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Synthetic genomics: a new venture to dissect genome fundamentals and engineer new functions

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Since the first synthetic gene was synthesized in 1970s, the efficiency and the capacity of made-to-order DNA sequence synthesis has increased by several orders of magnitude. Advances in DNA synthesis and assembly over the past years has resulted in a steep drop in price for custom made DNA. Similar effects were observed in DNA sequencing technologies which underpin DNA-reading projects. Today, synthetic DNA sequences with more than 10 000 bps and turn-around times of a few weeks are commercially available. This enables researchers to perform large-scale projects to write synthetic chromosomes and characterize their functionalities in vivo. Synthetic genomics opens up new paradigms to study the genome fundamentals and engineer novel biological functions.

Introduction

One of the major challenges in biological sciences was the determination of DNA sequences. In the beginning, only single DNA fragments were sequenced using the chain termination sequencing technique [1]. However, the Human Genome Project (GP-Read) accelerated the evolution of new sequencing techniques by having the ambitious goal to sequence the human genome within 15 years. The development of Next Generation Sequencing techniques today allows sequencing of a human genome within days. However, most eukaryotic genomes are not fully sequenced and new sequencing techniques are still being developed. As exemplary achievement of this development, in 2017 sequencing of one of the highly repetitive human centromeres was achieved [2]. Scientists are now performing well in reading genomes, a measurable output being the growing number of genome sequences in public databases. However, reading a book alone does not make a good writer, instead it requires one to start writing extensively and creatively to master the art and ultimately it leads to a better understanding of grammar and expression. In this case, one needs to write synthetic DNA sequences in order to better understand the grammar of life.

Writing DNA starts with short single-stranded fragments: the oligonucleotides. Since the development of the Polymerase Chain Reaction and the first complete synthesis of a gene, writing DNA in vitro has progressed impressively (Figure 1) [3,4]. Recent drops in DNA synthesis costs and the improved capability of synthesizing longer stretches of DNA allow the design and construction of whole synthetic chromosomes in the mega-base range. Recent publications report the construction of viral and microbial synthetic genomes, and the Sc2.0 project aims to generate the first synthetic eukaryotic genome. It is an open discussion how to define whether a chromosome or genome is synthetic. In this review, chromosomes and genomes are defined as synthetic when all building blocks of the final DNA molecule are generated by chemical synthesis. Chromosomes and genomes which are not completely synthesized are considered ‘engineered’ or ‘modified’ and are outside the scope of this review. We define synthetic genomics to be a new field where biology is being engineered at the genome level, and it is an intersection of synthetic biology and systems biology. This review neither aims to discuss assembly methods nor the dual-use character of synthetic genomics. The authors are fully aware of the potential dual use character, especially for the synthesis of viral genomes. However, these issues are discussed and reviewed extensively elsewhere [5–7].

Design concepts and assembly strategies for synthetic chromosomes

Computer-assisted design software (CADs) have been developed to ensure efficient and consistent design of synthetic DNA sequences at the genome scale [8**,9]. The design space of synthetic DNA is enormous and many (if not infinite) design blueprints are possible, as long as they can result in the viability of the cell, to achieve the design intention [10]. Initial projects aiming to synthesize a whole genome were conservative in
changes to the genetic content, but nonetheless resulted in the breakthrough in synthesizing, assembling and ultimately transplanting chromosome-scale synthetic DNA [11,12]. With increasing knowledge and progress in chromosome-scale DNA synthesis, the designs of synthetic sequences are becoming more complex and ambitious [8,13]. Many genome synthesis projects utilize a hierarchical genome assembly strategy starting with small building blocks which are assembled, by the technique of choice, to larger building blocks of around 50–100 kb. These fragments are used to further assemble the synthetic chromosome in a heterologous host or to replace the corresponding wild-type sequence in a stepwise manner. Each of the techniques have advantages and disadvantages (Box 1), and should be chosen carefully based on the use cases.

Synthesizing DNA goes viral

Although viruses and phages are not considered to be ‘alive’ they have a genome. They can reproduce themselves by leveraging the resources from a host. Viral genomes are rather small, with sizes between 1759 bps (Porcine circovirus [14]) and 1259 kb (Megavirus chilensis [15]) and can consist of DNA or RNA. The first complete synthesis of a viral RNA genome, the polio virus, was accomplished in 2002 [16]. The 7.5 kb synthesized cDNA genome was in vitro transcribed by RNA polymerase and can generate infectious virus particles after transfer into a cell free extract. Further viral RNA and DNA genomes were synthesized up to a size of 212 kb in recent years (Table 1). Synthesizing, as well as engineering variations of viral genomes to produce genome libraries, has an enormous potential for therapeutic applications. Vaccines and drugs could be quickly generated in
response to the emergency of a certain virus variant, which may help to prevent wider outbreaks [7].

**Synthesizing genomes of organelles**

Mitochondria in general, and the plastids of plants, contain a genome. Their sizes are rather small but show a huge variation in size and content. Studying these organelles is very interesting but challenging. Transformation of organelles must be done by bio-ballistic transformation [17]. The efficiency of synthetic DNA transformation is rather low. Mitochondria are the only organelles for which a complete organelle genome has been synthesized so far. The synthesis of the 16.3 kb mouse mtDNA genome was achieved by using 600 60mer oligonucleotides in four consecutive assembly rounds [18]. This step was predominantly the proof of principle for a DNA assembly method. However, it is an intriguing question why organelles still contain genetic content and have not migrated all necessary genes to the nucleus. There are exceptions in nature where the mitochondria do not contain any DNA [19].

**Synthesizing microbial genomes**

The genome of *Myoplasma capricolum* [12]. This is the first organism which is controlled by a synthetic genome, and is referred as *M. mycoides* JCVI-syn1.0. The genomic differences to *M. mycoides* are marginal and consist of designed ‘watermark’ sequences, 14 genes are deleted or disrupted and nineteen harmless polymorphisms were acquired during the building process.

This successful project was the starting point to generate a minimal *Mycoplasma* organism based on JCVI-syn1.0. Briefly, two independent teams failed to generate a viable cell, based on knowledge and genome synthesis, from scratch. However, multiple rounds of transposon mutagenesis and genome reduction finally generated *M. mycoides* JCVI-syn3.0 a minimal genome with a genome reduction of 50.8% in a design, build and test cycle manner [20,13**]. The 901 genes of *M. mycoides* JCVI-syn1.0 were reduced to 473 genes of which 149 are of unknown function and will give deeper insights into essentiality of genes [21].

An interesting ongoing project is the generation of a synthetic *Escherichia coli* genome. The genome of a previously diminished *E. coli* strain is redesigned in 87 ca. 50 kb segments to eliminate 7 codons in the coding sequence in a stepwise manner [97]. The 62 214 (5.4%) excluded codons are replaced by synonymous codons to maintain viability. The freed-up codons may be used to incorporate non-natural amino acids into proteins in the future. Absence of seven codons and corresponding tRNAs will, in addition, provide sufficient resistance to phages, rendering this strain of great general interest. Currently 55 of the 87 segments have been tested experimentally but the incorporation into a fully synthetic *E. coli* genome still needs to be proven functional.

**Synthesizing eukaryotic genomes**

As of today, there is no complete synthetic eukaryotic genome. However, the synthetic yeast genome — Sc2.0 project (www.syntheticyeast.org) aims to generate the first eukaryotic cell operated by a synthetic genome. The 16 chromosomes are synthesized in individual strains by teams of scientists within the Sc2.0 Consortium. The chromosomes are re-designed in a higher order of magnitude compared to any other existing write project [8**]. The major changes include the removal of most introns, transposons and repetitive elements. One central element of Sc2.0 design is the relocation of all tRNA genes to an independent 17th *Saccharomyces cerevisiae* chromosome, designated as the tRNA neochromosome. tRNA genes are heavily transcribed and therefore are hotspots of genomic instability caused by replication stress and transposon insertions. In addition, all non-essential genes are flanked by loxPsym sites which allow inducible large-scale genomic re-arrangements mediated by Cre-recombinase. This implemented genome rearrangement technique is therefore referred to as Synthetic Chromosome Assembly in Heterologous Organism.
Electron Microscopy and the Next Grand Challenge in Modern Biology

The successes in current genome synthesizing projects are leading to the next grand challenge in modern biological science: The Genome Project-write (GP-write). This project is a grand challenge using synthesis, gene editing and other technologies to understand, engineer and test living systems with the overarching goal to understand the blueprint for life provided by the Human Genome Project (HGP-read) [30**,31,32]. Therefore, a new international consortium was formed and first meetings were held in 2016 and 2017. The consortium is an open, interdisciplinary and international research group to focus efforts to realize GP-write.

GP-write has several goals, one being the development of new techniques and to accelerate the evolution of existing techniques with an overall goal to reduce synthesis costs by 1000-fold within ten years. Similar effects were achieved by HGP-read: today the cost of sequencing a human genome are magnitudes lower than the initial human genome sequence. The open nature of GP-write allows everyone to submit project proposals which will be evaluated by the Scientific Executive Committee. As of

### Table 1

Overview of finished synthesized chromosomes and genomes

| Year | Species | Size in kb (% of wt genome) | Highlights of the study | Reference |
|------|---------|----------------------------|-------------------------|-----------|
| 2002 | Polio virus | 7.5 | First infectious viral particles, based on a fully synthetic genome. | [16**] |
| 2003 | ϕX174 bacteriophage | 5.4 | Whole workflow: design, oligo synthesis, genome assembly and generation of infectious viral particles in fourteen days. | [34] |
| 2007 | Human endogenous retrovirus (HERV-K) consensus | 9.5 | Generation of a consensus genome of a human endogenous retrovirus which can replicate and is infectious. | [35] |
| 2008 | Bat severe acute respiratory syndrome (SARS)-like coronavirus (Bat-SCoV) | 29.7 | Until 2010 the largest synthetic replicating life form. | [36] |
| 2017 | Horsepox | 212 | First de novo synthesis of an in nature extinct orthopoxvirus of which close relatives causes smallpox in humans. | [7] |
| 2008 | Mycoplasma genitalium | 583 | Watermark sequences were inserted to identify synthetic DNA; disruption of one gene (MG408) to prevent pathogenicity and to do antibiotic selection. | [11] |
| 2010 | Mycoplasma mycoides JCVI-syn1.0 | 1079 | Watermark sequences were inserted to identify synthetic DNA; deletion or disruption of fourteen genes (one accidentally mediated by an IS1 E. coli transposon). | [12**] |
| 2016 | Mycoplasma mycoides JCVI-syn3.0 | 531.6 (49.3%) | Reduction of Mycoplasma mycoides JCVI-syn1.0 in a design, build, test manner based on transposon mutagenesis to reduce the genome from 901 to 473 genes. | [13**] |
| 2010 | Mus musculus mtDNA | 16.3 | Complete in vitro synthesis in four steps to produce a copy of the mouse mtDNA consensus genome. | [18] |
| 2011 | Saccharomyces cerevisiae synXII | 91.0 (101.9%)<sup>a</sup> | Applied design rules (if possible):<br>- Sympathetic replacement of TAG stop codons with TAA | [20] |
| 2012 | Saccharomyces cerevisiae synIII | 272.2 (86.2%)<sup>a</sup> | Synonymous recoding of tandem repeats in CDSs | [24**] |
| 2017 | Saccharomyces cerevisiae synV | 770.1 (94.7%)<sup>a</sup> | Deletion of genomic instability causing elements: introns, transposons, subtelomeric repeats and tRNA genes | [25] |
| 2017 | Saccharomyces cerevisiae synVII | 536.0 (92.9%)<sup>a</sup> | Insertion of loxPsym sites to 'SCRaMbLE' the genome | [26] |
| 2017 | Saccharomyces cerevisiae synXV | 242.7 (89.9%)<sup>a</sup> | Insertion of PCRtag sequences to identify synthetic DNA | [29] |
| 2017 | Saccharomyces cerevisiae synXII | 707.5 (94.9%)<sup>a</sup> | Insertion of PCRtag sequences to identify synthetic DNA | [27] |

<sup>a</sup> The size of synXII is slightly longer due to the insertion of 43 loxPsym sites (34 bps each).<br><sup>b</sup> Chromosome XII contains a cluster with >100 copies of the 9.1 kb rDNA operon which is not included in the shown size.

Rearrangement and Modification by LoxP-mediated Evolution (SCRaMbLE) and has already proven its functionality [20,22*,23*].

Recent publications report the synthesis and characterization of six Sc2.0 chromosomes and the right arm of synthetic chromosome IX (Table 1) which collectively correspond to 32% of the yeast genome [20,24**,25–29]. Strikingly, the individually synthesized chromosomes can be merged in a single cell by mating with a technique called endoreduplication intercross [5**]. Currently, the strain with the most synthetic chromosomes in one cell contains synIII, synV and synXII. With further progression of the Sc2.0 project more synthetic chromosomes will be finalised and ultimately merged to the final Sc2.0 strain.

**GP-write: a sneak preview into the future of synthetic genomics**

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Table 2

| Project title                                      | Project goals                                                                                           | Project lead(s)          |
|---------------------------------------------------|---------------------------------------------------------------------------------------------------------|--------------------------|
| UltraSafe Cell Line                               | The project aims to generate an Ultrasafe cell line by altering roughly 1% of the human genome. Some key goals are: Virus and prion resistance, removal of transposable elements, recoding of triplet repeats, recoding to a human consensus sequence in regard to SNPs and indels, implementing the bespoke SCRaMbLE system beside further alterations. | Jef Boeke & George Church |
| High-throughput HAC Design to                     | There is still a lack of understanding of the regulation of gene expression. The project will build two 1 mb regions of the human genome. The regions will be constructed as combinatorial libraries with different promoters and insulators to investigate ‘rules’ for optimal gene expression. | Pamela Silver & Jeffrey Way |
| Test Connections Between Gene Expression, Location and Conformation | This project tries to solve two questions based on Sc2.0 strains and implement the results into GP-write. Firstly: how can hybridization with natural occurring strains or organisms be excluded? Second: what shapes the contact maps of chromosomes? The latter will be answered by analysing multiple SCRaMbLed Sc2.0 strains to investigate chromatin structure based on re-arranged chromosomes | Jasper Rine |
| Safety and Containment; Chromatin and Chromosome Structure | Codon alteration is an important part of GP-write. The project aims to develop: Firstly, a rapid method for multiplex targeted genome modification; secondly, a respective rapid and robust screening system for living cells in 96-well format; thirdly, a strategy for rapid evaluation of heterogenic cell populations; and finally, a software to design the synthetic DNA fragments and evaluate viability of codon replacements. | Marc Lajoie |
| Empirically Designing Genomically Recoded Human Cell Lines | Efficient and precise engineering of the human genome is still a challenge. This project aims to create a complete pipeline for rapid engineering of human cells with an enrichment for homologous recombination repair. The project will also provide bioinformatic tools to optimize CRISPR based engineering. | Neville Sanjana |
| The Seven Signals Toolbox:                         | Cell differentiation is mainly driven by seven signal types. This project aims to generate a toolbox which allows the in vitro differentiation of GP-write cell lines. This is a crucial step for future applications in the field of cell therapies, tissue replacement or transplantation of organs. | Liam Holt |
| Leveraging Synthetic Biology to                   | fruition of gene expression. The project aims to define the Logic of Stem-Cell Programming. Cells will be transfected with various constructs to investigate the dependency of gene expression on re-arranged chromosomes and evaluate viability of codon replacements. |                          |
| Define the Logic of Stem-Cell Programming          | Efficient and precise engineering of the human genome is still a challenge. This project aims to create a complete pipeline for rapid engineering of human cells with an enrichment for homologous recombination repair. The project will also provide bioinformatic tools to optimize CRISPR based engineering. | Neville Sanjana |
| Precision Human Genome Engineering of Disease-AssOCIATED Noncoding Variants | Efficient and precise engineering of the human genome is still a challenge. This project aims to create a complete pipeline for rapid engineering of human cells with an enrichment for homologous recombination repair. The project will also provide bioinformatic tools to optimize CRISPR based engineering. | Neville Sanjana |
| Synthesizing a Prototrophic Human Genome          | This project postulates to introduce pathways for the nine amino acids and a variety of vitamins which cannot be synthesized by humans. These molecules derive from the diet. It investigates whether the milieu in the cell makes a prototrophic cell line feasible. If the project succeeds, further engineering would be performed and the first achievement would be a drastic cost reduction of cell line cultivation media. | Harris Wang |
| Through the Looking Glass:                        | Including the publics view and governance systems into GP-write is an important step. This project will generate a dialogue between scientists and the public. Incorporation of the society will enable acceptance and support for GP-write. | Todd Kuiken & Gigi Gronvall |
| Anticipating and Understanding Governance Systems and the Public’s Views on HGP-write | Synthetic Screening for Essential Intron and Retroelements in Human Cell and Animals | Yasunori Aizawa |
| Synthetic Screening for Essential Intron and Retroelements in Human Cell and Animals | This project aims to perform systematic screenings of intron and retroelements in the genome. Combinatorial variants of chosen genes will be investigated in a diploid background. The outcome will indicate if the removal of these elements, like in Sc2.0, is feasible in GP-write. | Max Berry |
| Isothermal Amplification Array                     | This project proposes a new method to synthesize DNA. It depends on two steps. Firstly: generation of short oligonucleotides by an isothermal amplification on an array. Secondly: the amplified oligonucleotides can anneal according to their design and nicks are sealed by a ligase. | Max Berry |
| Recombinase-Mediated Assembly                      | This project proposes a new method to assemble DNA fragments by utilizing a RecA-like recombinase (UvaX). The method should allow, with the Isothermal Amplification Array assembly from short oligos to chromosomem-sized DNA, with a significant labour reduction. | Max Berry |
| Synthetic Regulatory Genomics                     | The project aims to study regulatory variations of non-coding regions. The project will use multi-edited regulatory DNA sequences and analyse their function with multiple techniques. This project will give deeper insights into non-coding regions of complex genomes. | Matt Maurano |
| Concepts & Ethics in GP-write:                     | This project aims to build a model for deep analysis of concepts and ethics in GP-write, and aims include the dynamics of science and society. It aims to expand collaborations between sciences and the humanities and provide proper education and training. | Jeantine Lunshof |

* More detailed information can be found at [http://www.engineeringbiologycenter.org/](http://www.engineeringbiologycenter.org/).
January 2018 there are 13 pilot projects approved (Table 2). The projects cover many aspects of synthetic genomics, two highlights are the projects dedicated to the Concepts and Ethics in GP-write as well as Anticipating and Understanding Governance Systems and the Publics Views on HGP-write, which shows the importance to consider ethics and the publics views within GP-write.

One major remaining question is: What can we learn from GP-write? On one hand, there will be the ad hoc advances in enabling technologies, on the other hand there will be an immense gain of knowledge in biological sciences. Our knowledge of complex genomes is still limited. For instance, roughly 1% of the genome is responsible for all proteins in the cell. The remaining 99% are often referred as the ‘dark matter’ of the genome. Stepwise replacement of these elements, like in the Sc2.0 project, will potentially help us decipher the functions of the dark matter in the genome. On the application front, the pilot project to engineer a stable and safe cell line, has a profound implication for biomanufacturing and bioproduction (Table 2). GP-write still has a long way to go. However, the scientific community is curious about the outcome of the first pilot projects in the GP-write framework.

Conclusion
The initial genome writing projects summarized here show that individual native chromosomes and whole genomes can be replaced by chemically synthesized genomes. So far, the changes to DNA sequences are relatively modest but with growing knowledge of biological systems, the design will become more aggressive and adventurous which will lead us into previously unexplored territories. The exciting field of synthetic genomics will give new insights in basic research and will open new possibilities in applied science.

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Conflict of interest statement
Nothing declared.

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