thermore, while immunosubunit precursors-containing 15S proteasome assembly intermediates were detected in wild-type cells, under identical conditions 15S complexes were hardly detectable in PA28−−/− cells (Fig. 4B). Notably, PA28 was found to be associated with the immunosubunit precursor-containing 15S complexes, suggesting that PA28 is required for the incorporation of immunosubunits into proteasomes.

It has been demonstrated that immunosubunits are required for efficient antigen processing and that, in vitro, PA28 does not preferentially activate immunoproteasomes and does not alter proteasome substrate specificity (15). Thus, our findings strongly suggest that the inability to generate immune responses in PA28−−/− mice was a consequence of a decreased cellular level of immunoproteasomes. Taken together with the fact that, in PA28-expressing cells, immunoproteasomes have greater stability than proteasomes not containing immunosubunits (3, 12) and that PA28 interacts with the α-subunit ring of the proteasome (1), we hypothesize that, by inducing conformational changes of the proteasomal α-subunit ring (3, 16), PA28 promotes immunoproteasome assembly. PA28 might facilitate recruitment and assembly of the immunosubunit-containing proteasome β-subunit ring onto the α-subunit ring, which serves as a scaffold for the proteasome β-subunit ring formation and Upn1p-mediated maturation (17). As a consequence of increased cellular level of immunoproteasomes, MHC class I antigen processing and presentation are greatly enhanced by PA28.

**References and Notes**


9. Splenocytes were metabolically labeled for 60 min followed by a 60-min chase. NP-40 cell lysates were incubated at 31°, 37°, and 4°C for 45 min before being subjected to immunoprecipitation with anti-Kb or Dp. The amounts of class I molecules were quantified. It was found that while an equivalent amount of class I molecules was present in wild-type and PA28−−/− splenocytes before temperature challenge, the abundance of class I molecules in PA28−−/− cells after the temperature challenge decreased to ~80% of the wild-type levels.

10. A. Porgador, J. W. Yewdell, Y. Deng, J. R. Bennink, R. N. Germain, Immunity 6, 715 (1997). Ovalbumin-loaded LPS blasts from wild-type and PA28−−/− mice were stained with 25D1.16 and analyzed by flow cytometry. It was found that ~13% of the wild-type cells stained positive, whereas no staining of PA28−−/− cells was observed.


12. T. Preckel and Y. Yang, unpublished results.


18. Antisera specific to the proteasome subunit C9, LMP2, PA28α, and PA28β have been described (3). Antibodies specific to CD4, CD8, CD3-ε, TCR-αβ, CD23, CD25, CD28, CD45, and class I, and class II molecules were purchased from Pharmingen (San Diego, CA), 25D1.16 (10) and a PA28α-specific antiserum (2) were kindly provided by A. Porgador and M. Rechsteiner, respectively. Metabolic labeling, immunoprecipitation, immunoblotting, and sodium dodecyl sulfate–polyacrylamide gel electrophoresis (SDS-PAGE) were performed as described (3).


24. All animal experiments were conducted in accordance with institutional guidelines. We thank S. Sutton, J. Culver and S. Courtney for technical assistance and J.-F. Huang and G. Schoenhals for critical reading. The technical assistance of the DNA, vivarium, and peptide synthesis facilities of the R. W. Johnson Pharmaceutical Research Institute is gratefully acknowledged.

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**Global Transposon Mutagenesis and a Minimal Mycoplasma Genome**

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*Mycoplasma genitalium* with 517 genes is the smallest gene complement of any independently replicating cell so far identified. Global transposon mutagenesis 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. genitalium* and its close relative *M. pneumoniae* were determined by sequencing across the junction of the transposon and the genomic DNA. These junctions defined 1354 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.

One important question posed by the availability of complete genomic sequences (1–3) is how many genes are essential for cellular life. We are now in a position to approach this problem by rephrasing the question “What is life?” in genomic terms: “What is a minimal set of essential cellular genes?”

Interest in the minimal cellular genome predates genome sequencing [for a review, see (4)]. The smallest known cellular genome (5) is that of *Mycoplasma genitalium*, which is only 580 kb. This genome has been completely sequenced, and analysis of the se-
cludes orthologs of virtually every one of the 480 M. genitalium protein-coding genes, plus an additional 197 genes (8). There is a substantial evolutionary distance between orthologous genes in the two species, which share an average of only 65% amino acid sequence identity. The existence of these two species with overlapping gene content provided an experimental paradigm to test whether the 480 protein-coding genes shared between the species were already close to a minimal gene set. We applied transposon mutagenesis to these completely sequenced genomes, which permitted precise localization of insertion sites with respect to each of the coding sequences.

Populations of 200 to 1000 viable mycoplasma harboring independent transposon insertions were produced, and libraries of DNA fragments containing the junctions between the transposon and the chromosome were prepared and sequenced (9) (Table 1). Analysis of 2209 transposon junction fragments yielded 1354 different insertion sites. This data set is divided approximately equally between the two organisms. A total of 71% of the insertions were within genes in M. genitalium versus 61% in M. pneumoniae. This represents a substantial preference for intergenic insertion—because coding sequence constitutes 85% of the M. genitalium genome and 89% of the M. pneumoniae genome—and is consistent with the idea that intergenic sequences are less critical than protein-coding regions for viability. Transposon insertions have been identified in 140 different genes in M. genitalium and 179 different genes in M. pneumoniae.

The preference for insertion into the species-specific portion of the M. pneumoniae genome was striking (Fig. 1). The average density of distinct viable insertion events observed in M. pneumoniae–specific regions

Table 1. Summary of sequenced viable transposon insertion sites.

<table>
<thead>
<tr>
<th>Junctions sequenced*</th>
<th>1291</th>
<th>918</th>
</tr>
</thead>
<tbody>
<tr>
<td>Distinct sites†</td>
<td>685</td>
<td>669</td>
</tr>
<tr>
<td>Intergenic sites‡</td>
<td>199</td>
<td>261</td>
</tr>
<tr>
<td>Sites in genes§</td>
<td>484</td>
<td>408</td>
</tr>
<tr>
<td>Different genes‡</td>
<td>140</td>
<td>179</td>
</tr>
<tr>
<td>&quot;Disrupted&quot; genes¶</td>
<td>93</td>
<td>150</td>
</tr>
</tbody>
</table>

*Total numbers of interpretable sequences that include a junction between the end of the transposon and the mycoplasma genome sequence. †Numbers of unambiguously different positions in the genome sequence where a transposon junction was observed. The differences between these numbers represent multiple occurrences of identical insertion sites, which tended to cluster in certain genes, particularly in M. genitalium. ‡Annotated open reading frames and RNA genes in the two genomes, with respect to the total number of junctions sequenced. ¶Total number of annotated genes in which transposon insertions were observed. ‡Annotated open reading frames and RNA genes in the two genomes, with respect to the total number of junctions sequenced. ¶Total number of annotated genes in which transposon insertions were observed. See the text for a discussion of the criteria for counting "disrupted" genes.

Table 2. Estimating the total number of dispensable M. genitalium orthologs in M. pneumoniae. Among the 150 putative gene disruptions in M. pneumoniae, 57 are in genes that have orthologs in the M. genitalium genome, and 93 are in M. pneumoniae–specific genes. We have identified disruptive insertions in 47% (93/197) of the M. pneumoniae–specific genes. It is then a reasonable assumption that 47% of the dispensable M. pneumoniae genes common to both genomes have also been disrupted. This leads to an estimate that ~121 M. genitalium orthologs are dispensable and ~318 genes in total (121 + 197) are dispensable in M. pneumoniae. An analogous calculation using only those genes that have been hit more than once leads to an estimate of ~108 dispensable M. genitalium orthologs.

<table>
<thead>
<tr>
<th>Data from</th>
<th>Species-specific</th>
<th>Estimated total dispensable M. genitalium orthologs</th>
</tr>
</thead>
<tbody>
<tr>
<td>All genes disrupted in M. pneumoniae</td>
<td>93</td>
<td>121</td>
</tr>
<tr>
<td>Genes disrupted more than once in M. pneumoniae</td>
<td>42</td>
<td>108</td>
</tr>
</tbody>
</table>
large and suggests that this class of membrane proteins is important to the cell. Among the 19 genes encoding putative lipoproteins, we have identified potential disruptions in 13. There are several plausible interpretations of these seemingly incongruent facts, but perhaps the most likely is that the importance of these proteins is limited to fulfilling essential functions in the human host. This idea is substantiated by the occurrence of disruptions of several genes [M. genitalium/M. pneumoniae orthologs MG191 (MP014), MG192 (MP013), MG218 (MP527), and MG317 (MP388)] that are involved either directly or indirectly in mediating adherence to host cells (13).

The large group of genes with no functional assignment includes many genes with no known homologs outside of the mycoplasmas. As expected, many genes in this group have been disrupted (69 of 180). However, most of the 111 undisrupted genes of unknown function are apparently not dispensable and are expected to encode essential cellular functions.

Relatively few M. genitalium genes have functions related to biosynthesis and metabolism. This limited metabolic capacity has been compensated for by a proportionately greater dependence on transport of raw ma-

Fig. 1. Viable transposon insertions displayed on a composite M. pneumoniae–M. genitalium map. The M. pneumoniae genome is shown at a scale of 30 kb per line. Colored arrows above the line indicate annotated genes. Genes are colored according to their functional category, as indicated in the key. The genes are numbered sequentially from 1 to 677 as listed on Richard Herrmann’s Web page (http://mail.zmbh.uni-heidelberg.de/M_pneumoniae/genome/Sorted_Genes.html) and are referred to in the text as MP001 to MP677. Red triangles below the line indicate positions of transposon insertions documented in M. genitalium, mapped onto the M. pneumoniae genome (20). Red triangles above the line indicate positions of transposon insertions documented in M. pneumoniae. Regions of the genome that are M. pneumoniae–specific (absent from M. genitalium) are highlighted in pink directly on the line (20).
terials from the extracellular environment. Although the ability to generate nucleotides by salvage pathways has been retained, as have a limited number of biosynthetic and metabolic enzymes, it is evident that these pathways are nonessential in the laboratory, where the organism is apparently able to import nucleosides, amino acids, and other metabolites. Likewise, genes involved in the biosynthesis of cofactors [MG270 (MP450)], fatty acid and phospholipid metabolism [MG310 (MP395)], and heme conversion [MG118 (MP577)] appear to be dispensable.

Our data strongly support the idea that some metabolic pathways are essential. Glycolysis is thought to be the major source of adenosine triphosphate (ATP) and energy for *M. genitalium* and *M. pneumoniae*. We have not observed any disruptive insertions in any of the 10 genes involved in this pathway. Likewise, we have not identified any dispensable genes among the eight genes encoding ATP-proton-motive force interconversion activities.

ABC transporters are a heterotrimeric transport system made up of a specificity (ligand-binding) subunit, a permease, and an ATP-binding protein. ATP-binding subunits are distinct in that many appear to be “orphan” proteins, which are apparently overrepresented, compared to the other two subunits, in all genomes sequenced thus far. Likewise, there are specificity subunits with unknown partners within the *M. genitalium* genome, although their occurrence appears to be much more limited. Analysis of the *M. genitalium* genomic sequence data with less stringent searching parameters aimed at finding partners of the orphan specificity subunits led to the identification of potential transport partners (14). Because the sequence relatedness of these transporters was quite low, *M. genitalium* was thought to compensate for a reduced transporter spectrum by encoding transporters with broadened specificity (14). The current annotation of the *M. genitalium* genome lists 12 “orphan” ATP-binding proteins. We have obtained central insertions in only three of these genes [MG014 (MP136), MG390 (MP271), and MG467 (MP159)]. The fact that only 25% of the ATP-binding subunits in our data set tolerate insertions suggests that at least some of these orphan subunits do serve an essential function within the cell.

Two of the three subunits of an ABC phosphate transporter [MG410 (MP233) and MG411 (MP232)] have been putatively disrupted. Phosphate transport is thought to be an essential function. The insertion data for these two genes appear to be quite solid, in that MG410 (MP233) insertions were uncovered in both *M. genitalium* and *M. pneumoniae* and MG411 (MP232) contained multiple independent insertion events in *M. genitalium*. This finding forces us to consider the possibility that some as yet undefined transport system exists in these mycoplasmas that can compensate for mutations in the putative phosphate transporter.

Both *Mycoplasma* species examined in this study include two genes homologous to DNA pol III subunits [MG261 (MP460) and MG031 (MP120)]. We have found an insertion into MG261 (MP460), which supports the idea that it may function as a repair enzyme (3, 15) rather than being the main replication enzyme in the cell. The MG261 (MP460) insertion, together with insertions in the recA and uvrA excision repair genes, represents a special class of dispensable cellular functions. It is not necessarily surprising that cells can tolerate transposon insertions in these genes. It is almost certain that cells bearing such gene disruptions in nature would be quickly selected against. Although it is difficult to address this idea quantitatively, it poses a relevant question for consideration when attempting to define a minimal genome set for cellular life.

As expected, the number of transposon insertions recovered in genes involved in transcription was small. We have identified apparent disruptions in two of the five genes annotated as putative ATP-dependent RNA helicases [MG017 (MP134) and MG308 (MP397)]. The level of functional redundancy within this cellular role is somewhat unexpected and may reflect a broader role for these genes with regard to replication, repair, or transcriptional regulation.

A few insertions were in genes generally believed to be essential (12). Such events represent less than 1% of the total number of mapped insertion sites (1354). They include single putatively disruptive insertion events in two aminocycl RNA synthetase genes, the gene for ribosomal protein L28, the DNA replication genes dnaA and gidB, and a sigma factor gene. These unexpected findings forced us to consider explanations other than the dispensability of a function presumed to be essential. Functional assignments for some of these genes on the basis of sequence similarity may be incorrect. Also, some events that meet the criterion used in our analysis may not disrupt gene function. It is highly improbable that these events were recovered by cloning transposon junctions from nonviable cells. However, some cells might contain a functional duplicate copy of a gene in addition to the disrupted gene. It may also be that some functions can be supplied by unexpected uptake of enzymes or other compounds from the medium, or by cross-feeding. Conclusive proof of the dispensability of any specific gene requires cloning and detailed characterization of a pure population carrying the disrupted gene.

It is possible in some cases to verify the disruption of a specific gene function despite the presence of a transposon insertion mutant within a mixed population. For fructose-permease, insertions in both the *M. genitalium* gene (MG062) and the orthologous *M. pneumoniae* gene (MP077) were identified. We were able to detect the sequenced insertion events, using the polymerase chain reaction (PCR), in DNA from populations of cells grown in glucose-supplemented medium, but not when the medium was instead supplemented with fructose (Fig. 2).

The power of global transposon mutagenesis in our case benefited greatly from application to the fully sequenced and annotated genomes of the closely related *M. genitalium* and *M. pneumoniae*. Although *M. genitalium* may possess a unique genome, close to the minimal genome required for survival in its human host, it is clear from the results presented here that it contains a large number of genes that are dispensable under laboratory growth conditions. Our results imply that the 111 genes of unknown function that have not been disrupted in our experiments, the majority are essential. The presence of so many genes of unknown function among the essential genes of the simplest known cell suggests that all the basic molecular mechanisms underlying cellular life may not yet have been described. The essential gene set is not the same as the minimal genome. It is clear that genes that are individually dispensable may not be simultaneously dispensable. The data presented here suggest some specific experiments that could be carried out as a first step in the engineering of a cell with a minimal genome in the laboratory environment. One way to identify a minimal gene set for self-replicating life would be to create and test a cassette-based artificial chromosome, an experiment pending ethical review (16).

**Fig. 2.** Analysis of conditionally dispensable genes. PCR was done using one transposon-specific primer directed toward the chromosomal junction, and a gene-specific primer priming toward the particular insertion site to be detected (27). (A) Control experiment in which an insertion in a gene of unknown function [MG296 (MP417)] was detected in the same DNAs used in (B). (B) PCR primers were designed to detect the presence of an insertion in MG062 in pool C of *M. genitalium Tn4001* transformants. (C) PCR primers were designed to detect an insertion in the *M. pneumoniae* ortholog of MG062 (MP077) in pool E of *M. pneumoniae Tn4001* transformants. Lane labels G (glucose) and F (fructose) indicate the sugar used to supplement the growth medium (27).

### References and Notes


9. Transposon Tn4001, originally from Staphylococcus aureus, was propagated in Escherichia coli plasmid pSM2062 (17) and then introduced into M. genitalium and M. pneumoniae cells by electroporation (18). Approximately one in 10^5 to 10^6 M. pneumoniae cells and one in 10^7 to 10^8 M. genitalium cells were transformed to resistance to gentamycin (Gm). Cultures were split immediately after electroporation to generate eight separate populations for each species. Each population harboring cells representing –200 transposition events for M. genitalium and >1000 events for M. pneumoniae. These populations were allowed to recover in SP4 medium overnight, followed by growth in the presence of Gm for 2 to 3 weeks, resulting in the expansion of cell number by a factor of >10^6. This procedure was designed to make the subsequent cloning of transposition events from nonviable cells highly improbable. Genomic DNA was isolated from mid-log cultures; 2 μg of DNA was digested with Dra I. The genomic DNA restriction digests were diluted to 5 ng/μl and fragments were circularized using DNA ligase. Transpose junctions were amplified using inverse PCR with two primers specific for the end of the transposon Tn4001. Reaction products containing oligonucleotide-encoded Eco RI and Hind III sites were digested with these enzymes and cloned into the corresponding sites in the plasmid pUC18. DNA sequencing templates were prepared from selected colonies and sequences generated as described (2). Transpose junction sequences were aligned with the appropriate genomic sequence to establish the site in the genome of transposon insertion.

10. Tn4001 is a composite transposon with IS256 sequences at both ends. IS256 produces 8 and 9 base pair duplications of the target sequence at the insertion site [K. G. Dyke, S. Aubert, N. el Soh, Plasmid 28, 235 (1992); L. B. Rice, L. L. Carias, S. H. Marshall, Antimicrob. Agents Chemother. 39, 1147 (1995)].


12. Supplementary data can be found at Science Online (www.sciencemag.org/feature/data/1042937.shl).

13. The proteins involved in mycoplasma adhesion have been most extensively studied in M. pneumoniae. For a review see D. K. Krause, Mol. Microbiol. 19, 247 (1999).


20. The figure shows 600 M. genitalium sites that were unambiguously mapped onto the M. pneumoniae genome. The whole-genome alignment method of Delcher et al. (19) mapped 317 sites, but does not map sites close to insertion/deletion and rearrangement differences between the chromosomes. An additional 201 sites were mapped by searching the M. pneumoniae genome for matches to short sequences [200 to 400 base pairs] containing each M. genitalium insertion site. An additional 83 sites that were not mappable by the above methods (because of matches to several related M. pneumoniae sequences) were mapped to the corresponding position within the orthologous M. pneumoniae gene. The M. pneumoniae–specific regions are from (3), with minor modifications. Each pink highlighted region contains a block of M. pneumoniae–specific genes and arbitrarily includes the intergenic regions flanking it.

21. In the M. genitalium experiments, the growth medium is SP-4 supplemented with either glucose or fructose as indicated. In the M. pneumoniae experiment, the cells were grown in Hayflick’s medium supplemented with glucose or fructose. Cells were grown five passages (split 1 to 10) under the indicated condition before preparation of DNA for PCR templates. This is approximately equivalent to nine doublings.

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Functional Human Corneal Equivalents Constructed from Cell Lines

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Human corneal equivalents comprising the three main layers of the cornea (epithelium, stroma, and endothelium) were constructed. Each cellular layer was fabricated from immortalized human corneal cells that were screened for use on the basis of morphological, biochemical, and electrophysiological similarity to their natural counterparts. The resulting corneal equivalents mimicked human corneas in key physical and physiological functions, including morphology, biochemical marker expression, transparency, ion and fluid transport, and gene expression. Morphological and functional equivalents to human corneas that can be produced in vitro have immediate applications in toxicity and drug efficacy testing, and form the basis for future development of implantable tissues.

The cornea comprises three major cellular layers: an outermost stratified squamous epithelium, a stroma with keratocytes, and an innermost monolayer of specialized endothelial cells. The structure of the cornea allows it to serve as a barrier to the outside environment and as a major element in the optical pathway of the eye (1). The cornea is transparent, avascular, and immunologically privileged (2), making it an excellent candidate for tissue engineering for transplantation. Various researchers have attempted to fabricate artificial corneas or parts of corneas in vitro (3), but there have been no reports of successfully reconstructed human corneas that mimic the anatomy and physiology of the human cornea.

Our objective was to develop a morphological and functional equivalent of the human cornea. Human cell lines were developed from cells isolated from the individual cellular layers of the cornea. Most were immortalized (4) by infection with an amphotropic recombinant retrovirus containing HIV-16 genes E6 and E7 (5, 6); others were immortalized by transfection (7) with mammalian expression vectors containing genes encoding SV40 large T antigen, pSVneo (8), and adenovirus E1A 125S (9), separately or in combination. Immortalized cells had random chromosomal breaks, structural rearrangements, and, in several lines, aneuploidy (10); similar chromosomal anomalies associated with immortalization were reported in HPV E6/E7 immortalized vascular endothelial cells (6), although random structural rearrangements and aneuploidy are also present in normal human corneal cells (11). The immortalized cells also had significant telomerase activity (12) [associated with the immortalized phenotype (6)] compared to little or no activity in nonimmortalized cells.

Before use in corneal equivalents, cell lines were screened for morphological, biochemical, and electrophysiological similarities to freshly dissociated or low-passage corneal cells obtained from postmortem human corneas. Electrophysiological screening of epithelial cells by means of amphotericin-perforated patch clamp imaging (13) showed that immortalized cells had whole-cell currents similar to those of cultured corneal epithelial cells (Fig. 1, A to F) (14). Cells with altered phenotypes and physiology (transformed cells), however, showed anamolous currents (Fig. 1, G to I). Patch clamping was also used in screening keratocyte and