Microbial Genomics and Chromosome Organization

Prokaryotic Chromosome Organization

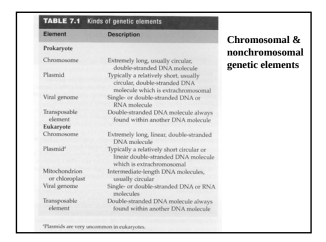
Generation of full genome sequences

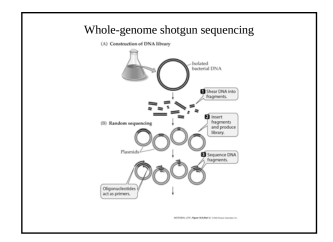
Genomic Structure & Functional Genomics

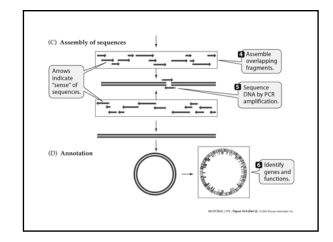
Genome size vs. No. of orfs

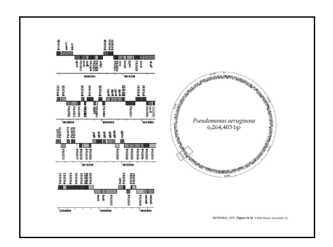
Minimal genome concept

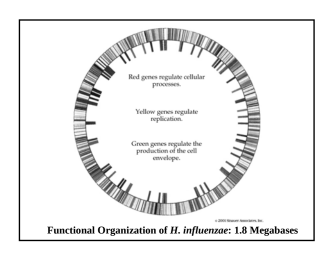
Lessons from full genomes

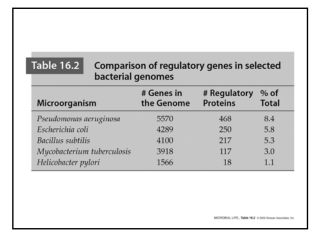


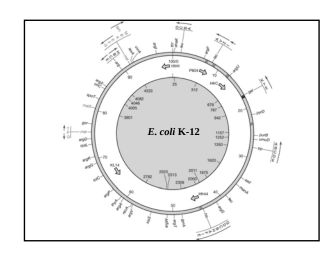












E. coli K-12 Genome Organization Lessons

Cured of F-plasmid and Lambda phage, 4.6 Mbs

88% of genome comprised in 4200 orfs
1% tRNAs and rRNAs
0.5% noncoding repetitive sequences
10% regulatory sequences, including promotors, operators, origin and terminus of DNA replication

70% single copy genes Some gene clusters, e.g., lac, trp, his, and NOT arg ~**6%** polycistronic mRNAs

E. coli K-12 Genome Organization Lessons (cont.)

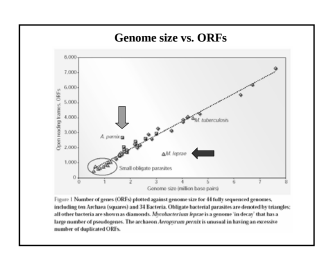
Gene orientation can be in both directions, however, highly expressed genes in the same direction as DNA replication forks travel.

For example, all 7 rrn operons Rem: transcription always 5' to 3'

Also found were several different cryptic, defective prophages and IS elements.

18% of genome from horizontal gene transfer, including large regions (~40Kb) known as islands of pathogenicity.

TABLE 15.2 Gene function in bacterial genomes							
	Percentage of genes on chromosome ^a in that category						
Functional categories	Escherichia coli	Haemophilus influenzae	Mycoplasma genitalium				
Metabolism	21.0	19.0	14.6				
Structural	5.5	4.7	3.6				
Transport	10.0	7.0	7.3				
Regulation	8.5	6.6	6.0				
Translation	4.5	8.0	21.6				
Transcription	1.3	1.5	2.6				
Replication	2.7	4.9	6.8				
Other, known	8.5	5.2	5.8				
Unknown	38.1	43.0	32.0				



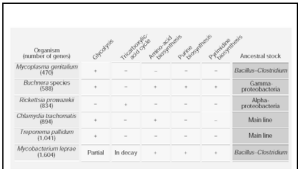


Figure 2 Many routes to intracellular adaptation. The differing presence (+) or absence (-) of $certain\ metabolic\ pathways\ in\ the\ streamlined\ genomes\ of\ parasitic\ bacteria\ shows\ how\ variable\ the$ process may be.

Global Transposon Mutagenesis and a Minimal Mycoplasma Genome

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Mycoplasma genālaim with 517 genes has the smallest gene complement of any independently replicating cell so far identified. Global transposon mu-tagenesis was used to identify nonessential genes in an effort to learn whether the naturally occurring gene complement is a true minimal genome under laboratory growth conditions. The positions of 2209 transposon insertions in the completely sequenced genomes of M. genitadiam and its close relative M. pneumoniae were determined by sequencing across the junction of the trans-poson and the genomic DNA. These junctions defined 1394 distinct sites of insertion that were not lethal. The analysis suggests that 265 to 350 of the 480 protein-coding genes of M. genitalium are essential under laboratory growth conditions, including about 100 genes of unknown function.

265 to 350 genes are the minimum necessary genome

The complete genome of the hyperthermophilic bacterium Aguifex aeolicus

Gerard Deckert**, Patrick V, Warren**, Terry Gaasterland; William G, Young*, Anna L, Lenox*, David E, Grahams, Ross Overboek; Marjory A, Snead*, Martin Keller*, Monette Auljay*, Robert Hubert, Robert A, Feldman*, Jay M. Short*, Gary J. Olsens & Ronald V. Swansoon*

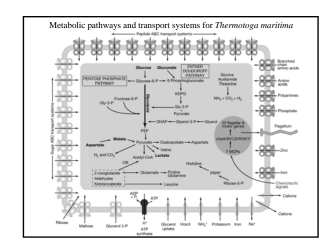
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Aquillar accident was one of the earliest diverging, and is one of the most therriophilic, bacteria known. It can grow on hydrog oxygen, carbon dioxide, and mineral salts. The complex metabolic machinery needed for A. accident to heroticon as a chemofilmountable hope in organism which sees an incorpance carbon source for his symbols and or incorpanic of deemical reservations are also as a constant of the immitted genome size. The uses of oxyging (altheit at very four concentrations) as an electron accept allowed by the presence of a complex respiratory appearates. Although this organism grows a 195° C, the extreme thermal lim the Bacteria, orly a few specific indications of thermophily are apparent from the genome. Here we describe the complete genome sequence of 15.51;35 bear pairs of this evolutionarity and physicologically interesting organism.



able 16.3 Distribution of genes of unknown function among selected bacterial genomes (Part 1)								
Organism Aeropyrum pernix K1	Genome Size (Mbp)	No. of ORFs (% coding)		Unknown Function		Unique ORF:		
		1,885	(89%)					
A. aeolicus VF5	1.50	1,749	(93%)	663	(44%)	407	(27%)	
A. fulgidus	2.18	2,437	(92%)	1,315	(54%)	641	(26%	
B. subtilis	4.20	4,779	(87%)	1,722	(42%)	1,053	(26%	
B. burgdorferi	1.44	1,738	(88%)	1,132	(65%)	682	(39%)	
Chlamydia pneumoniae AR39	1.23	1,134	(90%)	543	(48%)	262	(23%	
Chlamydia trachomatis MoP	1.07	936	(91%)	353	(38%)	77	(8%	
C. trachomatis serovar D	1.04	928	(92%)	290	(32%)	255	(29%)	
Deinococcus radiodurans	3.28	3,187	(91%)	1,715	(54%)	1,001	(31%	
E. coli K-12-MG1655	4.60	5,295	(88%)	1,632	(38%)	1,114	(26%	
H. influenzae	1.83	1,738	(88%)	595	(35%)	237	(14%	
H. pylori 26695	1.66	1,589	(91%)	744	(45%)	539	(33%)	
Methanobacterium thermotautotrophicum	1.75	2,008	(90%)	1,010	(54%)	496	(27%	

Organism Methanococcus jannaschii	Genome Size (Mbp)	No. of ORFs (% coding)		Unknown Function		Unique ORF	
		1,783	(87%)	1,076	(62%)	525	(30%
M. tuberculosis CSU#93	4.41	4,275	(92%)	1,521	(39%)	606	(15%)
M. genitalium	0.58	483	(91%)	173	(37%)	7	(2%
M. pneumoniae	0.81	680	(89%)	248	(37%)	67	(10%
N. meningitidis MC58	2.24	2,155	(83%)	856	(40%)	517	(24%
Pyrococcus horikoshii OT3	1.74	1,994	(91%)	589	(42%)	453	(22%
Rickettsia prowazekii Madrid E	1.11	878	(75%)	311	(37%)	209	(25%)
Synechocystis sp.	3.57	4,003	(87%)	2,384	(75%)	1,426	(45%)
T. maritma MSB8	1.86	1,879	(95%)	863	(46%)	373	(26%
T. pallidum	1.14	1,039	(93%)	461	(44%)	280	(27%)
Vibrio cholerae El Tor N1696	4.03	3,890	(88%)	1,806	(46%)	934	(24%)
Totals:	50.60	52,462	(89%)	22.358 (43%)		12,161	(23%)

Lessons from full genomes

Size range 600Kb to 12Mb

Vast number of putative genes with no known function.

Pathogenicity conferred by "Pathogenicity Islands" 44.5Kb in *Bacillus anthracis*

Symbiotic Island of >600Kb in *Sinorhizobium loti* including genes for nodulation and N-fixation

Adaptive gene losses in parasitic bacteria *Rickettsia* and *Chlamidia* are ATP thieves using the same "alien" ADP/ATP translocase

Microbial Genetic Exchange and Plasmids

Microbial Genetic Exchange is unidirectional!

Transformation Transduction Conjugation

Each requires Homologous Recombination

Types of plasmids

