Simplification of Boolean algebra through DNA Computing

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ABSTRACT

DNA Computing utilizes the properties of DNA for performing the computations. The computations include arithmetic and logical operations such as simplification of Boolean expression to its simplest form. Boolean function can be built from ANDs, ORs, and NOTs using minterm expansion. However, a practicing computer engineer will very rarely be satisfied with a minterm expansion, because as a rule, it requires more gates than necessary. The laws and identities of Boolean algebra will almost always allow us to simplify a minterm expansion. The efficiency of a logic circuit is high when the number of logic gates used to build it is small. However, minterm expression may be often simplified to a simpler Boolean expression, which can be implemented with fewer logic gates.

In this paper we introduced a new DNA computing algorithm for reducing any Boolean expression to its simplest form by using DNA strands. The major benefits of this method are its extraordinary information density, vast parallelism and ease of operation. In addition the most merit of this DNA Algorithm is its automation characteristics, and simple coding steps.

Keywords

DNA computing, Minterms, Simplify, DNA Strands, Boolean expression.

1. INTRODUCTION

DNA computing is new computation paradigms, which proposes the use of molecular biology tools to solve different mathematical problems. It is a form of computing which use DNA, biochemistry and molecular biology, instead of the traditional silicon-based computer technologies. DNA computing is interested in applying computer science methods and models to understand such biological phenomena and gain interest into early molecular evolution and origin of biological information processing. The primary advantage of DNA based computation is the ability to handle millions of operations in parallel. DNA computing is fundamentally similar to parallel computing in that it takes advantage of the many different molecules of DNA to try many different possibilities at once. DNA computing has two important features, which are Watson-Crick complimentarily and massive parallelism. Using the features, we solve some optimization problems, which usually need exponential time on silicon-based computers, in polynomial steps with DNA molecules.

However, for DNA computing to be applicable on a various range of problems for primitive operations, such as logic or arithmetic operations. A number of procedures have been proposed for the primitive operations with DNA molecules [1], [3], [4], [10], [11], [12], [14].

Boolean algebra:

Boolean algebra is algebra for the manipulation of objects that can take on only two values, typically true and false, although it can be any pair of values. Because computers are built as collections of switches that are either "on" or "off," Boolean algebra is a very natural way to represent digital information. In reality, digital circuits use low and high voltages, but for our level of understanding, 0 and 1 will suffice. It is common to interpret the digital value 0 as false and the digital value 1 as true.

Boolean Expression:

A Boolean expression on the Boolean variables $\{x_1, x_2, ..., x_n\}$ is an expression using those variables and the operations of a Boolean algebra. Every Boolean expression defines a Boolean function. Boolean function can be built from ANDs, ORs, and NOTs using minterm expansion.

Simplification of Boolean algebra:

The laws and identities of Boolean algebra will almost always allow us to simplify a minterm expression. The efficiency of a logic circuit is high when the number of logic gates used to build it is small. However, the sum-of-products (minterm) expression may be often simplified to a simpler Boolean expression, which can be implemented with fewer logic gates.

In this paper we introduced a new DNA algorithm for reducing any Boolean expression to its simplest form by using different combination of single strand DNAs. The rest of the paper is organized as follows: in section 2, we introduce the mathematical process to simplify Boolean expression, in section 3, we introduce the DNA algorithm to simplify the Boolean expression and section 4, we will give the example that how to simplify the Boolean Expression by using DNA computing following by conclusion.

2. SIMPLIFICATION OF BOOLEAN ALGEBRA

Some standardized forms are required for Boolean expression to simplify communication of the expression.

Sum-of-products (SOP): Example:

F(A,B,C,D) = AB + BCD + AD

The minterms in this sum correspond to those combinations of the values for which the function has a value of 1. This Boolean sum is sometimes called a **sum of products expansion** or **disjunctive normal form**.

We know that any Boolean function can be built from ANDs, ORs, and NOTs using minterm expansion. However, a practicing computer engineer will very rarely be satisfied with a minterm expansion, because as a rule, it requires more gates than necessary. The laws and identities of Boolean algebra will almost always allow us to simplify a minterm expansion. For example, the minterm expansion for a Boolean function f of three variables might be represented as follows:

 $\mathbf{f} = \mathbf{x'y'z'} + \mathbf{x'y'z} + \mathbf{x'yz'} + \mathbf{x'yz} + \mathbf{xyz'} + \mathbf{xyz}$

This would require a circuit with maximum gates: 12 ANDs, 5 ORs and 9 NOTs.

Using the identities of Boolean algebra, this minterm expansion can be simplified considerably:

$$\begin{split} f &= x'y'z' + x'y'z + x'yz' + x'yz + xyz' + xyz \\ &= x'y'(z'+z) + x'y(z'+z) + xy(z'+z) \text{ distributive law} \\ &= x'y' + x'y + xy \text{ complementarity & identity} \\ &= x'(y'+y) + xy \text{ distributive law} \\ &= x' + xy \text{ complementarity & identity} \\ &= x' + y \text{ redundancy} \end{split}$$

So, that big long minterm reduces down to x' + y which can be built with 1 OR and 1NOT. Therefore, we will look at a very simple technique that usually leads to a significant simplification of minterms. It won't always produce the simplest form, but it's close enough for most engineers considering the difficulty of the alternative method.

3. A NEW DNA ALGORITHM FOR SIMPLIFICATION BOOLEAN ALGEBRA

With the help of massive parallelism of DNA hybridization and the complementary Watson-Crick law, the optimal simplified expression can be found by basic molecular operations.

By means of the basic molecular operations such as merge, separate, denature, detect, etc., are used to simplify the Boolean expression to its simplest form.

For any given Boolean expression of 'n' variables in sum of products form, first of all we will see that if all the minterms contains 'n' variables or not. If not then we expand the expression with the possible combinations of variables left out in that specific minterm.

Algorithm:

Step 1: We choose 4n groups of oligonucleotides divided into four groups. The oligonucleotides of the first group represented variables x_1, x_2, \dots, x_n . The oligonucleotides of the second group represented variables $\overline{x_1}, \overline{x_2}, \dots, \overline{x_n}$ (where, x=1 if and only if x=0); the oligonucleotides of third and fourth group represented the complementary strands of the first group (that is $x_1', x_2', \dots, \overline{x_n}'$) and second group (that is $\overline{x_1}, \overline{x_2}', \dots, \overline{x_n}'$) respectively.

Step 2: We generate different 2^n combination of single strands DNA molecules for n variable where the oligonucleotides of third and fourth group are ligated according to the 2^n combinations of 'n' variables and placed those single stranded DNA molecules in individual 2^n test tubes.

Step 3: Now, the oligonucleotides of first and second groups are ligated according to minterms of the given expression. And AMPLIFY each minterms 2^n times.

<u>Step 4:</u> Then we MERGE the oligonucleotides of Step 2 and Step 3. The best paired strands are kept and remaining unpaired and semi-paired strands are separated from the test-tubes.

Step 5: The paired strands in each of the test-tube are denatured and keep the oligonucleotide which represent the complementary strands of minterms, in the test tubes and remaining strands are separated from the test tubes.

Step 6: Two minterms can be combined if they differ in exactly one literal. This means that their corresponding bit strings differ at exactly one bit position (Example: if 3 literals are present in each minterms then 000 can combined 001, 010, 100 etc.). So, the complementary strands of minterms which we get from step 5 are amplified n times, if one minterm can combined with n minterms, (if they differ in exactly one literal).

Step 7: Now, we have to generate the strands with combination of (n-1) variables and separate those strands in n different test tubes. Because n variables are present in each minterms. (Example: like 001 minterms having 3 combinations that are 00, 01 and 01seperated in three different test tubes).

Step 8: Now we have to combined those two minterms which are differ in exactly one bit position. (Example: like 001 minterms having 3 combinations that are 00, 01 and 01, seperated in three different test tubes. Now in that 3 test tube we merge the complementary strands of 0'0'1'.



Similarly, 001 differ 000 with exactly 1 bit position, so 000 minterms having 3 combination of 2 variables that are 00, 00 and

00, separated in three different test tubes and merge complementary strands of 0'0'0'.



Now we have to check that one combination is same for both the cases and that is 00).

Step 9: We store only common terms of two minterms (For above example, that is 00 because for both the cases common ds DNA is 00 and 0'0') and remaining portions are left out from the test tubes.

Step 10: We will denature the ds DNA which we get in step 9 and keep complementary part of the common terms (Example: for above case it is 0'0') and other are discarded from the test tubes.

Step 11: We store the result.

After getting result we will check any common terms are left between the minterms. If yes then go to next step otherwise go to step 18.

Step 12: Then we have to generate the strands with combination of (n-2) variables and separate those combinations in different test tubes.

Step 13: Now we have to combined those two minterms which are differ in exactly one bit position same as step 8.

Step 14: We store only common terms of two minterms and remaining portions are left out from the test tubes.

Step 15: We will denature the ds DNA and keep complementary part of the common terms in one test tube.

Step 16: We store the result and check that any common terms are left between the minterms which we get from step 15.

<u>Step 17:</u> If yes, then we have to generate the strands with combination of (n-3) variables and go to step 7. Process will continue until no common terms are left.

Step 18: End.

4. Example

For a given three variable expression:

$$Y = ABC + ABC + ABC + ABC + ABC + ABC$$

Here each minterms having equal number of variables. So there is no need of expansion.

Step 1: We choose 12 oligonucleotides which are divided into 4 groups. The oligonucleotides of the first group represents variables A, B, C; the oligonucleotides of the second group represents A, B, C; and the third and fourth group represent the complementary strands of first and second group respectively (Denoted as, A', B', C' and A', B', C').

Step 2: We generate 2^3 combinations of single strands DNA molecules for 3 variables. Where the oligonucleotides of third and fourth group are ligated according to the combinations of '3' variables and placed in individual test tubes.

	- -		-
A' B' C'	$A^{i}B^{i}C^{i}$	$A^{i}B^{i}C^{i}$	$A^{\prime}B^{\prime}C^{\prime}$
		-	
A' B' C'	A' B' C'	A'B'C'	A'B'C'

Figure 1. 2³ combination of ss DNA molecules are placed different test tubes

Step 3: Now, the oligonucleotides of first and second group are ligated according to minterms of the given expression. AMPLIFY 8 times for each minterms. Resulting DNA strands for minterms are as follows:

ABC	ABC	ABC	
- ABC	_ ABC	ABC	

Figure 2. ss DNA molecules representing minterms of a given Boolean Expression

Step 4: We MERGE the 8 sets of oligonucleotides which represent the minterms of Step 3 and with individual combinations which are already kept in 8 individual test tubes of Step 2. The best paired strands are kept and remaining unpaired and semi-paired strands are discarded from the test-tube.

ĀBC ĀBC	$\overline{\overline{A}} \overline{\overline{B}} C$ $\overline{\overline{A}}' \overline{\overline{B}}'C'$	ĀBĊ ĀBĊ	ĀВС Ā'B'C'	
Τo	T_1	T ₂	T_3	
АĒĈ	AĒC	A B Ĉ A' B' Ĉ'	ABC A'B'C'	
T₄	T₅	T_6	T7	

Figure 3. Merge each minterms with different combinations o	f
oligonucleotides	

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ĀBC ĀBC	ĀĒC Ā'Ē'C'	ĀBĊ ĀBĊ	Ā₿С Ā'₿'Ċ'
T ₀	T_1	T2	T ₃
		АВĊ A'B'Ċ'	ABC A'B'C'
		T_6	T 7

Figure 4. Unpaired and semi-paired strands are discarded

Step 5: The paired strands are denatured and the oligonucleotides which represent the complementary strands of each minterm are kept in the test tubes. Remaining strands are separated from the test tubes.

Ā'B'C'	A' Ē'C'	Ā'Ē'C'	Ā' B' C'
To	T ₁	T2	T ₃
		A'B'Ē'	A' B' C'
		T_6	T7

Figure 5. The paired strands are denatured and the complementary strands of each minterm are kept in test tubes

Step 6: Two minterms can be combined if they differ in exactly one literal. This means that their corresponding bit strings differ at exactly one bit position. For the above example, we can compare $T_0 - T_1$ and T_0 - T_2 , $T_1 - T_3$, $T_2 - T_3$ and $T_2 - T_6$ and finally $T_3 - T_7$ and $T_6 - T_7$ because in those test tubes the strands having bit string difference is exactly by 1 bit position.

The complementary strands of minterms which we get from step 5 are amplified.

AMPLIFY (T₀, T₀', T₀", T₀₀, T₀₀', T₀₀", T₀₀₀, T₀₀₀', T₀₀₀") AMPL1IFY (T₁, T₁', T₁", T₁₀, T₁₀', T₁₀₀, T₁₀₀', T₁₀₀") AMPL1FY (T₂, T₂', T₂", T₂₀, T₂₀', T₂₀", T₂₀₀, T₂₀₀', T₂₀₀") AMPLIFY (T₃, T₃', T₃", T₃₀, T₃₀', T₃₀", T₃₀₀, T₃₀₀', T₃₀₀") AMPLIFY (T₆, T₆', T₆", T₆₀, T₆₀', T₆₀", T₆₀₀, T₆₀₀") AMPLIFY (T₇, T₇', T₇", T₇₀, T₇₀', T₇₀", T₇₀₀, T₇₀₀")

	$\overline{\mathbb{A}'}_{\mathbf{B}'\mathbf{C}'}^{\mathbf{B}'\mathbf{C}'}$	Ā' Β'Ĉ' Τ ₀ '	Ā' В' Ĉ' Т2
Ā'Ē'C'	Ā'B'C', Ta		

Ā'Ē'C'	Ā' B' C'	Ā'Ē' C'	A'B' ⊄
T_2	T3'	T2''	T_6

Ā'B'C' A'B'C'		A' B' Ĉ'	A' B' C'	
T 3''	T7	Т _б ''	T 7'	

Figure6: Combined two minterms if they differ in exactly one literal

Step 7: Now, generate the strands with combination of 2 variables and separate those strands in three different test tubes and amplify those test tubes according to the minterms. For test tubes T_{0} , T_{0}' and T_{0}'' where, A=0, B=0, C=0, from that, we will take the combination of two variables and those are AB, BC, AC and put it into 1 test tube.

For T ₀ ,T ₀ ',T ₀ "	For T ₁ ,T ₁ ', T ₁ ''	For T ₂ ,T ₂ ', T ₂ ''	For T3 ,T3',T3''	For T ₆ ,T ₆ ', T ₆ ''	For T ₇ ,T ₇ ', T ₇ ''
ĀĒ BĒ	ĀC ĒC ĀĒ	ĀČ BČ ĀB	ĀC B C ĀB	АĊ ВĊ АВ	A C B C AB
T ₈	T9	T ₁₀	T11	T ₁₂	T ₁₃
1					

Figure7: Combination of 2 variables

 $\begin{array}{l} \text{SEPARATE} (T_8, T_{80}, T_{800}) \\ \text{AMPLIFY} (\underline{T_8}, T_{800}, T_{800}, T_8', \underline{T_{800}}', T_8'', T_{80}'', T_{800}'') [T_8: AB, \\ T_{80}: BC, T_{800}: AC] \\ \text{SEPARATE} (T_9, T_{90}, T_{900}) \\ \text{AMPLIFY} (\underline{T_9}, T_{90}, T_{900}, \underline{T_9}', T_{900}', T_9'', T_{90}'', T_{900}'') [T_9: B C, \\ T_{90}: AB, T_{900}: AC] \\ \text{SEPARATE} (T_{10}, T_{100}, T_{1000}) \\ \text{AMPLIFY} (T_{10}, T_{100}, T_{1000}, T_{10}', T_{1000}', T_{10}'', T_{100}'', T_{1000}'') \end{array}$

 $\begin{array}{l} \text{AMFLIF I (1_{10}, 1_{100}, 1_{100}, 1_{10}, 1_{100}, 1_{10}, 1_{100}, 1_{10}, 1_{100}, 1_{10}, 1_{100}$

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AMPLIFY (T₁₁, T₁₁₀, T₁₁₀, T₁₁', T₁₁₀', T₁₁₀', T₁₁", T₁₁₀", T₁₁₀") SEPARATE (T₁₂, T₁₂₀, T₁₂₀₀)

AMPLIFY (T₁₂, T₁₂₀, T₁₂₀₀, T₁₂', T₁₂₀', T₁₂₀₀', T₁₂", T₁₂₀", T₁₂₀₀") SEPARATE (T₁₃, T₁₃₀, T₁₃₀₀)

AMPLIFY (T₁₃, T₁₃₀, T₁₃₀₀, T₃', T₁₃₀', T₁₃₀₀', T₁₃", T₁₃₀", T₁₃₀₀")

Step 8: Now, we have to combine those two minterms which are differ in exactly one bit position.

Compare T₀ - T₁

MERGE $[T_0-T_8, T_{00}-T_{80}, T_{000}-T_{800} (For T_0 test tube)$



MERGE $[T_1-T_9, T_{10}-T_{90}, T_{100}-T_{900} (For T_1 test tube)]$



Figure8: Compare T₀ - T₁ and get result AB

Compare T_0 and T_1 test tubes, both the cases \overline{AB} is common.

Compare T₀' and T_{2:}

MERRGE $[\overline{T}_0' - T_8', \overline{T}_{00}' - T_{80}', T_{000}' - T_{800}' \text{ (For } T_0' \text{ test tube)}]$ MERGE [T₂-T₁₀, T₂₀-T₁₀₀, T₂₀₀-T₁₀₀₀ (For T₂ test tube)] Compare T_0' and T_2 test tubes, both the cases \overline{AC} is common.

Compare T₁' and T_{3:}

MERRGE $[T_1'-T_9', T_{10}'-T_{90}', T_{100}'-T_{900}'$ (For T_1' test tube)] MERGE $[T_3-T_{11}, T_{30}-T_{110}, T_{300}-T_{1100} (For T_3 test tube)$

Compare T_0' and T_2 test tubes, both the cases AC is common. Compare T₂' and T₃':

MERRGE $[\overline{T}_{2}'-T_{10}', \overline{T}_{20}'-T_{100}', T_{200}'-T_{1000}' \text{ (For } T_{2}' \text{ test tube)}]$ MERGE $[T_3'-T_{11}', T_{30}'-T_{110}', T_{300}'-T_{1100}' (For T_3' test tube)]$ Compare T_2' and T_3' test tubes, both the cases \overline{AB} is common. Compare T₂" and T₆':

 $\overline{\text{MERRGE} [T_2"-T_{10}", T_{20}"-T_{100}", T_{200}"-T_{1000}"} \text{ (For } T_2" \text{ test tube)}]$ MERGE $[T_6'-T_{12}', T_{60}'-T_{120}', T_{600}'-T_{1200}'$ (For T_6' test tube)] Compare T_2'' and T_6' test tubes, both the cases **BC** is common. Compare T₃" and T₇:

 $\overline{\text{MERRGE} [T_3"-T_{11}", T_{30}"-T_{110}", T_{300}"-T_{1100}" (For T_3" \text{ test tube})]}$ MERGE $[T_7-T_{13}, T_{70}-T_{130}, T_{700}-T_{1300} (For T_7 \text{ test tube})]$

Compare T_3'' and T_7 test tubes, both the cases **BC** is common. Compare T₆" and T₇':

 $\overline{\text{MERRGE} [\overline{T_6''} - T_{12}'', \overline{T_{60}}'' - T_{120}'', T_{600}'' - T_{1200}'' \text{ (For } T_6'' \text{ test tube)}]}$ MERGE [T₇'-T₁₃', T₇₀'-T₁₃₀', T₇₀₀'-T₁₃₀₀' (For T₇' test tube)] Compare T_3'' and T_7 test tubes, both the cases **AB** is common. Step 9: We store only common terms of two minterms others are separated from the test tubes. In common terms, we store only ds DNA part and remaining portions are left out by cutting through restriction enzymes.

Step 10: We will denature the ds DNA and keep complementary part of the common terms in a test tubes.

And the common terms are $\underline{A'B'+B'C'+B'C'+A'C'+A'C'+A'B'+A'B'}$ Step 11: We store the result and check if any common terms are left between the minterms, which we get from step 10. If yes then go to next step otherwise go to step 18.

After denaturation, we get those minterms terms having common terms, so again the same process will be continued. Store the minterms in different test tubes.

Ā'Ē' T0	$\bar{A}^{'C'} \\ T_1$	B' C' T2
Ā'Β' Το'	Ā'В' Т1'	A 'B' T2'
	В ⁻ С' Т₁''	

Figure9: Minterms that consist some common terms after first iteration

Step 12: Then, we generate the strands with combination of 1 variable and separate those combinations in different test tubes.

For T ₀	For T ₀ '	For T _l '	For T ₁	For T ₁ "	For T ₂	For T ₂ '
Ā	Ē	В	С	ē	С	A
ē	Ā	Ā	Ā	В	В	В
T ₃	T ₄	Τ _ó	Τ5	T ₇	T8	T9

Figure10: Combination of 1 variable

Step 13: We have to combine those two minterms which are differ in exactly one bit position. For the above example, we can compare, $T_0 - T_1$, $T_0 - T_1'$, $T_0 - T_1''$ and $T_0' - T_1$, $T_0' - T_1'$ and $T_1' - T_2'$. $T_1''-T_2, T_1-T_2$

The complementary strands of minterms which we get from step 10 are amplified.

AMPLIFY (T₀, T₀₁, T₀₂, T₀₃, T₀₄, T₀₅, T₀₆, T₀₇, T₀₈.)

AMPLIFY (T₀', T₀₁', T₀₂', T₀₃', T₀₄', T₀₅', T₀₆', T₀₇', T₀₈')

AMPLIFY (T₁, T₁₁, T₁₂, T₁₃, T₁₄, T₁₅, T₁₆, T₁₇, T₁₈)

- AMPLIFY (T₁', T₁₁', T₁₂', T₁₃', T₁₄', T₁₅', T₁₆', T₁₇', T₁₈') AMPLIFY (T₁", T₁₁", T₁₂", T₁₃", T₁₄", T₁₅", T₁₆", T₁₇", T₁₈")
- AMPLIFY (T₂, T₂₁, T₂₂, T₂₃, T₂₄, T₂₅, T₂₆, T₂₇, T₂₈)

AMPLIFY (T2', T22', T23', T24', T25', T26, 'T27, T28') Step 14: Now, we have to generate the strands with combination of 1 variable and separate those strands in 2 different test tubes and amplify those test tubes according to the minterms. SEPARATE (T₃, T₃₀) AMPLIFY (T₃, T₃₀, T₃', T₃₀', T₃", T₃₀") SEPARATE (T4, T40) AMPLIFY (T4, T40, T4', T40', T4", T40") SEPARATE (T5, T50) AMPLIFY (T₅, T₅₀, T₅', T₅₀', T₅", T₅₀") SEPARATE (T₆, T₆₀) AMPLIFY (T₆, T₆₀, T₆', T₆₀', T₆", T₆₀") SEPARATE (T7, T70) AMPLIFY (T7, T70, T7', T70', T7", T70") SEPARATE (T_{8.} T₈₀) AMPLIFY (T8, T80, T8', T80', T8", T80") SEPARATE (T₉, T₉₀) AMPLIFY (T9, T90, T9', T90', T9", T90") Now, those two minterms which differ in exactly one bit position are combined. Compare T₀ and T_{1:} MERRGE $[\overline{T}_0 - T_3, T_{01} - T_{30}]$ MERGE [T₁-T₅, T₁₁-T₅₀] Compare T_0 and T_1 test tubes, both the cases \overline{A} is common Compare T₀ and T_{1'}: $\overline{\text{MERRGE}} [\overline{T}_{02} - T_3', \overline{T}_{03} - T_{30}']$ MERGE [T₁'-T₆, T₁₁'-T₆₀] Compare T_0 and T_1 'test tubes, both the cases A is common Compare T₀' - T 1: MERRGE [T₀'-T₄, T₀₁'-T₄₀] MERGE [T₁₂- T₅', T₁₃- T₅₀'] Compare T_0' and T_1 test tubes, both the cases A is common Compare T₀' - T₁': MERRGE [T₀₂'-T₄', T₀₃'-T₄₀'] MERGE [T 12'- T6, T13'- T60] Compare T_0' and T_1' test tubes, both the cases A is common Compare T₁" - T 2: MERRGE [T₁"-T₇, T₁₁"-T₇₀] MERGE [T 2- T8, T21- T80] Compare T_1 "and T_2 test tubes, both the cases **B** is common <u>Compare T₀- T 1'':</u> MERRGE [T₀₄-T₃", T₀₅- T₃₀"] MERGE [T₁₂"- T₇', T₁₃"- T₇₀'] Compare T_1 and T_0 test tubes, both the cases C is common Compare T₂'- T 1': MERRGE $[T_{14}'-T_{6}', T_{15}'-T_{60}']$ MERGE [T₂'- T₉, T₂₁'- T₉₀] Compare T_2 'and T_1 ' test tubes, both the cases is **B** common <u>Compare T₁- T 2:</u> MERRGE [T₁₅-T₅", T₁₅- T₅₀"] MERGE [T₂₂- T₈, T₂₃- T₈₀] Compare T_1 and T_2 test tubes, both the cases C is common Result is: $\overline{A} + \overline{A} + \overline{A} + \overline{A} + B + B + C + C$

After denaturation, we get those minterms terms having common terms, so again the same process will be continued. Store the minterms in different test tubes and repeat the steps as above. The final result will be A+B

5. CONCLUSION

The purpose of this chapter was to show a mathematical application of DNA computing. Hence, we consider DNA as Arithmetic-Logic Unit, where human operators implement biochemistry procedures to perform mathematical operations. The power of the DNA Computing consists in the capability to represent, and compute, huge binary numbers, or highly small ones which are impossible to consider in a conventional computer. In other words, we are able to calculate mathematical operations with unlimited decimal digits. It is worthy noticing that "unlimited" does not mean "endless", but unfixed number of bits. If we consider that 50g of DNA contains 10³³ molecules, it is clear that in few grams of DNA we can encode a great deal of molecular bits. In this paper we introduced a new DNA computing algorithm for reducing any Boolean expression to its simplest form by using DNA strands. This feature is a beautiful remedy for computational problems, which all depend on the fixed number of bits reserved to the representation in conventional computing.

6. REFERENCES

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