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Dissecting biological “dark matter” with single-cell genetic analysis of rare and uncultivated TM7 microbes from the human mouth

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We have developed a microfluidic device that allows the isolation and genome amplification of individual microbial cells, thereby enabling organism-level genomic analysis of complex microbial ecosystems without the need for culture. This device was used to perform a directed survey of the human subgingival crevice and to isolate bacteria having rod-like morphology. Several isolated microbes had a 16S rRNA sequence that placed them in candidate phylum TM7, which has no cultivated or sequenced members. Genome amplification from individual TM7 cells allowed us to sequence and assemble >1,000 genes, providing insight into the physiology of members of this phylum. This approach enables single-cell genetic analysis of any uncultivated minority member of a microbial community.

environmental microbiology | metagenomics | microfluidics | single-cell analysis

The earth contains enormous microbial diversity. Microbes colonize a wide variety of environmental niches, creating complex ecosystems and communities. Despite the marvelous progress in microbiology over the past century, we have only scratched the surface of this microbial world. It has been estimated that <1% of bacterial species have been axenically cultured, and fewer than half of the recognized bacterial phyla include cultivated representatives (1). This can be viewed as biology’s “dark matter” problem: just as astronomers can only indirectly infer the existence of a large amount of as-yet-undetected mass in the universe, microbiologists can only estimate microbial diversity by using techniques such as comparative 16S ribosomal RNA (rRNA) gene analysis (2), community DNA hybridization efficiency (3), and metagenomic gene inventories (4, 5). Although these techniques are useful, the cell, which is the ultimate unit of biological organization, is lost as a distinct informational entity.

Two general approaches have been used in addressing this problem. The first is to work on simple communities that contain only a few microbial species, in which case genome sequences can be reconstructed computationally after sequencing bulk DNA purified from the community (4). The second approach is to isolate individual cells by fluorescence-activated cell sorting, micromanipulation, or serial dilution, followed by genomic DNA amplification using techniques such as multiple-strand displacement amplification (MDA) (6, 7). The latter approach has been used successfully to perform genomic analysis of the cultivated and abundant marine bacterium *Prochlorococcus* MIT9312 (7). However, this approach remains difficult for two primary reasons: (i) the confidence needed to assert the presence of single cells in microliter volumes and (ii) the meticulous reagent cleaning and sample handling required to suppress background amplification in microliter-volume MDA (6). Those hurdles become even greater when complex environmental samples are

used. The number of species present requires substantial reagent consumption and expensive postamplification screening, and the probability of contamination is much higher because of the presence of free DNA. Neither approach has been validated with a complex ecosystem.

We have designed and fabricated a microfluidic chip to address these limitations. This device provides the ability to perform parallel isolation of single bacteria by steering them to any one of eight individually addressable chambers, followed by lysis and amplification of their individual genomes in 60-nl volumes. By using nanoliter volumes, the specific template concentration is increased by three orders of magnitude, as suggested previously (8, 9). To demonstrate the potential of this approach in microbial ecology, we performed a selective survey of microbes found in the human subgingival crevice, followed by whole-genome amplification (WGA) and high-throughput sequencing. The 16S rRNA gene-based phylogeny of several of these microbes placed them within the candidate phylum TM7, for which no cultivated or sequenced members exist (10), thereby providing unique genetic information about oral representatives of the TM7 phylum.

Results and Discussion

The microfluidic strategy for microbe isolation and genome amplification (Fig. 1) was validated on *Escherichia coli*. More than two dozen amplifications on single *E. coli* cells were performed, with a success rate of >90%. Subsequent PCR analysis of 10 genomic loci distributed over the *E. coli* chromosome showed that the amplification achieved excellent coverage and was able to amplify sequences with equal efficacy independent of their location on the genome [see supporting information (SI) Fig. 4]. Control experiments with only culture fluid in the chamber showed no significant amplification.

We then demonstrated the ability to select, isolate, and amplify the genomes of single bacteria from the human oral microbiota. The number of species in the human mouth is estimated to be ≈700 (11, 12). Because of the challenges of removing intact biofilm samples, rather than performing a comprehensive survey of this complex community our purpose

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Abbreviation: WGA, whole genome amplification.

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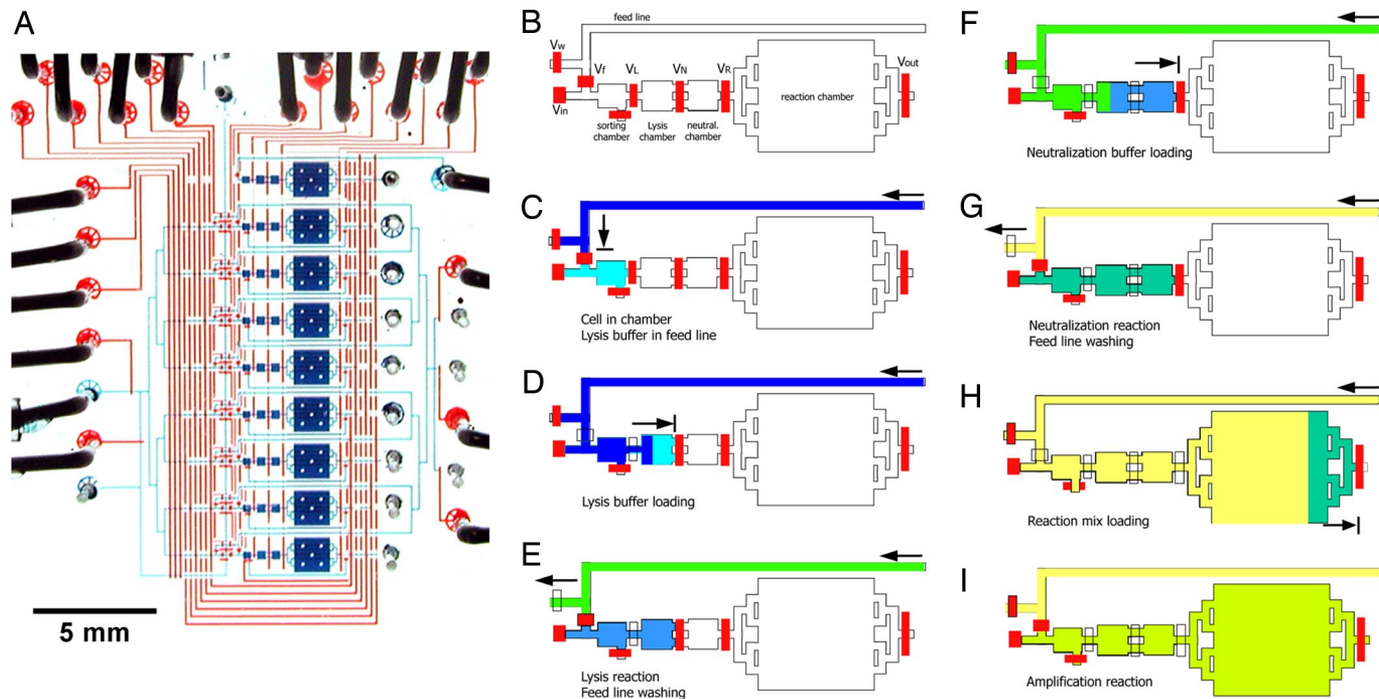


Fig. 1. Single-cell genome amplification device. (A) Photograph of a single-cell isolation and genome amplification chip capable of processing eight samples in parallel. To visualize the architecture, the channels and chambers have been filled with blue food coloring, and the control lines to actuate the valves have been filled with red food coloring. (Scale bar, 5 mm.) (B) Schematic diagram of a single amplification unit. The feed line is used to bring reagents into the chambers when the V_R valve is open and to the waste when the V_W valve is open. The V_{in} valve allows deposition of a single bacterium into the sorting chamber. The lysis (3.5 nl), neutralization (3.5 nl), and reaction chambers (50 nl) are used in sequence and are separated by individual valves V_L , V_N , and V_R , respectively. Valve V_{out} allows recovery of the amplified genomic material from the chip into an individual microfuge tube. (C) After a cell is trapped in the chamber, the feed line is filled with lysis buffer. (D) The lysis buffer is used to push the cell into the lysis chamber. (E) While the lysis buffer is mixing with the cell solution by diffusion, the feed line is flushed. (F) Neutralization buffer is loaded into the feed line and used to push the cell lysate into the neutralization chamber. (G) While the neutralization reaction is mixing by diffusion, the feed line is flushed. (H) The WGA reagents are loaded into the feed line and used to push the neutralized cell lysate into the reaction chamber. (I) The amplification reaction proceeds in a closed system comprising sorting, lysis, neutralization, and reaction chambers.

was instead to target an unexplored phylum and a relatively rare subset of the oral microbiota, TM7. By selecting microbial cells with a rod-like morphotype, we expected to enrich for the candidate phylum TM7 (13, 14). Little is known about the TM7 lineage. On the basis of comparative analyses of 16S rRNA genes, TM7 is one of a number of prominent “candidate” bacterial phyla lacking any cultivated representatives but comprising >50 phylotypes (1). rRNA gene sequences from the TM7 phylum have been found in a variety of habitats, ranging from deep sea hydrothermal vents to the healthy human mouth (10, 13, 14). In addition, sequence types within this phylum have been associated with chronic periodontitis in humans (13, 14). Fluorescence *in situ* hybridizations specific for TM7 showed that 0.7–1.9% of the subgingival microbiota belong to the TM7 phylum (13). A significant subset of this phylum has a peculiar morphology, characterized by long, thick filaments (up to $50 \times 4 \mu\text{m}$), making these cells good candidates for a morphotype-based selection (10, 13).

To identify the isolated rod-like cells, we performed PCR on the 16S rRNA gene from amplified genomic DNA, using primer sequences conserved across most species of the bacterial domain. Positive results for 16S rDNA PCR were obtained for 34 of 35 captured, single cells. After gel purification, 30 of these amplicons were directly sequenced; 28 gave unique sequences that were compared against the National Center for Biotechnology Information database by using BLAST (15). Fig. 2 shows a phylogenetic tree based on 16S rRNA gene sequences of most recognized bacterial phyla, with annotations for isolates from the present survey. The 28 sequences from this study are associated with five different bacterial phyla, with most se-

quences located in the phylum *Fusobacteria* and specifically related to the genus *Leptotrichia*.

We identified four members of the phylum TM7 from the amplified cells, of which three were closely related to a known oral TM7 clone (>99.6%; GenBank accession no. AY144355) (14) and a fourth clone related to a more distant lineage in the phylum (97.3%; AY134895) (16). To verify that the genome of a unique sequence type was amplified, the 16S rRNA amplicon of one TM7 sample (TM7a) was cloned, and 24 clones were sequenced; 23 of these had >99.5% sequence identity to the directly sequenced PCR product. To provide insight into the biology of the TM7 phylum and to investigate the ability to recover whole genome sequences from single uncultivated cells, we used the amplified genomic DNA from this sample for pyrosequencing and genome assembly. The resulting genome sequence data set was loaded into the Integrated Microbial Genomes with Microbiomes (IMG/M) database (17) to facilitate comparative analysis.

The assembly of TM7a genomic sequence resulted in the generation of 3,245 genes and gene fragments distributed across 1,825 scaffolds, totaling 2.86 megabases (Mb). Genome size estimates based on approaches such as the Lander–Waterman equation (18) or the characterization of known, conserved, single-copy genes (19) rely on random sampling of the genome. Single-cell amplification introduces a bias in read sampling such that we could not reliably estimate TM7 genome size. The assembly was fairly fragmented, with only 60% of the genes on multigene scaffolds. This suggested that there is multiple representation of some genes in the assembly and that the actual number of sampled genes in TM7a is somewhat smaller. If one

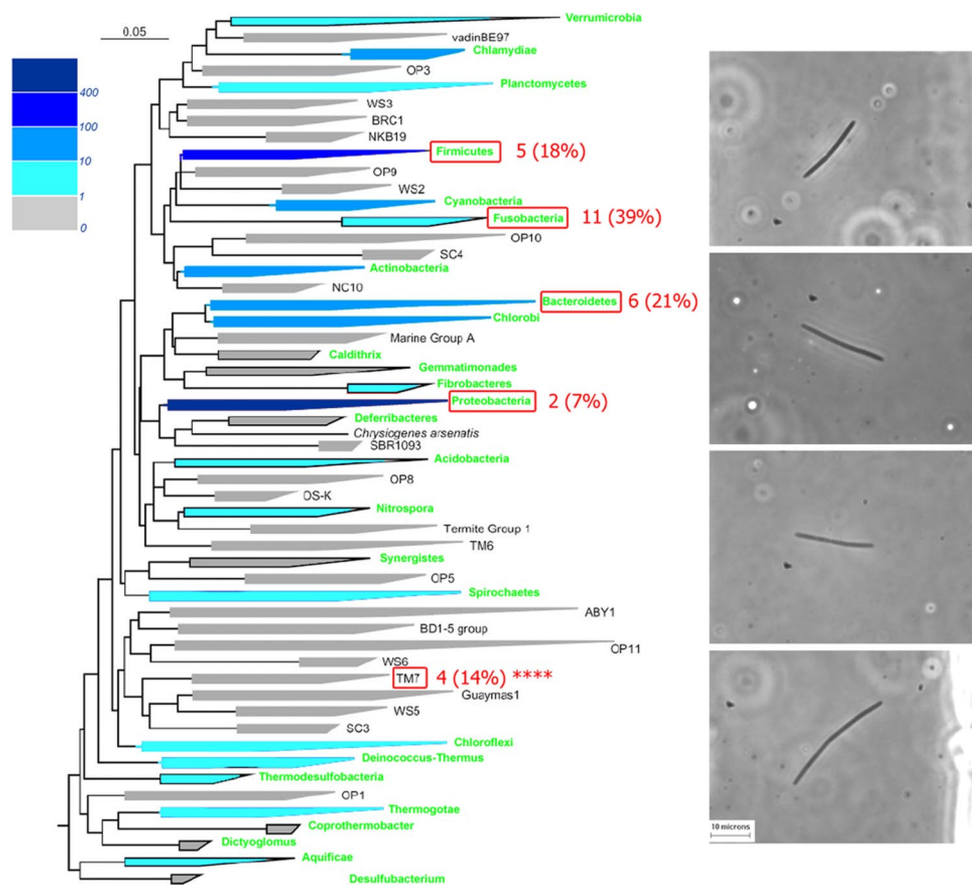


Fig. 2. Results of rod-like morphotype survey. (Left) Phylogenetic tree showing bacterial phyla based on 16S rRNA gene analysis (adapted from ref. 1). Green text indicates that at least one member of the phylum has been cultivated; different shades of blue indicate the number of genome sequencing projects in a particular phylum that were completed or in progress as of May 2006. Red numbers and percentages indicate the results of our single-cell survey of the human subgingival crevice, in which filamentous bacteria with rod-like morphotypes were isolated and lysed and their genomes were amplified. [Reprinted with permission from ref. 1 (Copyright 2003, Annual Reviews, www.annualreviews.org).] (Right) Optical micrographs of the four TM7 cells isolated in this survey.

applies a more conservative filter and only includes genes from large contigs (defined as those having three or more genes), then one is left with 1,474 genes on 288 scaffolds. This is probably a better estimate of the number of unique sampled genes in TM7a. Approximately 43% of genes were assigned a predicted function based on homology to published sequences, and 44% of the genes were mapped to clusters of orthologous groups (Table 1). We tested the validity of the assembly by choosing five regions of the genome with an average size of 1 kb. We designed PCR primers and successfully amplified all five regions from aliquots of the amplified TM7a genomic DNA (see SI Fig. 5).

Sequence similarity-based mapping showed that most of the TM7a genes are not closely related to genes from representatives of any known phyla. For example, 80% of the predicted TM7 proteins have <60% sequence identity to proteins from other sequenced organisms (Table 2). With this approach, a full third (33%) of the TM7 genes have <30% protein sequence identity to genes from any known phylum. This result is consistent with other cases of genome sequencing in previously uncharacterized phyla. For example, *Rhodospirella baltica* was the first sequenced representative of the *Planctomycetes* phylum, and 89% of its proteins have <60% identity to proteins from other known organisms; 20% have no matches with >30% identity. In contrast, a survey of 13 bacterial species in phyla with multiple sequenced representatives showed that, on average, only 15% of the proteins have <60% identity to proteins in other organisms, and 3% are unassigned at the 30% cutoff (see SI Table 3).

Although the majority of genes in the TM7a assembly are only distantly related to genes found in other organisms, a minority have a relatively high sequence similarity (>60% identity) to genes found in members of the classes *Bacilli*, *Clostridia*, or *Fusobacteria*. The presence of these genes may be the result of

Table 1. Statistics characterizing TM7a assembly and annotation, derived from IMG/M

Item	No.	% of total
DNA bases		
Total	2,864,887	100.0
Coding	1,160,954	40.5
G + C	981,862	34.3
DNA scaffolds	1,825	100.0
Genes		
Total	3,245	100.0
Protein coding	3,160	97.4
With function prediction	1,389	42.8
Without function prediction	1,771	54.6
Assigned to enzymes	530	16.3
Connected to KEGG pathways	400	12.3
Not connected to KEGG pathways	2,760	85.1
In clusters of orthologous groups	1,422	43.8
In protein families	1,221	37.6

KEGG, Kyoto Encyclopedia of Genes and Genomes (www.genome.jp/kegg).

salvage pathways (cysteine synthase and glycine hydroxymethyltransferase). We identified several genes coding for glycosyl hydrolase family enzymes distantly related to α -amylases and oligo-1,6-glucosidases, suggesting that oral TM7 cells may be capable of using oligosaccharides as growth substrates. Arginine is another potential growth substrate because of the presence of genes from the arginine deiminase pathway (arginine deiminase, ornithine carbamoyltransferase, and carbamate kinase). We also identified genes for ABC transporters that are likely responsible for oligopeptide uptake, suggesting that TM7 cells may be capable of using other amino acids as well.

It is an open question whether these bacteria have attributes associated with virulence and might be capable of contributing to oral disease. We noted the presence of genes for type IV pilus biosynthesis, including one similar to that which encodes the *Vibrio vulnificus* type IV pilin (21). Although type IV pili may facilitate the adherence of bacteria to epithelial cells, and contribute to biofilm formation, in Gram-positive cells, type IV pili have been shown to be responsible for an unusual communal form of gliding motility (22). TM7 cells from a sludge bioreactor appeared to have typical Gram-positive cell envelopes by electron microscopy (10). Therefore, if the TM7 are Gram-positive, their type IV pili may be involved in gliding motility.

We also investigated genes that might participate in cell envelope biosynthesis and found a gene predicted to encode a novel sortase, distantly related to those of *Firmicutes* and *Actinobacteria*, and a gene predicted to encode a UDP-*N*-acetylmuramyl tripeptide synthetase related to those of the bifidobacteria, suggesting a specific relationship of the TM7 cells to the Gram-positive lineages (see SI Fig. 6). Interestingly, in bifidobacteria the latter enzyme is predicted to add an atypical amino acid (ornithine or lysine instead of the more common diaminopimelate) to the growing peptidoglycan chain producing an A4 α / β type peptidoglycan. This peptidoglycan type has been implicated in chronic granulomatous inflammation (23) and may serve as a virulence factor for oral TM7.

In conclusion, we have isolated single bacterial cells from a complex human microbial community and sequenced their DNA to provide genetic insights into the TM7 phylum. The cell selection process described here used morphology as the basis for selection of the targeted bacteria. It would also have been possible to achieve the same results from an unbiased survey of the environmental sample; this simply would have required processing of a larger number of cells. Given that the cells were isolated from a complex bacterial biofilm (24) with no manipulation other than pipetting and dilution, many environmental microbial ecosystems should be amenable to this technique. We predict that, as genomes from the microbial dark matter are sampled by using techniques such as single-cell amplification, a much richer tapestry of microbial evolution will emerge.

Materials and Methods

Microfluidic Chip Fabrication. Microfluidic chips (Fig. 1) were fabricated as described previously (25), using the “push up” geometry with the following adjustments. The flow molds contained two layers: one for feeding lines and valves (SPR220; 7 μ m high) and one for the reaction chambers (SU8 2025; 25 μ m high). The control molds contained two layers: one layer for hydration channels under the reaction chamber (SU8 2015; 10 μ m high) and one for the control lines (SU8 2025; 25 μ m high).

Sample Collection and Isolation. Samples were collected from periodontal pockets by scraping subgingival tooth surfaces of a healthy individual (male, 40 years of age) after 5 days without tooth brushing. These biofilm specimens were dispersed, suspended, and washed twice in 1 \times PBS buffer and then resuspended in 1 \times PBS 0.2% Tween 20 before loading onto the chip. The chip was placed on an optical microscope, and the sample

was pumped through a sorting channel. When a single rod-shaped cell or a filament with the appropriate morphology (13) was visually detected in front of each processing unit, an isolation valve was closed and the cell was examined with a higher magnification. If the cell satisfied the selection criteria, the sorting valve was opened and the cell was pumped into the sorting chamber. Otherwise, the isolation valve was reopened and another cell was selected. This operation was repeated for seven processing units of the chip; the eighth unit was used for a negative control, having only suspension fluid inside. The chip also contains an independent processor with a separate, nonaddressable input that was filled with a mixture of lysed cells as a positive control. Every template chamber was then carefully checked for the number of bacterial cells, and a high-magnification image was recorded for every cell (Fig. 2). Of 42 processing units (six chips) used, 35 contained only one visible cell or filament.

Cell Lysis and WGA. Lysis, neutralization, and WGA were performed with the REPLI-g kit (Qiagen, Valencia, CA), using the recommended protocol except for on-chip WGA, for which the reaction mix was supplemented by 0.2% Tween 20 and one additional volume of polymerase. Once all of the chambers were loaded with cells, a 1-h-long lysozyme treatment was applied, using 1 \times PBS with 0.2% Tween 20 and 100 units/ μ l lysozyme (Epicentre, Madison, WI). This procedure was performed by taking advantage of the gas-permeability of polydimethylsiloxane to dead-end fill the feeding lines with the lysis buffer (Fig. 1C) and by opening the feeding valve to push the contents of the sorting chamber into the lysis chamber (Fig. 1D). Lysis and DNA denaturation reagents were allowed to incubate for 30 min. During this time, the feeding lines were washed first with air and then with the neutralization buffer (Fig. 1E). After completion of the lysis, the feeding valve was reopened, and neutralization buffer was pushed into the unit by dead-end filling of the neutralization chamber (Fig. 1F). After 15–20 min, the feeding line was washed again, this time with the WGA reaction mix (Fig. 1G). The feeding valve was reopened, and the reaction mix was used to dead-end fill the reaction chamber. With each WGA reaction isolated by closed valves, the chip was placed on a hotplate set at 32°C. The on-chip amplification took place for 10 to 16 h, after which samples were retrieved from the chip. The amount of amplified DNA after this step was estimated to be \approx 50 ng. A second, off-chip amplification was performed with the REPLI-g kit to obtain micrograms of DNA, the amount required for sequencing.

16S rRNA Gene Amplification, Cloning, and Sequencing. Gene PCR of 16S rRNA was performed on amplified genomic DNAs by using broad-range bacterial primers 8FM (5'-AGAGTTTGATCMTGGCTCAG-3'; adapted from ref. 26) and 1391R (5'-GACGGGCGGTGTGTRCA-3'; adapted from ref. 11). These primers amplify approximately >90% of the full-length bacterial 16S rRNA coding sequence. PCR mixtures were composed of 1 \times PCR buffer II (Applied Biosystems, Foster City, CA), 1.5 mM MgCl₂, 0.05% Triton X-100, 20 mM tetramethylammonium chloride, 0.1 mM of each deoxyribonucleoside triphosphate, 0.4 μ M of each primer, 2.5 units of AmpliTaq DNA polymerase (Applied Biosystems), and 1 μ l of amplified DNA in a final volume of 50 μ l. PCRs included 5 min at 95°C and 35 cycles of 30 sec at 94°C, 30 sec at 55°C, and 90 sec at 72°C, followed by 8 min at 72°C. PCRs were sequenced (Genaway, Hayward, CA) directly after purification from agarose gel by using the QIAquick gel extraction kit (Qiagen) or after cloning by using the TOPO-TA cloning kit (Invitrogen, Carlsbad, CA).

Genome Sequencing and Assembly. Pyrosequencing (454; Life Sciences, Branford, CT) was performed on randomly amplified

genomic material from three TM7 cells: TM7a, TM7b, and TM7c. Each sequencing run yielded 10–39 Mb of raw data composed of \approx 100-bp reads. The reads were assembled by using the 454 Newbler assembler and Forge whole-genome shotgun assembler (D. Platt, unpublished data). An initial assembly treating the coverage as a classic Poisson distribution indicated that the coverage of these genomes was quite uneven and that some regions were not joined because of either excess or very low coverage. The data were reassembled with Forge, using “metagenomic assumptions.” In this configuration, the assembler relaxes the Poisson depth assumption, which allows for much deeper coverage and exploration of low-coverage, less-certain overlaps between reads. All single-read, more highly error-prone contigs were excluded from the assembly. Genes were predicted on contigs greater than or equal in length to an average Sanger read (750 bp) by using fgenesb, as described previously (27), and then loaded into the IMG/M system (17) to facilitate comparative analysis.

Identification of Putative TM7 Genes. Putative TM7 genes were identified by comparing contigs and reads from the three TM7 data sets. Contigs >750 bp from cell TM7a or TM7b with a match (blastn, *e*-value $10e10$, low-complexity filter off) to one or more reads from a different cell (TM7b or TM7a, respectively)

were assigned to the TM7 metagenome if the match was >90% identity, had an alignment length >90% of the read length, and was at least 50 bp. This reciprocal comparison was also conducted on TM7b and TM7c. The rationale behind this binning is that TM7 cells with >99% 16S rRNA identity would be the only source of orthologs (between data sets) with >90% sequence identities, because contaminating exogenous DNA would presumably be randomly “sampled” from the oral microbiota. This strict identity threshold likely means numerous, more divergent TM7 orthologs would have been excluded. A total of 386 contigs with a combined length of 963 kbp were identified as putatively originating from TM7 genomes. These contigs encoded 850 ORFs, of which 481 could be assigned a putative function.

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