genome rich in AT pairs, it is possible to obtain a highly accurate sequence with single-pass coverage. However, any regions with single sequence coverage that contained ambiguities were again sequenced with an alternative sequencing chemistry. (ii) Areas with more than single sequence coverage that contained ambiguities or G-C compressions were also sequenced again with an alternative sequencing chemistry. The combination of sequence redundancy together with the application of an alternative sequencing chemistry in areas with ambiguities is, we believe at least as accurate, if not more so, than double-stranded coverage. By these criteria we have reduced the number of nucleotide ambiguities [International Union of Biochemistry (IUB) codes] in the sequence to less than 1 in 19,000. The same approaches used to resolve ambiguities were also applied to areas where apparent frameshifts were indicated. Sixty potential frameshifts were identified by comparison to entries in peptide databases. Although some of these potential frameshifts are undoubtedly real, others may reflect the hundreds of frameshifts present in GenBank sequences from public databases (30). They may also represent biologically significant phenomena such as insertions or deletions in insertion elements, or in tandem repeats often associated with virulence genes (31).

We also considered comparison of our sequence to existing GenBank H. influenzae Rd sequences as a method for evaluating sequence accuracy as reported for yeast chromosome VIII (32). Unlike yeast, only a limited number of H. influenzae sequences are in GenBank (38 H. influenzae Rd accessions) and these are not necessarily of high

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accuracy. The results of such a comparison show that our sequence is 99.67 percent identical overall to those GenBank sequences annotated as H. influenzae Rd. Two problems were apparent with this type of comparison. Sequences could differ because of strain variation, which is poorly annotated in the GenBank entries. It is also difficult to evaluate the significance of differences as the accuracy of the GenBank entries was impossible to assess. We compared GenBank accession M86702 (strA resistance gene) to our sequence and found the identity to be 94.7 percent over 545 bp. There are 24 single base pair mismatches relative to our sequence as well as an insertion and a deletion. Comparison of our sequence to GenBank accession L23824 (adenylate cyclase) shows a 99.7 percent match over 2960 bp. There are nine single base pair mismatches and one insertion. In this case the mismatches all fall in the noncoding flanking regions. While we cannot speak to the accuracy of these GenBank sequences, we are very confident of our sequences in these regions because of the $3 \times$ to $9 \times$ coverage with high-quality sequence data. Thus, a comparison of our sequence to sequences in GenBank annotated as H. influenzae Rd is not a meaningful way to evaluate the accuracy of the sequence.

Although it is extremely difficult to assess sequence accuracy, we wanted to provide an approximation of accuracy based on frequency of shifts in open reading frames, unresolved ambiguities, overall quality of raw data, and fold coverage. We estimate our error rate to be between 1 base in 5000 rRNAs, as and 1 base in 10,000.

We also attempted to estimate the costsites, replica of the complete sequencing of the genome composition Reagent and labor costs for construction of chemical ar small insert and λ libraries, template prep many of the aration and sequencing, gap closure, se include a de quence confirmation, annotation, and preplobvious sequ aration for publication were summed and The H. ir divided by the genome length. Sequencinglar chromoso projects that require up front mappingall G+C n should include the cost of construction emately 38 p the clone maps for sequencing. Not includepercent; G, 1 ed were costs associated with developmenG+C conten of technology and software that will be usewith several for future sequencing projects. The estimatglobal structu ed direct cost was 48 cents per finished bas5000 bp, the pair. Because of the techniques developeeven except t during this project any future genomes G+C and sev). The G+(this size should cost less.

Data and software availability. The Hix rRNA op influenzae genome sequence has been deportophage. Ger ited in the Genome Sequence DataBao proteins er (GSDB) with the accession number L4202re located at and is termed version 1.0. The nucleotic 1.59 Mbp of sequence and peptide translation of eacenome has a predicted coding region with identified stænt than ave and stop codons have also been accessionercent G+C.

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by GSDB. We consider annotation, accuracy checking, and error resolution to be ongoing tasks. As outlined above, there are _predicted coding regions with potential frameshift errors in the sequence. As these are resolved, they will be deposited with GSDB. We also expect the annotation of the sequence to increase over time and be updated in GSDB.

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Additional data are available on our World Wide Web site (http://www.tigr.org). An expanded version of Table 3 has links to the database accessions that were used to identify the predicted coding regions, additional sequence similarity data, and coordinates of the predicted coding regions. The alignments between the predicted coding regions and the database sequences are also available. The data can also be queried by gene identification number, putative identification, matching accession, and role. The entire sequence and the sequences of all predicted coding regions and their translations, including those having frameshifts, are also available. This Web site will be -maintained as an up-to-date source of H. the influenzae genome sequence data, and we encourage the scientific community to forour ward their results for inclusion (with proper noattribution) at this site. ng-

The software developed at TIGR that is the described in the article is still under development. However, TIGR will work with asother genome centers to make its software proavailable upon request. lon

Genome analysis. We have attempted nes, of ppredict all of the coding regions and nate dentify genes, transfer RNAs (tRNAs) and RNAs, as well as other features of the 000 MA sequence (such as repeats, regulatory cost ites, replication origin sites, and nucleotide imposition), with the realization that biome. themical and biological conformation of n of many of these will be an ongoing task. We repnclude a description of some of the most sebyious sequence features. rep-

and The H. influenzae Rd genome is a circucing I chromosome of 1,830,137 bp. The overping I G+C nucleotide content is approxin of lately 38 percent (A, 31 percent; C, 19 lud- trcent; G, 19 percent; T, 31 percent). The nent <code>____HC</code> content of the genome was examined used ith several window lengths to look for nat- Jobal structural features. With a window of base 100 bp, the G+C content is relatively pped wen except for seven large regions rich in s of θ +C and several regions rich in A,+T (Fig.

). The G+C-rich regions correspond to H. x rRNA operons and a cryptic mu-like pos- tophage. Genes for several proteins similar Base proteins encoded by bacteriophage mu 2023 re located at approximately position 1.56 tide 1.59 Mbp of the genome. This area of the each mome has a markedly higher G+C constart nt than average for H. influenzae (~ 50 oned excent G+C compared to \sim 38 percent for

the rest of the genome).

The minimal origin of replication (oriC) in E. coli is a 245-bp region defined by three copies of a 13-bp repeat at one end (sites for initial DNA unwinding) and four copies of a 9-bp repeat (sites for DnaA binding, the first step in replication) at the other (33). An approximately 280-bp sequence containing structures similar to the three 13-bp and four 9-bp repeats defines the putative origin of replication in H. influenzae Rd. This region lies between sets of ribosomal operons rmF, rmE, rmD and rmA, rmB, rmC. These two groups of ribosomal operons are transcribed in opposite directions and the placement of the origin is consistent with their polarity for transcription. Termination of E. coli replication is marked by two 23-bp termination sequences located \sim 100 kb on either side of the midway point at which the two replication forks meet. Two potential termination sequences sharing a 10-bp core sequence with the E. coli termination sequence were identified in H. influenzae. These two regions are offset approximately 100 kb from a point approximately 180° opposite of the proposed origin of *H*. influenzae replication.

Six rRNA operons were identified. Each contains three subunits and a variable spacer region in the order: 16S subunit-spacer region—23S subunit—5S subunit. The subunit lengths are 1539, 2653, and 116 bp, respectively. The G+C content of the three ribosomal subunits (50 percent) is higher than that of the genome as a whole. The G+C content of the spacer region (38 percent) is consistent with the remainder of the genome. The nucleotide sequence of the three rRNA subunits is completely identical in all six ribosomal operons. The rRNA operons can be grouped into two classes based on the spacer region between the 16S and 23S sequences. The shorter of the two spacer regions is 478 bp (rrnb, rmE, and rmF) and contains the gene for tRNA^{Glu}. The longer spacer is 723 bp (rmA, rmC, and rmD) and contains the genes for tRNA^{Ile} and tRNA^{Ala}. The two sets of spacer regions are also completely identical across each group of three operons. Other tRNA genes are present at the 16S and 5S ends of two of the rRNA operons. The genes for tRNAArg, tRNAHis, and tRNA^{Pro} are located at the 16S end of *rmE* while the genes for $tRNA^{Trp}$ and $tRNA^{Asp}$ are located at the 5S end of rmA.

The predicted coding regions were initially defined by evaluating their coding potential with the program GENEMARK (34) based on codon frequency matrices derived from 122 H. influenzae coding sequences in GenBank. The predicted coding region sequences (plus 300 bp of flanking sequence) were used in searches against a database of nonredundant bacterial proteins

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(NRBP) created specifically for the annotation. Redundancy was removed from NRBP at two stages. All DNA coding sequences were extracted from GenBank (release 85), and sequences from the same species were searched against each other. Sequences having more than 97 percent identity over regions longer than 100 nucleotides were combined. In addition, the sequences were translated and used in protein comparisons with all sequences in Swiss-Prot (release 30). Sequences belonging to the same species and having more than 98 percent similarity over 33 amino acids were combined. NRBP is composed of 21,445 sequences extracted from 23,751 GenBank sequences and 11,183 Swiss-Prot sequences from 1099 different species.

A total of 1743 predicted coding regions was identified. Searches of the predicted coding regions for H. influenzae were performed against NRBP with BLAZE (35) run on a Maspar MP-2 massively parallel computer with 4096 microprocessors. BLAZE translates the query DNA sequence in the three plus-strand reading frames and identifies the protein sequences that match the query. The protein-protein matches were aligned with PRAZE, a modified Smith-Waterman (23) algorithm. In cases where insertions or deletions in the DNA sequence produced a potential frameshift, the alignment algorithm started with protein regions of maximum similarity and extended the alignment to the same database match in alternative frames by means of the 300-bp flanking region. Unidentified predicted coding regions and the remaining intergenic sequences were searched against a dataset of all available peptide sequences from Swiss-Prot, the Protein Information Resource (PIR), and GenBank. Identification of operon structures is expected to be facilitated by experimental determination of promoter and termination sites.

Each putatively identified H. influenzae gene was assigned to one of 102 biological role categories adapted from Riley (36). Assignments were made by linking the protein sequence of the predicted coding regions with the Swiss-Prot sequences in the Riley database. Of the 1743 predicted coding regions, 736 have no role assignment. Of these, no database match was found for 389, while 347 matched "hypothetical proteins" in the database. Role assignments were made for 1007 of the predicted coding regions. Each of the 102 role categories was grouped into one of 14 broader role categories (Table 2). A compilation of all the predicted coding regions, their identifiers, a three-letter gene identifier, and percent similarity are presented in Table 3 (foldout). An annotated complete genome map of H. influenzae Rd is presented in Fig. 2 (fold-out). The map places each predicted

coding region on the H. influenzae chromosome, indicates its direction of transcription and color codes its role assignment. Role assignments are also represented in Fig. 1.

A survey of the genes and their chromosomal organization in H. influenzae Rd makes possible a description of the metabolic processes H. influenzae requires for survival as a free-living organism, the nutritional requirements for its growth in the laboratory, and the characteristics that make it different from other organisms specifically as they relate to its pathogenicity and virulence. The genome would be expected to have complete complements of certain classes of genes known to be essential for life. For example, there is a one-to-one correspondence of published E. coli ribosomal protein sequences to potential homologs in the H. influenzae database. Likewise, as shown in Table 3, an aminoacyl tRNA synthetase is present in the genome for each amino acid. Finally, the location of tRNA genes was mapped onto the genome. There are 54 identified tRNA gencs, including representatives of all 20 amino acids.

In order to survive as a free-living organism, H. influenzae must produce energy in the form of ATP via fermentation or electron transport. As a facultative anaerobe, H. influenzae Rd is known to ferment glucose, fructose, galactose, ribose, xylose, and fucose (37). As indicated by the genes identified in Table 3, transport systems are available for the uptake of these sugars by the phosphoenolypyruvate-phosphotransferase system (PTS), and by non-PTS mechanisms. Genes that specify the common phosphate-carriers enzyme I and Hpr (ptsI and ptsH) of the PTS system were identified as well as the glucose-specific crr gene. We have not, however, identified the gene-encoding, membrane-bound, glucosespecific enzyme II. The latter enzyme is required for transport of glucose by the PTS system. A complete PTS system for fructose was identified.

Genes encoding the complete glycolytic pathway and for the production of fermen-

tative end products were identified. Also identified were genes encoding functional anaerobic electron transport systems that depend on inorganic electron acceptors such as nitrates, nitrites, and dimethyl sulfoxide. Genes encoding three enzymes of the tricarboxylic acid (TCA) cycle appear to be absent from the genome. Citrate synthase, isocitrate dehydrogenase, and aconitase were not found by searching the predicted coding regions or by using the E. coli enzymes as peptide queries against the entire genome in translation. This provides an explanation for the large amount of glutamate (1 g/liter) that is required in defined culture media (38). Glutamate can be directed into the TCA cycle by conversion to α -ketoglutarate by glutamate dehydrogenase. In the absence of a complete TCA cycle, glutamate presumably serves as the source of carbon for biosynthesis of amino acids from precursors that branch from the TCA cycle. Functional electron transport systems that depend on oxygen as a terminal electron acceptor are available for the production of adenosine triphosphate.

Previously unanswered questions regarding pathogenicity and virulence can be addressed by examining certain classes of genes such as adhesins and the lipo-oligosaccharide biogenesis genes. Moxon and coworkers (31) have obtained evidence that a number of these virulence-related genes contain tandem tetramer repeats that undergo frequent addition and deletion of one or more repeat units during replication such that the reading frame of the gene is changed and its expression thereby altered. It is now possible, by means of the complete genome sequence, to locate all such tandem repeat tracts (Fig. 2) and to begin to determine their roles in phase variation of such potential virulence genes.

Haemophilus influenzae Rd has a highly efficient, DNA transformation system. The DNA uptake sequence site, 5' AAGTGC-GGT, present in multiple copies in the genome, is necessary for efficient DNA uptake (39). It is now possible to locate all of these

Table 4. Two-component systems in H. influenzae Rd. ID, identity; Sim, similarity.

Identification	Location	Best	ld	Sim	Length
number		match*	(%)	(%)	(bp)
HI0220 HI0267 HI1707 HI1378 HI0726 HI0837 HI0884 HI1379 HI1379 HI1378	239,378 299,541 1,781,143 1,475,017 777,934 887,011 936,624 1,475,502 1,781,799	Sensors arcB narQ basS phoR Regulators narP cpxR arcA phoB basR	39.5 38.1 27.7 38.1 59.3 51.9 77.2 52.9 43.5	63.9 68.0 51.5 61.6 77.0 73.0 87.8 71.4 59.3	200 562 250 280 209 229 236 228 219

"In all cases, the best match was to a gene of E. coli-

sites and describe their distribution with respect to genic and intergenic regions (40). Fifteen genes involved in transformation have already been described and sequenced (41). Six of the genes, comA to comF, comprise an operon that is under positive control by a 22-bp, palindromic, competence regulatory element (CRE) located approximately one helix turn upstream of the promoter. It is now feasible to locate additional copies of CRE in the genome and discover potential transformation genes under CRE control (42). In addition, other global regulatory elements may be discovered with an ease not previously possible.

One well-described system for gene regulation in bacteria is the "two-component" system composed of a sensor molecule that detects an environmental signal and a regulator molecule that is phosphorylated by the activated form of the sensor. The regulator protein is generally a transcription factor that, when activated by the sensor, turns on or off expression of a specific set of genes. It has been estimated that *E*. coli harbors 40 sensor-regulator pairs (43). The H. influenzae genome was searched with representative proteins from each family of sensor and regulator proteins with TBLASTN and TFASTA. Four sensor and five regulator proteins were identified with similarity to proteins from other species (Table 4). There appears to be a corresponding sensor for each regulator protein except CpxR. Searches with the CpxA protein from E. col identified three of the four sensors listed in Table 4, but no additional significant matches were found. It is possible that the sequence similarity is low enough to be undetectable with TFASTA. All of the regulator proteins present fall into the OmpR subclass (43). No representatives of the NtrC class of regulators were found. This class of proteins interacts directly with the sigma-54 subunit of RNA polymerase, which is absent from H. influenzae, and which plays a major role in the regulation of a large number of operons in E. coli and other enterobacteria. The absence of the Ntr network in H. influenzae suggests significant differences in the regulatory processes between these two groups of organisms.

Some of the most interesting questions that can be answered by a complete genome sequence relate to the genes or pathway that are absent. The nonpathogenic H. m fluenzae Rd strain varies significantly from the pathogenic serotype b strains. Many @ the differences between these two strainappear in factors affecting infectivity. For example, we have found that the eight genes that make up the fimbrial gene cluster (44) involved in adhesion of bacteria to host cells are absent in the Rd strain. The pepN and purE genes, which flank the fimbrial cluster in H. influenzae type b strains

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Haemophilus influenzae type b pepN hifA hifY3 H hifY1 hifY2 hifB purE Haemophilus influenzae Rd DepN purE

Fig. 3. A comparison of the region of the *H. influenzae* chromosome containing the eight genes of the fimbrial gene cluster present in *H. influenzae* type b and the same region in *H. influenzae* Rd. The region is flanked by *pepN* and *purE* in both organisms. However, in the noninfectious Rd strain the eight genes scover of the fimbrial gene cluster have been excised. A 172-bp spacer region is located in this region in the Rd r CRE strain and continues to be flanked by the *pepN* and *purE* genes.

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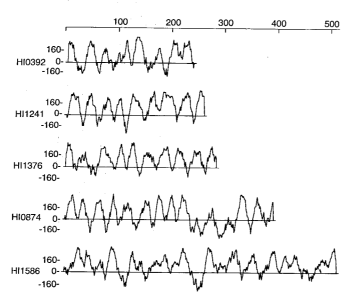
On a broader level, we determined e tha a reg which E. coli proteins are not in H. influted blenzae by taking advantage of a nonredunregulation and set of protein-coding genes from E. on factioli, namely the University of Wisconsin , turn Genome Project contigs in GenBank: 1216 genespredicted protein sequences from GenBank ors 4 accessions D10483, L10328, U00006, ifluer, U00039, U14003, and U18997 (45). The senta minimum threshold for matches was set so or an that even weak matches would be scored as an positive, thereby giving a minimal estimate gulate of the E. coli genes not present in H. influrity tlenzae. We used TBLASTN to search each Ther of the E. coli proteins against the complete or fogenome. All BLAST scores greater than CpxR100 were considered matches. Altogether E. cd627 E. coli proteins matched at least one sted ilregion of the H. influenzae genome and 589 ifican proteins did not. The 589 nonmatching at there examined and found to conbe untain a disproportionate number of hyporegulatical proteins from E. coli. Sixty-eight Omplercent of the identified E. coli proteins of there matched by an H. influenzae sequence . Thiwhereas only 38 percent of the hypothetical th theoreteins were matched. Proteins are annoheras

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A total of 389 predicted coding regions did not display significant similarity with a six-frame translation of GenBank release 87. These unidentified coding regions were compared to one another with FASTA. Two previously unidentified gene families were identified. Two predicted coding regions without database matches (HI0589 and HI0850) share 75 percent identity over almost their entire lengths (139 and 143 amino acid residues respectively). A second pair of predicted coding regions (HI1555 and HI1548) encode proteins that share 30 percent identity over almost their entire lengths (394 and 417 amino acids respectively). These similarities suggest that there may be previously unidentified gene families present in these regions.



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Another analysis that can be applied to the unidentified coding regions is hydropathy analysis, which indicates the patterns of potential membrane-spanning domains that are often conserved between members of receptor and transporter gene families, even in the absence of significant amino acid identity. The five best examples of unidentified predicted coding regions that display potential transmembrane domains with a periodic pattern that is characteristic of membrane-bound channel proteins are shown in Fig. 4. Such information can be used to focus on specific aspects of cellular function that are affected by targeted deletion or mutation of these genes.

We have learned some important lessons concerning overall strategy from the H. influenzae sequencing project that should reduce the effort required for future bacterial genome sequencing projects. For example, the small insert library and the large insert library should be constructed and end-sequenced concurrently. It is essential that the sequence fragments used for the assembly are of the highest quality. The sequences should be rigorously checked for vector contamination. Although it is important that sequence read lengths be long enough to span most small repeats, they must also be highly accurate. Our raw sequence data contained on average less than 1.5 percent uncertainties. The use of high quality individual sequence fragments and a rigorous assembly algorithm essentially eliminated difficulty with achieving closure. The success of whole genome shotgun sequencing offers the potential to accelerate research in a number of areas. Comparative genomics could be advanced by the availability of an increased number of complete genomes from a variety of prokaryotes and eukaryotes. Knowledge of the complete genomes of pathogenic organisms could lead to new vaccines. Information obtained from the genomes of particular organisms could have industrial applications. Finally, this strategy has potential to facilitate the sequencing of the human genome.

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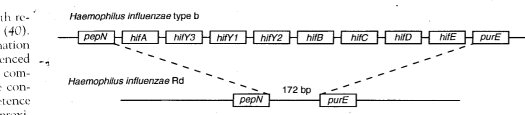


Fig. 3. A comparison of the region of the H. influenzae chromosome containing the eight genes of the fimbrial gene cluster present in H. influenzae type b and the same region in H. influenzae Rd. The region is flanked by pepN and purE in both organisms. However, in the noninfectious Rd strain the eight genes of the fimbrial gene cluster have been excised. A 172-bp spacer region is located in this region in the Rd strain and continues to be flanked by the pepN and purE genes.

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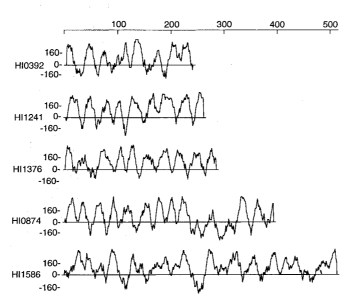
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RESEARCH ARTICLE

Another analysis that can be applied to the unidentified coding regions is hydropathy analysis, which indicates the patterns of potential membrane-spanning domains that are often conserved between members of receptor and transporter gene families, even in the absence of significant amino acid identity. The five best examples of unidentified predicted coding regions that display potential transmembrane domains with a periodic pattern that is characteristic of membrane-bound channel proteins are shown in Fig. 4. Such information can be used to focus on specific aspects of cellular function that are affected by targeted deletion or mutation of these genes.

We have learned some important lessons concerning overall strategy from the H. influenzae sequencing project that should reduce the effort required for future bacterial genome sequencing projects. For example, the small insert library and the large insert library should be constructed and end-sequenced concurrently. It is essential that the sequence fragments used for the assembly are of the highest quality. The sequences should be rigorously checked for vector contamination. Although it is important that sequence read lengths be long enough to span most small repeats, they must also be highly accurate. Our raw sequence data contained on average less than 1.5 percent uncertainties. The use of high quality individual sequence fragments and a rigorous assembly algorithm essentially eliminated difficulty with achieving closure. The success of whole genome shotgun sequencing offers the potential to accelerate research in a number of areas. Comparative genomics could be advanced by the availability of an increased number of complete genomes from a variety of prokaryotes and eukaryotes. Knowledge of the complete genomes of pathogenic organisms could lead to new vaccines. Information obtained from the genomes of particular organisms could have industrial applications. Finally, this strategy has potential to facilitate the sequencing of the human genome.

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- 15. Haemophilus influenzae Rd KW20 DNA was prepared by extraction with phenol. A mixture (3.3 ml) containing 600 µg of DNA, 300 mM sodium acetate, 10 mM tris-HCl, 1 mM Na-EDTA, and 30 percent glycerol was sonicated (Branson Model 450 Sonicator) at the lowest energy setting for 1 minute at 0°C with a 3-mm probe. The DNA was precipitated in ethanol and redissolved in 500 μ l of tris-EDTA (TE) buffer to create blunt ends; a 100-µl portion was digested for 10 minutes at 30°C in 200 µl of BAL 31 buffer with 5 units of BAL 31 nuclease (New England BioLabs). The DNA was extracted with phenol, precipitated in ethanol, redissolved in 100 μl of TE buffer, and fractionated on a 1.0 percent low melting agarose gel. A fraction (1.6 to 2.0 kb) was excised, extracted with phenol, and redissolved in 20 μl of TE buffer. A two-step ligation procedure was used to produce a plasmid library in which 97 percent of the recombinants contained inserts, of which >99 per-cent were single inserts. The first ligation mixture (50 μ l) contained 2 μg of DNA fragments, 2 μg of Sma I + bacterial alkaline phosphatase pUC18 DNA (Pharmacia), and 10 units of T4 ligase (Gibco/BRL), and incubation was at 14°C for 4 hours. After extraction with phenol and ethanol precipitation, the DNA was dissolved in 20 µl of TE buffer and separated by electrophoresis on a 1.0 percent low melting agarose gel. A ladder of ethidium bromide-stained linearized DNA bands, identified by size as insert (i), vector (v), v+i, v+2i, v+3i, and so on, was visualized by 360-nm ultraviolet light, and the v+i DNA was excised and recovered in 20 μl of TE. The v+i DNA was blunt-ended by T4 polymerase treatment for 5 minutes at 37°C in a reaction mixture (50 µl) containing the linearized v+i fragments four deoxynucleotide triphosphates (dNTPs) (500 µM each) and 9 units of T4 polymerase (New England BioLabs) under buffer conditions recommended by the supplier. After phenol extraction and ethanol precipitation, the repaired v+i linear pieces were dissolved in 20 μl of TE. The final ligation to produce circles was carried out in a 50- μ l reaction containing 5 μ l of v+i DNA and 5 units of T4 ligase at 14°C overnight. The reaction mixture was heated for 10 minutes at 70°C and stored at -20°C.
 - 16. A 100-μl portion of Epicurian Coli SURE 2 Supercompetent Cells (Stratagene 200152) was thawed on ice and transferred to a chilled Falcon 2059 tube on ice. A 1.7- μ l volume of 1.42 M β -mercaptoethanol was added to the cells to a final concentration of 25 mM. Cells were incubated on ice for 10 minutes. A 1-µI sample of the final ligation mix was added to the cells and incubated on ice for 30 minutes. The cells were heat-treated for 30 seconds at 42°C and placed back on ice for 2 minutes. The outgrowth period in liquid culture was omitted to minimize the preferential growth of any given transformed cell. Instead, the transformed cells were plated directly on a nutrient rich SOB plate containing a 5-ml bottom layer of SOB agar (1.5 percent SOB agar consisted of 20 g of tryptone, 5 g of yeast extract, 0.5 g of NaCl, and 1.5 percent Difco agar/liter). The 5-ml bottom layer was supplemented with 0.4 ml of ampicillin (50 mg/ml) per 100 ml of SOB agar. The 15-ml top layer of SOB agar was supplemented with 1 mt of X-gal (2 percent), 1 ml of MgCl₂ (1 M), and 1 ml of MgSO₄ (1 M) per 100 ml of SOB agar. The 15-ml top layer was poured just before plating. Our titer was approximately 100 colonies per 10-µl aliquot of transformation.
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 - For the unamplified λ library, *H. influenzae* Rd KW20 19. 20.
 - DNA (>100 kb) was partially digested in a reaction mixture (200 μ l) containing 50 μ g of DNA, 1× Sau3A

I buffer, and 20 units of Sau3A I for 6 minutes at 23°C. The digested DNA was extracted with phenol and fractionated on a 0.5 percent low melting agarose gel at 2 V/cm for 7 hours. Fragments from 15 to 25 kb were excised and recovered in a final volume of 6 μ l. We used 1 μ l of fragments with 1 μ l of DASHII vector (Strategene) in the recommended ligation reaction. One microliter of the ligation mixture was used per packaging reaction as recommended in the protocol with the Gigapack II XL Packaging Extract (Stratagene, 227711). Phage were plated directly without amplification from the packaging mixture (after dilution with 500 μl of recommended SM buffer and treatment with chloroform). [SM buffer contains (per liter) 5.8 g of NaCl, 2 g of MgSO₄ \cdot H₂O, 50 ml of 1 M tris-HCl, pH7.5, and 5 ml of a 2 percent solution of gelatin.] The yield was about 2.5×10^3 plaque forming units (PFU) per microliter. The amplified library was prepared essentially as above except the λ GEM-12 vector was used. After packaging, about 3.5×10^4 PFU were plated on the restrictive NM539 host. The lysate was harvested in 2 ml of SM buffer and stored frozen in 7 percent dimethyl sulfoxide. The phage titer was approximately 1 × 10⁹ PFU/ml. 21. M. D. Adams, *et al.*, *Nature* **368**, 474 (1994).

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- 25. Oligonucleotides were labeled by combining 50 pmol of each 20-mer and 250 mCi of $[\gamma^{-32}P]$ adenosine triphosphate and T4 polynucleotide kinase. The labeled oligonucleotides were purified with Sephadex G-25 superfine (Pharmacia). A portion containing 10⁷ counts per minute of each was used in a Southern hybridization analysis of H. influenzae Rd chromosomal DNA digested with one frequently cleaving endonuclease (Ase I) and five less-frequent ones (Bgl II, Eco RI, Pst I, Xba I, and Pvu II). The DNA from each digest was fractionated on a 0.7 percent agarose gel and transferred to nylon (Nytran Plus) membranes (Schleicher & Schuell). Hybridization was carried out for 16 hours at 40°C. To remove nonspecific signals, we sequentially washed each blot at room temperature with increasingly stringent conditions up to 0.1× saline sodium citrate and 0.5 percent SDS. Blots were exposed to a Phosphorimager cassette (Molecular Dynamics) for several hours; hybridization patterns were compared visually. S. Altschul et al., J. Mol. Biol. 215, 403 (1990)
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- 28. Standard amplification by polymerase chain reaction (PCR) was performed in the following manner. Each reaction (57 μ l) contained a 37- μ l mixture of 16.5 μ l of H_2O, 3 μ l of 25 mM MgCl_2, 8 μ l of a dNTP mix (1.25 mM each dNTP), 4.5 μ l of 10× PCR core buffer II (Perkin-Elmer N808-0009), and 25 ng of *H. influenzae* Rd KW20 genomic DNA. The appropriate two primers (4 μ l, 3.2 pmol/ μ l) were added to each reaction. A preliminary incubation (hotstart) was per-formed at 95°C for 5 minutes followed by a 75°C hold. During the holding period, Amplitaq DNA polymerase (Perkin-Elmer N801-0060, 0.3 μ l in 4.3 μ l of $\rm H_2O, 0.5~\mu l$ of 10× PCR core buffer II) was added to each reaction. The PCR profile was 25 cycles of 94°C for 45 seconds, then denature; 55°C for 1 minute, then aneal; 72°C for 3 minutes, then extension. All reactions were performed in a 96-well format on a Perkin-Elmer GeneAmp PCR System 9600. Long-range PCR was performed as follows: Each reaction contained a $35.2-\mu$ mixture of 12.0 μ of $\rm H_2O,\,2.2~\mu l$ of 25 mM magnesium acetate, 4 μl of a dNTP mixture (200 μM final concentration), 12.0 μl of 3.3× PCR buffer, and 25 ng of *H. influenzae* Rd KW20 genomic DNA. The appropriate two primers (5 μ l, 3.2 pmol/ μ l) were added to each reaction. A preliminary incubation (hot start) was performed at

94°C for 1 minute. Then rTth polymerase (Perkin Elmer N808-0180) (4 units per reaction) in 2.8 µl c 3.3× PCR buffer II was added to each reaction. The PCR profile was 18 cycles of 94°C for 15 seconds denature; 62°C for 8 minutes, anneal and extern followed by 12 cycles 94°C for 15 seconds, den ture; 62°C for 8 minutes (increase 15 per cycle) anneal and extend; and 72°C for 10 minutes, finextension. All reactions were done in a 96-well for mat on a Perkin-Elmer GeneAmp PCR Syster

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 - 47. Supported in part by a core grant from Hurran Ge nome Sciences and an American Cancer Society grant (NP-838C) (to H.O.S.). Reagents for sequence ing reactions and the synthesis of the oligonucleo tides were a gift from the Applied Biosystems Div sion of Perkin-Elmer. We thank T. Burcham of Ac plied Biosystems for his contribution in the cavelop ment of the TIGR EDITOR software; M. Riley. Marin Biological Laboratory, Woods Hole, for making he

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