

SEMINAR REPORT

On

Bio Computers

Submitted by:

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for the award of the degree

of

Bachelor of Technology



DEPARTMENT OF COMPUTER SCIENCE AND ENGINEERING
COLLEGE OF ENGINEERING , TRIVANDRUM

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CERTIFICATE



College of Engineering, Trivandrum

This is to certify that the seminar entitled **Bio Computers** done by **Vinod Kumar S (13400061)** of the **Department of Computer Science and Engineering, College of Engineering, Trivandrum** in partial fulfillment of the requirements for the award of the degree of Bachelor of Technology in Computer Science and Engineering under Kerala University is a Bonafide record of work carried out by him.

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Place : Trivandrum

Date : 25th July, 2016

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ABSTRACT

Biocomputing is one of the upcoming field in the areas of molecular electronics and nanotechnology. The idea behind blending biology with technology is due to the limitations faced by the semiconductor designers in decreasing the size of the silicon chips, which directly affects the processor speed. Biocomputers consists of biochips unlike the normal computers, which are silicon-based computers. This biochip consists of biomaterial such as nucleic acid, enzymes, etc.

The power of a biocomputer is that it acts as a massively parallel computer and has immense data storage capability. Thus, it can be used to solve NP-complete problems with higher efficiency. The possibilities for biocomputers include developing a credit card size computer that could design a super-efficient global air-traffic control system. The basic idea behind biocomputing is to use molecular reactions for computational purposes.

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Abbreviations

DNA	Deoxyribo Nucleic acid
RNA	Ribo Nucleic acid
HPP	Hamiltonion Path Problem
NP	Nondeterministic Polynomial time
PCR	Polymerase Chain Reaction
DES	Data Encryption Standard

1 Introduction

Biocomputers use systems of biologically derived molecules- such as DNA and proteins to perform computational calculations involving storing, retrieving and processing data.

DNA (Deoxyribo Nucleic Acid) computing, also known as molecular computing is a new approach to massively parallel computation based on groundbreaking work by Adleman. DNA computing was proposed as a means of solving a class of intractable computational problems in which the computing time can grow exponentially with problem size (the ‘NP - complete’ or non-deterministic polynomial time complete problems) . A DNA computer is basically a collection of specially selected DNA strands whose combinations will result in the solution to some problem, depending on the problem at hand. Technology is currently available both to select the initial strands and to filter the final solution.

DNA computing is a new computational paradigm that employs (bio) molecular manipulation to solve computational problems, at the same time exploring natural processes as computational models. In 1994, Leonard Adleman at the Laboratory of Molecular Science, Department of Computer Science, University of Southern California surprised the scientific community by using the tools of molecular biology to solve a different computational problem.

2 DNA Computing

2.1 DNA Fundamentals

DNA (deoxyribonucleic acid) is a double stranded sequence of 4 nucleotides. The four nucleotides that compose a strand of deoxyribonucleic acid are as follows: Adenine (A), guanine (G), Cytosine (C) and Thymine (T): they are typically referred to as bases. The order of those bases is what determines DNA’s instructions, or ordering.

DNA performs two major functions:

- self replication.
- codes for production of proteins.

Each deoxyribonucleotide consists of 3 components:

- a sugar — deoxyribose.
 - five carbon atoms: 1’ to 5’.
 - hydroxyl group (OH) attached to 3’ carbon.
- a phosphate group.
- a nitrogenous base.

The chemical structure of deoxyribonucleic acid consists of a specific bond of 2 linear sequences of bases. This bond follows a property of Complementarity: adenine bonds with T (A-T) and vice-verse (T-A), cytosine bonds with G (C-G) and vice-verse (G-C). This is called Watson-Crick complementarity. The four nucleotides adenine (A), guanine (G), cytosine (C) and thymine (T) compose a strand of DNA. The polarity of DNA strands is determined by its two different ends: the 3’end, and the 5’end. The double helix is an anti-parallel (two strands of opposite polarity) bonding of two complementary strands.

2.2 Basic Operations on DNA

DNA is the major data storage molecule in living cells. There are highly specific enzymes that can either duplicate the information in DNA molecules or transmit this information to other DNA molecules. Instead of using electrical impulses to represent bits of information, the DNA computer uses the chemical properties of these molecules by examining the patterns of combination or growth of the molecules or strings. DNA can do this through the manufacture of enzymes, which are biological catalysts that are analogous to ‘software’ for computing data.

A single strand of DNA is string that consists of four different symbols A, C, G, T. Mathematically this means we have at our disposal a letter alphabet, $\Sigma = \{A\ T\ G\ C\}$ to encode information. In a DNA computer, computation takes place in test tubes or on a glass slide coated in 24K gold. The input and output are both strands of DNA, whose genetic sequences encode certain information. A program on a

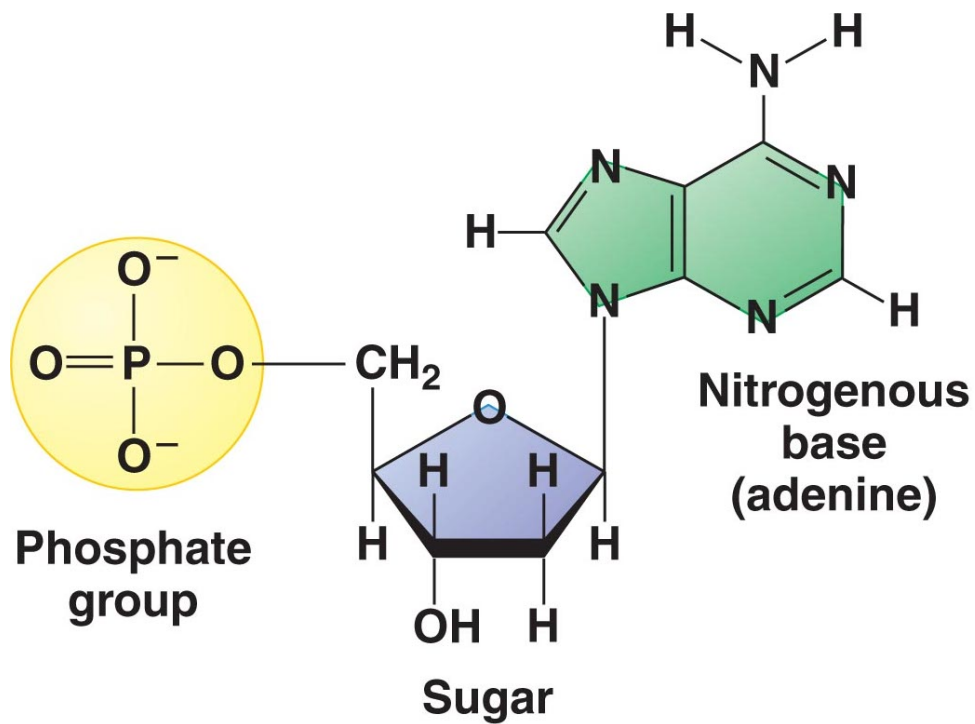


Figure 1: DNA Bases

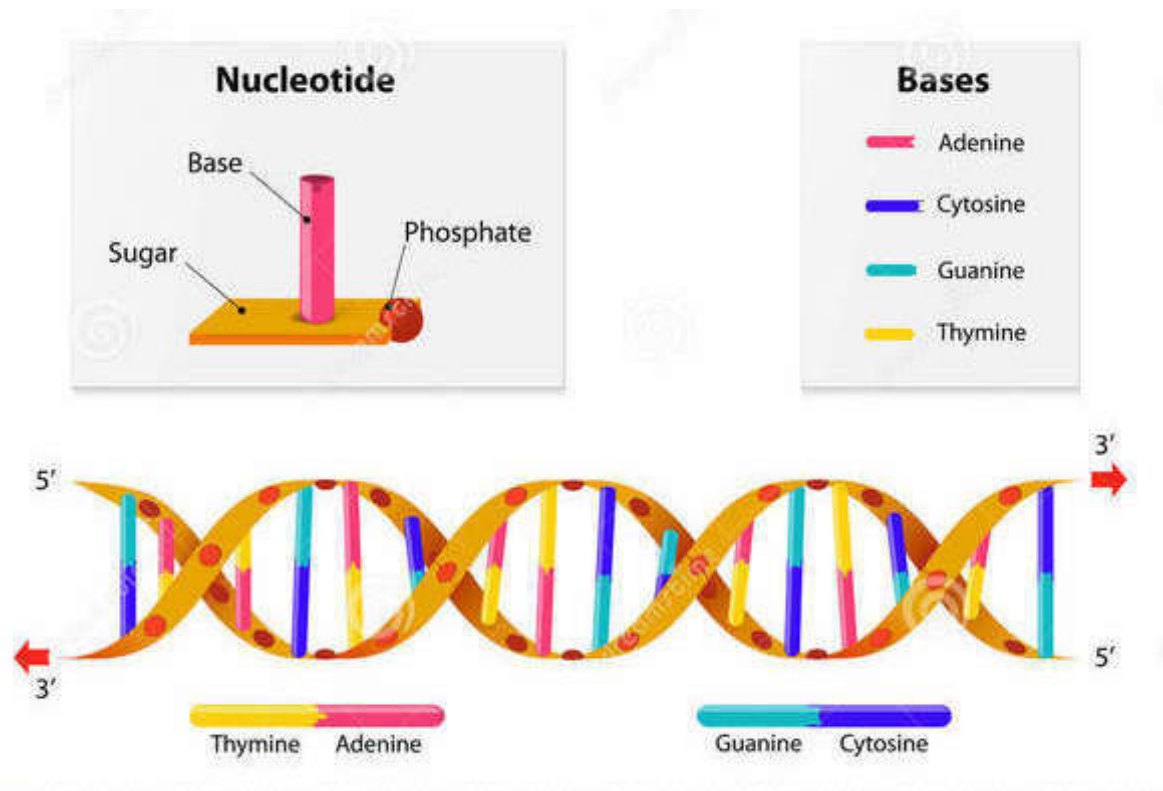


Figure 2: DNA Structure

DNA computer is executed as a series of biochemical operations, which have the effect of synthesizing, extracting, modifying and cloning the DNA strands.

DNA computation are based on various combinations of the following primitive bio-operations:

- **Synthesizing**- A desired polynomial length strand is generated.
- **Mixing**- Combining the content of two test tubes to perform union operation.
- **Annealing**- bond together two single-stranded complementary DNA sequences by cooling the solution.
- **Melting**- break apart a double-stranded DNA into its single-stranded complementary components by heating the solution.
- **Amplifying (copying)** - make copies of DNA strands by using the Polymerase Chain Reaction (PCR). The DNA polymerase enzymes perform several functions including replication of DNA. The replication reaction requires a guiding DNA single strand called template and a shorter oligonucleotide called a primer, that is annealed to it.
- **Separating**- Strands are separated by length using gel electrophoresis.
- **Extracting**- those strands that contain a given pattern as a substring by using affinity purification.
- **Ligating**- paste DNA strands with compatible sticky ends by using DNA ligases. Indeed, another enzyme called DNA ligase, will bond together, or “ligate”, the end of a DNA strand to another strand.
- **Cutting**- DNA double-strands at specific sites by using commercially available restriction enzymes. One class of enzyme, called restriction endonucleases, will recognize a specific short sequence of DNA, known as a restriction site. Any double-stranded DNA that contains the restriction site within its sequence is cut by the enzyme at that location.
- **Substituting**- substitute, insert or delete DNA sequences by using PCR site-specific oligonucleotide mutagenesis.
- **Marking**- single strands by hybridization: complementary sequences are attached to the strands, making them double-stranded. The reverse operation is **unmarking** of the double-strands by denaturing, that is, by detaching the complementary strands. The marked sequences will be double-stranded while the unmarked ones will be single-stranded.
- **Detecting and Reading**- given the contents of a tube, say “yes” if it contains at least one DNA strand, and “no” otherwise. PCR may be used to amplify the result and then a process called **sequencing** used to actually read the solution.

In short, the DNA computer encodes the problem to be solved in the DNA using the four values A, T, C, G. The solution to a problem is obtained as a DNA strand.

Every possible sequence can be chemically created in a test tube on trillions of different DNA strands and the correct sequences can be filtered out using genetic engineering tools.

2.3 DNA Computing Example - The Hamiltonian Path Problem

In 1994 Leonard M. Adleman showed how to solve the Hamilton Path Problem, using DNA computation.

Hamiltonian Path Problem:

A directed graph G with designated nodes *start* and *end* is said to have a Hamiltonian path if and only if there exists a sequence of compatible one-way edges e_1, e_2, \dots, e_n that begins at *start*, ends at *end* and enters every other node exactly once. A simplified version of this problem, known as the traveling salesman problem, poses the following question: given an arbitrary collection of cities through which a salesman must travel, what is the shortest route linking those cities?

This problem is difficult for conventional computers to solve because it is a “non-deterministic polynomial time problem”. These problems, when the instance size is large, are intractable with conventional computers, but can be solved using massively parallel computers like DNA computers. NP problems are intractable with deterministic (conventional/serial) computers, but can be solved using non-deterministic (massively parallel) computers. A DNA computer is a type of non-deterministic computer. The Hamiltonian Path problem was chosen by Adleman because it is known as “NP-complete”.

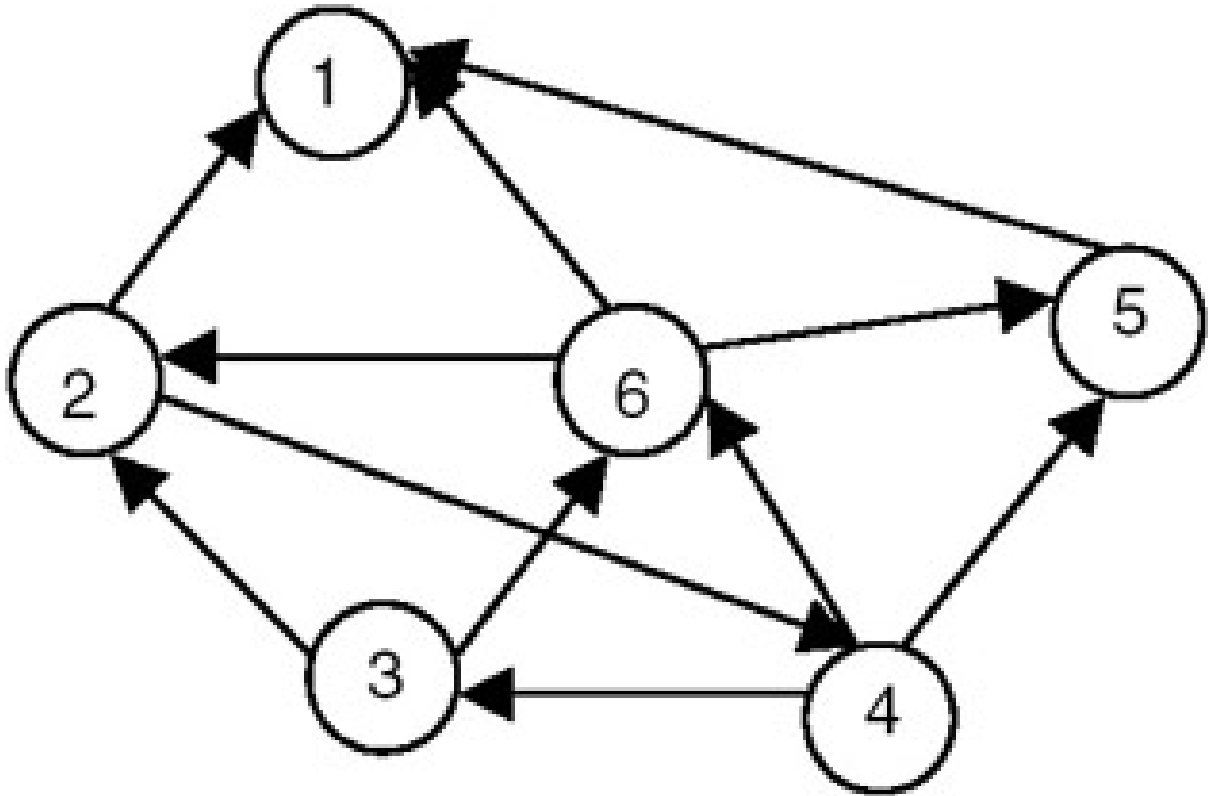


Figure 3: Directed graph with node 4 as source(start) and node 1 as destination node(end)

Adleman's Algorithm

Input: A directed graph G with n vertices, and designated vertices start and end.

Step 1: Generate paths in G randomly in large quantities.

Step 2: Reject all paths that

- do not begin with start.
- do not end in end.

Step 3: Reject all paths that do not involve exactly n vertices.

Step 4: For each of the n vertices v :

- reject all paths that do not involve v .

Output: YES, if any path remains; NO, otherwise.

For step 1, each node of the graph is encoded into a random 20-base strand of DNA. Then, for each edge of the graph, a different 20-base oligonucleotide was generated that contains the second half of the source code plus the first half of the target node.

To implement step 2, the product of step 1 was amplified by PCR using oligonucleotide primers representing start and end and ligase enzyme. This amplified and thus retained only those molecules encoding paths that begin with start and end with end. 10^{14} computations are carried out in a single second.

For implementing step 3, gel electrophoresis is used for the separation and recovery of DNA strands of the correct length. The desired path, if it exists, would pass through all four nodes, each of which was assigned a length of 20 bases.

Step 4 is accomplished by successive use of affinity purification for each node other than the start and end nodes.

The solution strand has to be filtered from the test tube:

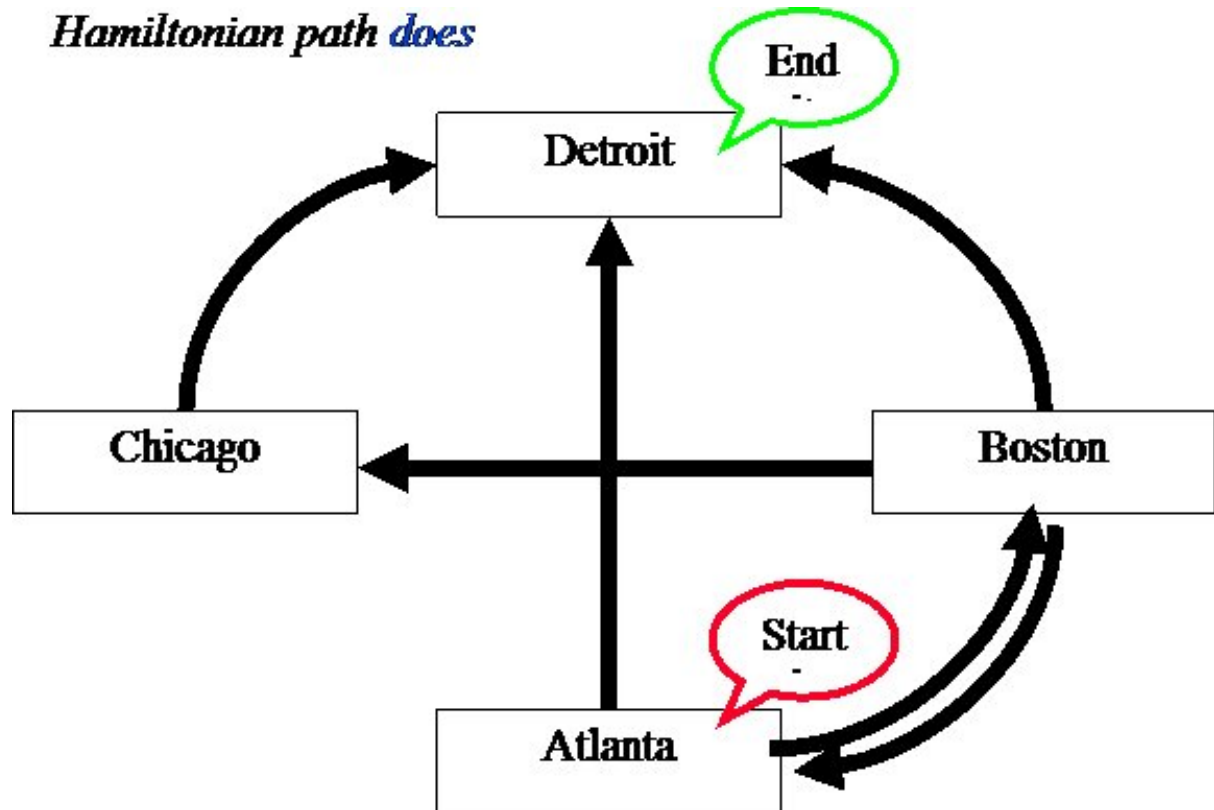


Figure 4: Simplified Graph - Hamiltonion Path is Atlanta-Boston-Chicago-Detroit

CITY	DNA NAME	COMPLEMENT
ATLANTA	ACTTGCAG	TGAACGTC
BOSTON	TCGGACTG	AGCCTGAC
CHICAGO	GGCTATGT	CCGATACA
DETROIT	CCGAGCAA	GGCTCGTT
FLIGHT	DNA FLIGHT NUMBER	
ATLANTA - BOSTON	GCAGTCGG	
ATLANTA - DETROIT	GCAGCCGA	
BOSTON - CHICAGO	ACTGGGCT	
BOSTON - DETROIT	ACTGCCGA	
BOSTON - ATLANTA	ACTGACTT	
CHICAGO - DETROIT	ATGTCCGA	

Figure 5: DNA for vertices and edges

GCAG TCGG ACTG GGCT ATGT CCGA

Atlanta → Boston → Chicago → Detroit

For a graph with n vertices, there are a possible $(n-1)!$ permutations of the vertices between beginning and ending vertex. To explore each permutation, a conventional computer must perform $O(n!)$ operations to explore all possible cycles. However, the DNA computing model only requires the representative oligos. Once placed in solution, those molecules will anneal in parallel, providing all possible paths in the graph at roughly the same time. That is equivalent to $O(1)$ operations, or constant time. In addition, no more space than what was originally provided is needed to contain the constructed paths.

3 Bio Computers

3.1 DNA Data Storage

A DNA storage system consists of a DNA synthesizer that encodes the data to be stored in DNA, a storage container with compartments that store pools of DNA that map to a volume, and a DNA sequencer that reads DNA sequences and converts them back into digital data. Figure 6 shows an overview of the integrated system.

The basic unit of DNA storage is a DNA strand that is roughly 100-200 nucleotides long, capable of storing 50-100 bits total. Therefore, a typical data object maps to a very large number of DNA strands. The DNA strands will be stored in “pools” that have stochastic spatial organization and do not permit structured addressing, unlike electronic storage media. Therefore, it is necessary to embed the address itself into the data stored in a strand. This way, after sequencing, one can reassemble the original data value.

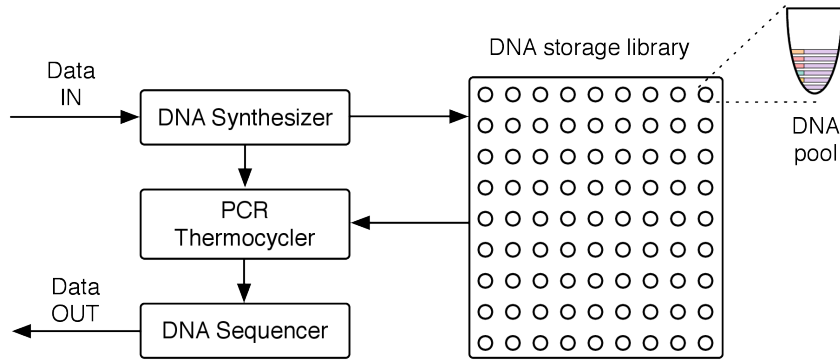


Figure 6: DNA Storage System Overview

A storage system needs a way to assign identification tags to data objects so they can be retrieved later. A simple key-value architecture is used, where a *put(key, value)* operation associates value with key, and a *get(key)* operation retrieves the value assigned to key. To implement a key-value interface in a DNA storage system, there is : (1) a function that maps a key to the DNA pool (in the library) where the strands that contain data reside; and (2) a mechanism to selectively retrieve only desired portions of a pool (i.e, random access), since the DNA container will likely store significantly more data than the desired object.

Random access is implemented by mapping a key to a pair of PCR primers. At write time, those primers are added to the strands. At read time, those same primers are used in PCR to amplify only the strands with the desired keys.

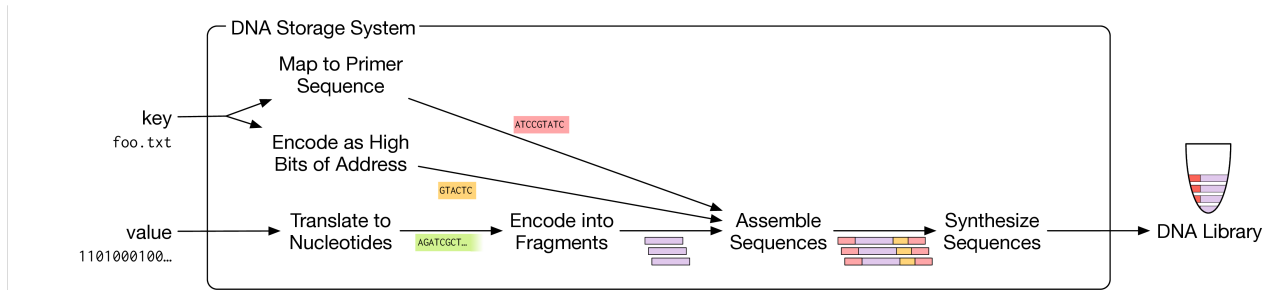


Figure 7: DNA Storage System Write

Figure 7 shows flowcharts for the write (*put*) and Figure 8 read (*get*) processes in more detail. The write process (Fig. 7) takes as input the key and value to store. It uses the key to obtain the PCR primer sequences, compute the high part of the address, and to determine the pool in the DNA library where the resulting strands will be stored. The low part of the address indexes the multiple strands generated

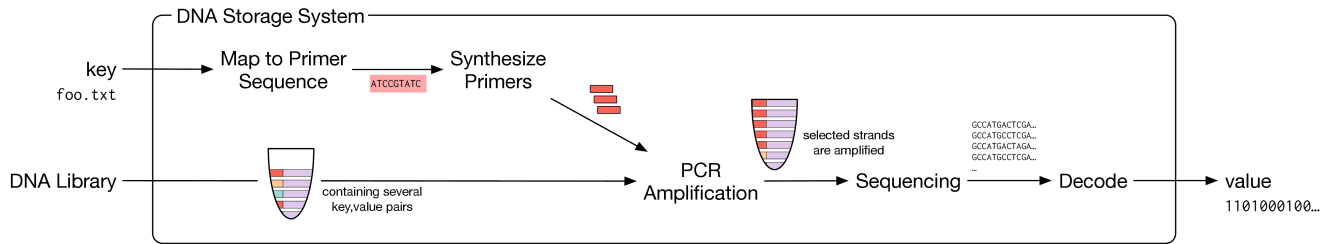


Figure 8: DNA Storage System Read

by chunking the value (see Sec. 4.2). Next, it encodes the data addresses, payloads and error detection codes, and attaches the primer target sequences, to produce final DNA sequences for the synthesizer to manufacture. The resulting DNA molecules are stored in the storage library for archival.

The read process (Fig. 8) takes as input a key. It uses the key to obtain the PCR primer sequences that identify molecules in the pool associated with that key. Next, the storage system physically extracts a sample from the DNA pool that contains the stored data, but likely also includes large amounts of unrelated data. The sample and the PCR primers are sent to the PCR thermocycler, which amplifies only the desired strands. The resulting pool goes to the DNA sequencer, which ultimately produces the digital data readout. Note that this process might be iterative since it may require multiple samples and sequencing steps to extract all the data associated with the desired keys. The DNA synthesizer is used for both producing the DNA strands that hold data payload as well as synthesizing the PCR primers used to amplify data during the random access read process.

The read process removes a sample of DNA from the pool, and so cumulative reads reduce the quantity of DNA available for future operations. But DNA is easy to replicate, and so the pools can easily be replenished after read operations if necessary. If successive amplification is problematic, a pool can also be completely resynthesized after a read operation.

Researchers at Microsoft and the University of Washington have reached an early but important milestone in DNA storage by storing a record 200 megabytes of data on the molecular strands. The impressive part is not just how much data they were able to encode onto synthetic DNA and then decode. It's also the space they were able to store it in. Once encoded, the data occupied a spot in a test tube “much smaller than the tip of a pencil.”

3.2 Cell to Cell Communication

Many viruses are capable of packaging and transmitting non-viral genetic material. For example, “transvestite” bacteriophage lambda particles can package and transduce phage T7 genomes or other genetic material. However, phage lambda and many viruses actively destroy the host cell in releasing virus particles.

Bacteriophage M13 (M13) is a filamentous phage whose progeny particles are secreted from infected cells. M13 is selected as the cell-communicating virus. It's the ideal specimen: It doesn't kill the host cell, it is possible to vary the length of DNA that can be packed, and it can be engineered to get its DNA into mammalian cells. The M13 communication system is like a wireless information network for cells to send and receive messages. M13 wraps up strands of DNA (programmed by scientists) and sends them out in proteins that infect cells and release the DNA messages once they have gained entry. Scientists can send whatever they want in the DNA, everything from a sentence in a book to a sequence that encodes fluorescent protein. The M13 system dramatically increases the amount of data that can be transmitted at one time compared to previous cell-to-cell communication systems, roughly 80,000 bits compared to one bit with the sugar molecule system. M13 can also transmit data over long ranges. The technology could be used in tissue engineering as well as in creating artificial organs and biomaterials that have no direct analog in nature. This is referred to as “Biological Internet”.

3.3 Transcriptor - Biological Transistor

A biological transistor is made from genetic material — DNA and RNA — in place of gears or electrons. The team calls its biological transistor the “transcriptor”. The biological transistor developed by Jerome

Bonnet and colleagues could be used inside living cells to record when cells have been exposed to certain external stimuli, or even to turn on and off cell reproduction as needed. The creation of the transcriptors enable bio computers to study and reprogram living systems, monitors environments and improve cell therapeutics.

In electronics, a transistor controls the flow of electrons along a circuit. Similarly, in biologics, a transcriptor controls the flow of a specific protein, RNA polymerase, as it travels along a strand of DNA. A group of natural proteins, called integrases, is repurposed to realize digital control over the flow of RNA polymerase along DNA, which in turn allowed to amplify genetic logic. Using transcriptors, it is possible to create what is known in electrical engineering as logic gates that can derive true-false answers to virtually any biochemical question that might be posed within a cell. These transcriptor-based logic gates as “Boolean Integrase Logic,” or “BIL gates” for short. Transcriptor-based gates alone do not constitute a computer, but they are the third and final component of a biological computer that could operate within individual living cells.

“AND” and “OR” are just two of the most basic Boolean logic gates. An “AND” gate, for instance, is “true” when both of its inputs are true — when “a” and “b” are true. An “OR” gate, on the other hand, is true when either or both of its inputs are true. In a biological setting, the possibilities for logic are as limitless as in electronics, it is possible to test whether a given cell had been exposed to any number of external stimuli — the presence of glucose and caffeine, for instance. BIL gates will make that determination and store that information so that it could easily identify those which had been exposed and which had not. By the same token, it can command the cell to start or stop reproducing if certain factors were present. And, by coupling BIL gates with biological Internet, it is possible to communicate genetic information from cell to cell to orchestrate the behavior of a group of cells. The transcriptor achieves a key similarity between the biological transistor and its semiconducting cousin: signal amplification.

4 Comparison with Conventional Computers

Similarities

- **Transformation of Data**

Both DNA computers and electronic computers use Boolean logic (AND, OR, NAND, NOR) to transform data. The logical command “AND” is performed by separating DNA strands according to their sequences, and the command “OR” is done by pouring together DNA solutions containing specific sequences. For example, the logical statement “X or Y” is true if X is true or if Y is true. To simulate that, the scientists would pour the DNA strands corresponding to “X” together with those corresponding to “Y”. Following is an example of how a Bio Chemical Inverter works. Working of inverter, the concentration of a particular messenger RNA (mRNA) molecule represents a logic signal. In the first case, the input mRNA is absent and the cell transcribes the gene for the output mRNA using RNA polymerase (RNAP) molecules. In the second case, the input mRNA is present and the cell translates the input mRNA into the input protein using ribosomes. A digital inverter that consists of a gene encoding the instructions for protein B and containing a region (P) to which protein A binds. When A is absent (left)—a situation representing the input bit 0, the gene is active and B is formed—corresponding to an output bit 1. When A is produced (right)—making the input bit 1, it binds to P and blocks the action of the gene—preventing B from being formed and making the output bit 0.

- **Manipulation of Data**

Electronic computers and DNA computers both store information in strings, which are manipulated to do processes. Vast quantities of information can be stored in a test tube. The information could be encoded into DNA sequences and the DNA could be stored. To retrieve data, it would only be necessary to search for a small part of it - a key word, for example - by adding a DNA strand designed so that its sequence sticks to the key word wherever it appears on the DNA.

- **Computation Ability**

All computers manipulate data by addition and subtraction. A DNA computer should be able to solve a satisfiability problem with 70 variables and 1,000 AND-OR connections. To solve it, assign various DNA sequences to represent 0's and 1's at the various positions of a 70 digit binary

number. Vast numbers of these sequences would be mixed together, generating longer molecules corresponding to every possible 70-digit sequence

Differences

	Conventional	Biological
Components and Materials	Inorganic, e.g. silicon	Biological, e.g.DNA
Data representation	Base 2	Base 4
Size	Large	Small
Processing scheme	Sequential and limited massively parallel	Massively parallel
Toxic components?	Yes	No
Energy efficient?	No	Yes

Table 1: Table to test captions and labels

5 Advantages and Disadvantages of bio computers

Advantages

- Performs millions of operations at same time.
 - Good for parallel computing.
- Ability to use large amounts of working memory.
 - 1 gram of DNA can hold 1 x 10¹⁴ MB of data Or 145 trillion CDs.
 - 1 CD is 800 MB.
- Cheaper.
- Lightweight.
 - 1 lb of DNA has more computing power than all computers ever made.
- Low power used to keep in original state.
- Has ability to solve hardest problems in a matter of weeks.
- Environmentally friendly.
 - Clean, readily available materials.

Disadvantages

- Molecular operations are not perfect.
- DNA computing involves a relatively large amount of error.
- As size of problem grows, probability of receiving incorrect answer eventually becomes greater than probability of receiving correct answer.
- Sometimes there are errors in the pairing of DNA strands.
- Simple problems solved faster on electronic computers.
- Human assistance is required.

- No universal method of data representation.
- Time consuming lab procedures.
- DNA has a half-life.
 - Solutions could dissolve away before end result is found.
- Information can be untransmittable.
 - Current DNA algorithms compute successfully without passing any information from one processor to the next in a multiprocessor connection bus.

6 Applications

6.1 Medical Application

Biological computers are made inside a patient's body. The mere information of the patient's body is called a blueprint along which lines the biological computer would be manufactured. Once the computer's genetic blueprint has been provided, the human body will start to build it on its own using the body's natural biological processes and the cells found in the body. Through boolean logic equations, we can easily use the biological computer to identify all types of cellular activity and determine whether a particular activity is harmful or not. The cellular activities that the biological computer could detect can even include those of mutated genes and all other activities of the genes found in cells. As with conventional computers, the biological computer also works with an output and an input signal. The main inputs of the biological computer are the body's proteins, RNA and other specific chemicals that are found in the human cytoplasm. The output on the other hand could be detected using laboratory equipment.

6.2 Solving NP-Problems

The primary advantage offered by most proposed models of DNA based computation is the ability to handle millions of operations in parallel. The massively parallel processing capabilities of DNA computers may give them the potential to find tractable solutions to otherwise intractable problems, as well as potentially speeding up large, but otherwise solvable, polynomial time problems requiring relatively few operations.

A working DNA based computer might hold an advantage over conventional computers when applied to decomposable problems, those problems that are able to be divided into separate, non-sequential tasks, because they can hold so much data in memory and conduct so many operations at once. However, due to the length of time required to conduct the biochemical operations, non-decomposable problems, those requiring many sequential operations, are likely to remain much more efficient on a conventional computer.

6.3 Cryptography

DNA computing techniques have already been theoretically applied to a real life problem: breaking the Data Encryption Standard (DES). Although this problem has already been solved using conventional techniques in a much shorter time than proposed by the DNA methods, the DNA models are much more flexible, potent and cost effective.

DES is a method of encrypting 64-bit messages with a 56-bit key, used extensively in the United States. Electronic keys are normally a string of data used to code and/or decode sensitive messages. By finding the appropriate key to a set of encrypted messages, one can either read encoded messages or pose as the such messages. Using a special purpose electronic computer and differential cryptanalysis, it has been shown that the key to DES can be found in several days. However, to do so would require 2^{43} examples of corresponding encrypted and unencrypted messages (known as plain-text/cipher-text pairs) and would slow down by a factor of 256 if the strength of the encrypting key was increased to 64-bits. It is proposed that DES could be broken using a DNA based computer and a search algorithm similar to Adleman's original technique. This procedure would be expected to take 4 months, but would only need a single plain-text/cipher-text pair or an example of cipher text with several plain text candidates to be successful. The feasibility of applying DNA computation to this problem was also addressed using a

more refined algorithm (the sticker model approach) which enabled the researchers to suggest that they could solve the problem using less than a gram of DNA, an amount that could presumably be handled by a desk top sized machine. Both models would likely be more cost and energy effective than the expensive electronic processors required by conventional means, but are entirely theoretical. The first model ignores error rates incurred through laboratory techniques and the inherent properties of the DNA being used. The second model requires an error rate approaching .0001, with higher rates substantially affecting the volume of DNA required. Despite these assumptions, these models show that existing methods of DNA computation could be used to solve a real life problem in a way that is both practical and superior to methods used by conventional computers. It is also demonstrated that such benefits can be obtained despite error rates that would be unacceptable in an electronic computer and that may be unavoidable in a molecular one.

7 Conclusion

While a desktop PC is designed to perform one calculation very fast, DNA strands produce billions of potential answers simultaneously. This makes the DNA computer suitable for solving “fuzzy logic” problems that have many possible solutions rather than the either/or logic of binary computers. In the future, some speculate, there may be hybrid machines that use traditional silicon for normal processing tasks but have DNA co-processors that can take over specific tasks they would be more suitable for.

Implanting computer programs into living creatures may not be far away. In the past few years, scientists have taken the first steps towards creating a host of cellular robots that are programmed to carry out tasks such as detecting and cleaning up environmental pollutants, tracking down cancer cells in a body and manufacturing antibiotics or molecular-scale electronic components. These researchers have imported notions of electrical engineering—digital logic, memory and oscillators—into the realm of biology.

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