Generating a synthetic genome by whole genome assembly: \( \phi X174 \) bacteriophage from synthetic oligonucleotides

Hamilton O. Smith, Clyde A. Hutchison III†, Cynthia Pfannkoch, and J. Craig Venter‡

Institute for Biological Energy Alternatives, 1901 Research Boulevard, Suite 600, Rockville, MD 20850

Contributed by J. Craig Venter, November 3, 2003

We have improved upon the methodology and dramatically shortened the time required for accurate assembly of 5- to 6-kb segments of DNA from synthetic oligonucleotides. As a test of this methodology, we have established conditions for the rapid (14-day) assembly of the complete infectious genome of bacteriophage \( \phi X174 \) (5,386 bp) from a single pool of chemically synthesized oligonucleotides. The procedure involves three key steps: (i) gel purification of pooled oligonucleotides to reduce contamination with molecules of incorrect chain length, (ii) ligation of the oligonucleotides under stringent annealing conditions (55°C) to select against annealing of molecules with incorrect sequences, and (iii) assembly of ligation products into full-length genomes by polymerase cycling assembly, a nonexponential reaction in which each terminal oligonucleotide can be extended only once to produce a full-length molecule. We observed a discrete band of full-length assemblies upon gel analysis of the polymerase cycling assembly product, without any PCR amplification. PCR amplification was then used to obtain larger amounts of pure full-length genomes for circularization and infectivity measurements. The synthetic DNA had a lower infectivity than natural DNA, indicating approximately one lethal error per 500 bp. However, fully infectious \( \phi X174 \) virions were recovered after electroporation into Escherichia coli. Sequence analysis of several infectious isolates verified the accuracy of these synthetic genomes. One such isolate had exactly the intended sequence. We propose to assemble larger genomes by joining separately assembled 5- to 6-kb segments; \( \approx 60 \) such segments would be required for a minimal cellular genome.

Chemical synthesis of life in the laboratory has been a standing challenge to synthetic organic chemistry since Wöhler’s synthesis of urea in 1828 (1), and the doctrine of spontaneous generation was put to rest by an address by Louis Pasteur in 1864. With an understanding of the genetic role of DNA, much work has focused on the synthesis of oligonucleotides and genes. The synthesis of the 207-bp gene for tyrosine suppressor tRNA in 1979 by Khorana and 17 coworkers (2) was a monumental undertaking. Since then, the automated DNA synthesizer has been developed based on fundamental advances in synthetic methods from the laboratories of Letsinger (3, 4) and Caruthers (5, 6).

In 1999 we described a minimal prokaryotic genome based on results from random whole genome transposon mutagenesis that inactivated one gene per cell (7). By using this approach, \( \approx 300 \) essential genes for self-replicating cellular life were described, and we proposed to make a synthetic chromosome to test the viability of this hypothesis (7). Before attempting synthesis of a microbial chromosome, we commissioned an independent bioethical review of our proposed scientific plan (8). After \( \approx 1 \) year of deliberation, the reviewers concluded that we were taking a reasonable scientific approach to an important biological question. The broader implications of the creation of life in the laboratory can now be considered a realistic possibility. However, there are several technical barriers to the synthesis of microbial chromosome-sized stretches of DNA that are hundreds of thousands to millions of nucleotides long, the most notable being the contamination of the oligonucleotides by truncated species. Although such oligonucleotides are highly useful as primers for PCR amplification and DNA sequencing, only small (a few hundred base pairs) synthetic genes can generally be accurately and directly synthesized without multiple repair/selection steps. For example, the recent report (9) of the assembly of a partially active poliovirus from cloned synthetic segments of DNA from which polio genomic RNA (7,440 bases) could be transcribed was quite complex and took many months to accomplish. First, segments 400–600 bp long were individually assembled and cloned, and 5–15 isolates of each were sequenced to find one that was correct or readily correctable by oligonucleotide mutagenesis. These segments were then assembled into three larger segments of the polio genome, recombined, and finally assembled to produce a full-length product. This slow process would not be practical for synthesizing a 300,000-bp chromosome. We have now improved the methodology for synthesis of multigene segments of a genome as a step toward synthesis of a cellular genome. As a test of this methodology we have established conditions for global assembly of the infectious genome of bacteriophage \( \phi X174 \) (5,386 bp) from a single pool of chemically synthesized oligonucleotides. \( \phi X174 \) presents no known hazard because it infects only certain enteric bacteria and is not a human, plant, or animal pathogen. Therefore, its choice for synthesis serves to separate safety issues from ethical considerations and other potential risks associated with synthetic genomes.

\( \phi X174 \) (\( \phi X \)) is the prototypical minute icosahedral bacteriophage that was first characterized in the Sinseheimer laboratory beginning in the 1950s. The \( \phi X \) virion contains a circular single-stranded DNA that replicates through a double-stranded replicative form of DNA (RF). Its chromosome was the first DNA molecule purified to homogeneity (10), and, consequently, \( \phi X \) DNA has been used in many landmark experiments. It was the first viral DNA shown to be infectious in 1961, and in 1967 Goulian, Kornberg, and Sinsheimer (11) demonstrated that \( \phi X \) DNA synthesized with DNA polymerase, using the intact genome as a template, was infectious. This feat was hailed as “life in the test tube” (12). The \( \phi X174 \) genome was also the first DNA completely sequenced by Sanger et al. in 1978 (13). The sequence describes a remarkably compact gene organization with several cases of genes expressed by translating overlapping regions of the DNA in two different reading frames. Thus, \( \phi X \) provides a historic and demanding test bed for accurate synthetic chromosomes.

We set out to develop a procedure by which any 5- to 6-kb segment of DNA can be quickly and easily assembled from a

Abbreviations: RF, replicative form of DNA; PCA, polymerase cycling assembly; syn-\( \phi X \), synthetic \( \phi X174 \).

†Present address: Department of Microbiology and Immunology, University of North Carolina, Chapel Hill, NC 27599-7290.

‡To whom correspondence should be addressed. E-mail: jcventer@tcag.org.

single pool of synthetic oligonucleotides with high fidelity. In 1995, Stemmer et al. (14) used a polymerase cycling reaction to assemble a 2.7-kb plasmid from a single pool of oligonucleotides that was able to replicate in Escherichia coli. Synthesis of the larger ϕX genome, however, provides a more stringent test for optimizing oligonucleotide assembly methods. Its compact genetic organization makes it relatively intolerant to mutation compared with a plasmid, which may require only a replication origin and a drug resistance marker. We report here a procedure (Fig. 1) that utilizes sequential ligation and polymerase cycling reactions to accomplish the assembly of the ϕX genome in order of magnitude more rapidly than with previous methods (9) and with great savings in the sequence analysis of intermediate products and in human effort. This method should make it possible to assemble a minimal microbial genome from as few as 60 synthetic 5- to 6-kb segments.

Materials and Methods

Synthetic Oligonucleotides. Oligonucleotides were obtained from Integrated DNA Technologies (Coralville, IA) in 96-well format and normalized to 100 μM each. Based on the accompanying mass spectrometer data and information from the supplier, we ascertained that, on average, each 42-mer contained ~95% truncated species. To purify the oligonucleotides, 10 μl of each of the top-strand (or bottom-strand) oligonucleotides were pooled in two separate pools (top-1 to -130 and bottom-131 to -259), dried, and dissolved in 50 μl of water. Twenty microliters of concentrated pool plus 20 μl of formamide was heated to 95°C for 2 min, and 100 μl aliquots were loaded into 1.5-mm × 15-mm slots of a preparative 12% sequencing gel and electrophoresed at 1,300 V for ~4 h. The bands, which migrated close to the xylene cyanol marker, were visualized with a handheld 254-nm UV lamp and excised. The gel was extruded through a tuberculin syringe into 0.7 ml of TE buffer, frozen at −20°C overnight, eluted at 37°C on a rotating wheel for 1 h, and filtered through a glass, wool-plugged, 1-ml Eppendorf pipette tip. The recovered oligonucleotides were ethanol-precipitated and dissolved in 50 μl of water.

Phosphorylation of Oligonucleotides. The oligonucleotides were phosphorylated before ligation. A 100-μl reaction mixture containing 20 μl of purified top (or bottom) oligonucleotides, 10 μl of 10X T4 polynucleotide kinase buffer (NEB, Beverly, MA), 1 mM ATP, and 40 units of T4 polynucleotide kinase (NEB) was incubated at 37°C for 1 h. The reaction was terminated by phenol-chloroform extraction and ethanol precipitation. The phosphorylation reaction was repeated a second time. After extraction and precipitation, the oligonucleotides were dissolved in 50 μl of water.

Ligation Reactions. The 100-μl ligation reaction mixture contained 10 μl of top 5’P oligonucleotides, 10 μl of bottom 5’P oligonucleotides, 10 μl of 10X Taq ligation buffer (NEB), and 60 μl of water. The mixture was heated to 95°C for 2 min and slow-cooled over 30 min to 55°C. Ten microliters of Taq ligase (40 units/μl, NEB) was added, and incubation was continued at 55°C for 18 h. The reaction was terminated by phenol-chloroform extraction and ethanol precipitation. The products were dissolved in 90 μl of TE buffer.

Polymerase Cycling Assembly (PCA) of Oligonucleotides. PCA was carried out in reaction mixtures (50 μl) containing 5 μl of 10X Advantage 2 buffer, 5 μl of 10X dNTP mixture, 1 μl of 50X HF polymerase mixture (Clontech Advantage HF 2 PCR Kit, catalog no. K1914-1), and 0.2, 0.5, or 1 μl of the Taq ligation product. The polymerase mixture contains both an N-terminal deletion mutant of TaqDNA polymerase that lacks 5’-3’exonuclease activity and Deep Vent® polymerase (NEB) with 3’-exonuclease proofreading activity. Cycling parameters were 94°C for 15 sec, slow cool at −0.1°C/sec to 55°C, annealing at 55°C for 2 min, and extension at 72°C for 6 min. Thirty-five cycles were carried out followed by finishing at 72°C for 5 min.

PCR Amplification of Synthetic ϕX174 (synϕX) DNA Molecules Produced by PCA. The 25-μl PCR mixtures contained 2.5 μl of 10X Advantage 2 buffer, 2.5 μl of 10X dNTPs, 0.5 μl of HF polymerase mixture, 0.5 μl of each second-stage PCA product, and 1 μl each of 10 μM top-1 and bot-259 oligonucleotides (unpurified). PCR parameters were 94°C for 15 sec, 55°C for 30 sec, and 72°C for 6 min for 25 cycles, finishing with 72°C for 5 min. The PCR mixtures were pooled, phenol-chloroform-extracted, ethanol-precipitated, and redissolved in 10 μl of TE buffer.

Conversion of Linear synϕX DNA Molecules to Infectious Circular Molecules. The pooled PCR-amplified synϕX DNA was cleaved with PcrI, and the linear DNA was gel-purified to yield ~1 μg of linear synϕX. The linear synϕX molecules were circularized by ligation under dilute conditions (~1 μg/ml) with T4 ligase.
DNA Sequencing. Plaques were picked directly into 50-μl PCRs with a 10-μl micropipette tip and subjected to 30 cycles of amplification consisting of 10 sec at 94°C, 30 sec at 55°C, and 6 min at 72°C. Fifty microliters of a mixture of 40 μl of shrimp alkaline phosphatase (1 unit/μl), 8 μl of exonuclease I (20 units/μl), and 752 μl of water added to each PCR mixture, and digestion was at 37°C for 45 min and 45°C for 15 min, followed by 72°C for 15 min to inactivate the enzymes. The syndX DNA was gel-purified before carrying out standard sequencing reactions and sequencing on a 3730 XL sequencer (Applied Biosystems). Both strands were sequenced, and depth was generally >2-fold.

Rationale and Theoretical Considerations

The steps we used to accomplish the synthesis of infectious φX double-stranded RF DNA from a single pool of oligonucleotides are summarized in Fig. 1. In developing this strategy, we considered how impurities present in oligonucleotide preparations can result in assembly errors and mutations in the final product. Thus, we were led to purify the oligonucleotides before assembly. We also analyzed and determined the theoretical endpoints and limitations of the two basic assembly steps: ligation and PCA.

Oligonucleotide Purity. If automated DNA synthesizers produced pure oligonucleotides of the programmed sequence, then assembly of long double-stranded DNA molecules would be straightforward. In reality, only ~50% of the molecules in preparations such as those used in our work have the correct chain length. The population of other molecules includes both truncated species capped at the growing end and uncapped molecules containing errors (mostly deletions). These incorrect molecules will either block assembly of the oligonucleotides or result in mutations in the assembled DNA. Because, on average, one of every two molecules is correct, the probability that a strand of our φX genome would completely assemble correctly by random selection from 130 unpurified oligonucleotides is (1/2)^130 or ~10^-39. We estimated that, even with selection for infectivity, it was essential to reduce the number of incorrect oligonucleotides to ~10% to allow detection of correct molecules. Because purification of each oligonucleotide would be time-consuming and laborious, and because all our synthetic oligonucleotides were of equal chain length, we chose to gel-purify pooled oligonucleotides. We pooled the nucleotides from each strand in two separate pools to minimize the likelihood of annealed structures that could interfere with gel purification on the basis of chain length.

Ligation. We chose Taq ligation as the first step in our φX assembly, because ligation under stringent annealing conditions (55°C) would diminish the possibility of incorrect pairing and might also select against ligation of oligonucleotides containing mutations. In principle, it should have been possible to obtain full-length φX genomes by ligation. A major reason why this was not attained is that the concentrations of all of the oligonucleotides in the ligation mixture are not equal. Growing assemblies terminate prematurely when particular oligonucleotides become exhausted. Fig. 2 shows the results of a computer simulation of the ligation reaction. Both the fraction of full-length product and the average chain length of assemblies drop rapidly as the ligation reaction. Both the fraction of full-length product and the average chain length of assemblies drop rapidly as the

PCA. The PCA reaction is a thermocycling polymerase reaction, similar to a PCR but without a pair of primers present in excess compared with template (14). At each cycle, DNA is melted and overlapping single strands reanneal. If the 3' ends of reannealed strands are such that they can be extended by using the opposite strand as template, then polymerase extends the strands to form duplex molecules. DNA strands continue to elongate at each cycle until they either are full-length or can no longer be
The synthetic genome was extended. It should be noted that PCA is not an amplification reaction. Only limited total synthesis can occur, and, because the polymerase mixture we used contains no 5′ exonuclease, no DNA is degraded. It is difficult to analyze the kinetics of the process; however, the final end products are simple to describe (Fig. 2B). Molecules present in the starting mixture can be extended in the 5′–3′ direction to the end of the genome just once, over a number of cycles. Only the molecules containing the 5′ ends of each strand can be extended to full length. The total increase in mass of the DNA is limited to \( \sim \frac{3n}{2} \), and the fraction of final mass that can achieve full length is \( \sim \frac{2}{n} \), where \( n \) is the number of oligonucleotides on one strand (Fig. 2B). We assumed it would be unlikely that either the ligation or the PCA reactions are a major source of errors, because no synthesis occurs in the former and synthesis is limited in the latter (although some errors could occur by deamination or depurination during these steps). Most errors can be attributed to incorrect synthesis of the oligonucleotides. If our oligonucleotides are 90% pure, then each synthetic chain would, on average, contain \( \sim 13 \) mutations. The fraction of correct syn\( \text{dX} \) molecules would be \( (0.9)^{130} \) or \( \sim 10^{-6} \).

**Results**

The oligonucleotides were designed to synthesize a \( \text{dX} \) genome with exactly the sequence reported by Sanger et al. in 1978 (13) [several database entries exist with exactly this sequence, NC_001422 (GenBank database), J02482, and V01128]. In designing the oligonucleotide set, we adopted the strategy of appending sequence to either end of the \( \text{dX} \) sequence to make a slightly larger molecule that could be cleaved to size with \( \Phi 29 \) and then circularized to produce infectious DNA. We appended 5′-TAACGCTTGCA to the left end and one G plus a randomly generated 73-nucleotide sequence to the right end, thereby restoring a \( \Phi 29 \) site at each end. Starting at the left end, 130 oligonucleotides (top-1 to top-130), each 42 bases long, were consecutively generated. Similarly, starting on the bottom strand at position 22, oligonucleotides numbered bot-131 to bot-259 were generated (see Fig. 1, upper left). These oligonucleotides were pooled and gel-purified as described in Materials and Methods.

We phosphorylated the oligonucleotides and mixed the top and bottom pools. To improve the ligation fidelity we used \( \text{Taq} \) ligase at 55°C for 18 h. A sample of the ligation reaction was analyzed on an Invitrogen 2% E-gel (Fig. 3E). The average size of the double-stranded products was \( \sim 700 \) bp, with a small fraction extending to 2 or 3 kb (Fig. 3E, lane N). Fig. 3E, lane D, shows a sample denatured in formamide before loading on the gel. The single-stranded products ranged in size, with the largest being \( \sim 1 \) kb.

To obtain full-length \( \text{dX} \) DNA molecules, we diluted 0.2-, 0.5-, and 1-μl samples of the ligation product to 50 μl and subjected them to 35 cycles of PCR as shown in Fig. 3A. With the 0.2- and 0.5-μl samples, single-stranded assemblies approaching full length or nearly full length were produced, whereas assemblies from the 1-μl sample were shorter. PCR of ligation samples exceeding 1 μl yielded even shorter PCR products (data not shown). Apparently, dilute conditions favor the annealing of only two partially overlapping strands at a time followed by fill-in synthesis, whereas concentrated conditions favor multibranchanded, annealed structures that interfere with fill-in synthesis.

To improve the yield of full-length \( \text{dX} \), diluted samples of the first stage of PCR assembly were subjected to an additional 35 cycles of PCR. A small fraction of the products was now visible as full-length single strands (Fig. 3B) and as full-length duplexes (Fig. 3C). The presence of products of size apparently greater than full-length suggests either that some incorrect assemblies had accumulated during the later PCR cycles or that branched structures might have formed by reannealing either during electrophoresis (Fig. 3B) or during the final cycle of PCR (Fig. 3C).

After viewing the intensity of the bands and realizing that only 1/25th of the PCR reaction mixture was analyzed on the gel, we estimated that several nanograms of full-length product were produced. Because 1 ng of \( \text{dX} \) RF is \( \sim 1.7 \times 10^{9} \) molecules, we must have generated \( >10^{10} \) independent syn\( \text{dX} \) molecules, and we can infer that some of these would have the correct sequence.
A small sample (0.5 μl) of each second-stage PCA reaction was amplified by PCR by using top-1 and bot-259 oligonucleotides as primers (Fig. 3D). The PCR products were pooled and cleaved with PstI, and the linear DNA was gel-purified to yield ~1 μg of linear synφX. The linear synφX molecules were circularized by ligation under dilute conditions (<1 μg/ml) in preparation for infectivity testing. Gel analysis showed that ~50% of the linear synφX were converted to circular molecules (data not shown).

One microliter of synφX ligation product (an estimated 3–5 ng of circular molecules based on ethidium bromide staining intensity) was electroporated into DH10B cells (Invitrogen), divided into two aliquots (A and B), and plated for infectious particles as described in Materials and Methods. The synφX-A lysate yielded 194 plaques (Fig. 4), and the synφX-B lysate yielded 181 plaques per 100 μl plated. Some variation in plaque size was observed. The plaque yield from 3 to 5 ng of the synφX product was 2 × 10^4 of the yield from 1 ng of commercial NEB RF DNA; therefore, we estimate that the synφX product is ~5 × 10^3 of that for natural φX, leading us to conclude that there are 9–10 inactivating mutations in the synthetic genome. This is in reasonable agreement with our estimate of 90% purity for the gel-purified oligonucleotides.

Several plaques were picked from each plate directly into PCR mixtures for amplification and sequencing. The A plaques are independent of the B plaques, but individual plaques from A, or from B, are not necessarily independent, because they may have arisen from the same infected cell. Representative sequences from four plaques were compared in Fig. 5. Phage DNA from plaque B3 gave a sequence that was identical to GenBank accession no. J02482. The B1 DNA sequence contained one silent T → C transition, two silent G → A transitions, one G → A transition at position 4170 (resulting in a Gly-to-Ser amino acid change in the gene H protein), and one T → G transition at position 3606 (resulting in a Ser-to-Ala change in the gene H protein). DNA from plaque A4 contained a T → C silent mutation at position 1045 and A8 contained a G → A silent mutation at position 446, a C → T mutation at position 4399 (resulting in an Ala→Val amino acid change in gene A protein), and a C → A change at position 5144 (resulting in a Gln-to-Lys change in gene B and silent changes in genes A and A*). Infectious titers of synthetic phage from plaque B3 were indistinguishable from commercially available phage.

**Discussion**

Synthesis of a molecule with the same properties as a naturally occurring compound has traditionally been used as evidence for correctness of a proposed molecular structure. Therefore, it is interesting to consider whether synthesis of an infectious φX genome proves the correctness of the Sanger sequence (13) on which our synthesis was based. Clearly, the sequence is accurate enough to specify an infectious phage. One synφX isolate (B3) with wild-type plaque morphology is identical in sequence to the Sanger sequence. In a study of convergent evolution using φX, Bull et al. (15) determined that the wild-type φX used in their experiments differed at five positions from the Sanger sequence (13). In the present work we have also sequenced four different preparations of natural φX. It is interesting that none of these sequences is identical to the Sanger sequence (Fig. 6), differing from it by three to five single base substitutions in addition to the single base change associated with the am3 mutation, which is present in our natural DNA preparations but absent from the Sanger sequence as presented in the GenBank database (13). SynφX-A4, A8, and B3 are consequently closer to the Sanger sequence (13) than any of the natural φX DNA preparations (10) we have sequenced. It should be noted that the commercial φX DNA present in the laboratory while synthesis was ongoing differs at five positions (including the am3 site) from the Sanger sequence. None of those differences are found in our synφX sequences, demonstrating that this isolate is synthetic rather than

**Fig. 4.** Plaques of synφX-A. There appear to be several plaque morphologies: small plaques with sharp borders, medium-sized plaques, and large plaques with fuzzy borders.
a contaminant. The natural sequences closest to the Sanger sequence are from DNA preparations contemporaaneous with the original sequencing (Fig. 6). Our sequences were determined by using template DNA prepared in the late 1970s from dX brought back from a sabbatical in the Sanger laboratory by one of the authors (C.A.H.). It is difficult to conclude whether any of the three differences between the Sanger sequence and contemporaneous dX preparations represents sequencing errors. Only two of these differences are shared by all four of the natural dX sequences and the sequence of Bull et al. (15), and these are both silent changes. syndX-B3 is identical to the Sanger sequence (13) at both positions and is fully infectious. The quality of a synthetic genome sequence depends in part on the accuracy of the original DNA sequence from which it is derived. It is truly remarkable how well the first sequence of a DNA genome holds up to close scrutiny a quarter of a century later.

We have considered the source of mutations limiting the accuracy of assembled synthetic DNA. The predominant impurities in oligonucleotide preparations are molecules shorter than the desired product that are expected to result in deletions after assembly. In our experiments, selection for viable dX strongly selects against such framework mutations, and none were seen in the viable syndX isolates. The observed mutations are all single-base substitutions, predominantly transitions [changes from one pyrimidine to the other (C→T) or from one purine to the other (A→G)]. The two obvious sources of these substitutions are enzymatic replication errors and preexisting base substitutions in the synthetic oligonucleotide preparations. The mutation frequency expected from enzymatic errors is simply due to the 25 cycles of PCR carried out before circularization of the genome. PCA reactions extend molecules present in the original ligation mixture to full length only a single time, and so are not expected to contribute significantly to the mutation frequency. The error rate for PCR using Taq polymerase has been measured to be \(10^{-5}\) (mutation frequency/bp/duplication) and the HP polymerase mixture used in these experiments is reported to have significantly higher fidelity. This leads us to estimate an average of approximately one PCR-induced mutation per genome. The observed rate appears to be somewhat higher, leading us to speculate about the possible origin of base substitutions in the oligonucleotide preparations.

The predominant transition type observed (G.C to A.T, which is \(G\)→C changes) could arise by deamination of C to produce U. PCR amplification would then copy U in DNA as T to produce normal DNA that is insensitive to the uracil N-glycosylase of E. coli. The phosphoramidite of U might exist as a contaminant of the C phosphoramidite used in oligonucleotide synthesis, or deamination might occur at any stage before PCR amplification of the full-length dX genome. We have demonstrated the rapid, accurate synthesis of a large DNA molecule based only on its published genetic code. The accuracy of our final product was demonstrated by DNA sequencing and phage infectivity. Our methods will permit serial synthesis of gene cassettes containing four to seven genes in a highly robust manner. However, without selectivity, these cassettes will contain mutations (≈2 per kb) derived from errors contained in the oligonucleotide pool. Thus, without error correction, they would currently be unsuitable for assembly into a chromosome for an entire organism. Selection for infectivity, such as we have used, or for ORFs in single genes provides advantages for synthesizing viruses or short sequences. However, when our method of synthesis is coupled with DNA sequencing and repair by site-directed mutagenesis, it will enable rapid production of accurate cassettes that can be assembled into larger genomes. The capabilities of DNA synthesis have lagged far behind our ability to determine sequences during the past 30 years. If this gap can be closed, then limitless possibilities for the application of synthetic methods to the study and practical application of genomics will emerge. There are many reasons to synthesize DNA chemically, rather than clone natural sequences, one of which is to prove correctness (or incorrectness) of a sequence, because many published sequences contain errors. The natural DNA may be unavailable to the experimenter for various reasons, including an uncooperative laboratory, an environmental sample that has been used up, an archaeological sample in short supply, or a sequence from badly degraded DNA or from an extinct organism. Also, the sequence could be deduced (rather than experimental) from an ancestral sequence, a designer protein, or a fusion of domains from different proteins. There may be a hazard associated with the source of the natural sequence, or the target sequence may be RNA or a protein sequence rather than a DNA sequence. The sequence may need to be reengineered to alter the codon usage (or the code) for a particular host; to alter closely spaced regulatory signals or protein initiation (ribosome binding sites), promoters, or transcription terminators; to introduce restriction sites; or to allow convenient construction of a family of related (but different) constructs. Synthesis may become the easiest way to get a sequence as methods are refined. A desired construct may require the assembly of many pieces from different sources. Synthesis obviates the need to develop a special strategy for each construct, providing complete flexibility of design. The combination of improved oligonucleotide synthesis combined with the methods described here will enable rapid, accurate synthesis of genomes of self-replicating organisms that will serve as a basis for understanding minimal cellular life. Synthetic genomics will become commonplace and will provide the potential for a vast array of new and complex chemistries altering our approaches to production of energy, pharmaceuticals, and textiles.

This work was supported by grants from the U.S. Department of Energy and the J. Craig Venter Science Foundation.