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Modeling and Simulation of Genome Evolution Using Linear Boolean Functions Associated with One Dimensional Cellular Automata

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Abstract- Structural and functional behavior of genomes could be studied using one dimensional binary-valued three neighborhood cellular automata updating rules. These updating rules are linear Boolean functions, and they are applied to the adjoint sequences of adenine, (A), Thymine (T), Guanine (G) and Cytosine (C) corresponding to the characteristic sequence of a genome. This paper proposes the use of linear Boolean functions, and demonstrates the textural or fractal behavior of genome evolution in terms of nucleotide adjoints.

Keywords: linear boolean functions, cellular automata, genome evolution.

GJCST-G Classification: J.3



Strictly as per the compliance and regulations of:



Now the adjoints of this sample sequence of length 40 are given below.

A(n) = 0110010010011000011000011000000000000000
 T(n) = 0001001100000000000001000001010011001100
 G(n) = 1000100000000110000100000100000000000011
 C(n) = 0000000001100001100010100010101100110000

A cellular automaton is an idealized parallel processing system consisting of an array of numbers (1-D, 2-D and more) realized using updating rules based on certain neighborhood. For example, a one-dimensional cellular automaton would consist of a finite length array as shown below.

III. CELLULAR AUTOMATA AND LINEAR BOOLEAN FUNCTIONS

A cellular automaton is an idealized parallel processing system consisting of an array of numbers (1-D, 2-D and more) realized using updating rules based on certain neighborhood. For example, a one dimensional cellular automaton would consist of a finite length array as shown below.

---	---	---	i-1	i	i+1	---	---	---
-----	-----	-----	-----	---	-----	-----	-----	-----

Consider an ith cell in the array. This cell has a neighbor i-1 on its left and another i+1 on its right. All three put together is called a three-neighborhood. One can assign a site (cell) variable ξ_{i-1} , ξ_i , and ξ_{i+1} to the three-neighborhood cells. At a particular instant of time, these variables take on numerical values, say either a 0 or a 1. In such a case, the variables are denoted as ξ_{ti-1} , ξ_{ti} , and ξ_{ti+1} . The value of the ith cell at the next instant of time is evaluated using an updating rule that involves the present values of the ith, (i-1)th and (i+1)th cells. This updating rule is essentially a linear Boolean function of three variables. One can construct 256 linear Boolean functions as updating rules of one-dimensional three-neighborhood binary-valued cellular automata. Each rule defines an automaton by itself. So, one-dimensional binary-valued three-neighborhood cellular automata (123CA) rules could be used to model adjoints of a genome sequence. The first thirty linear Boolean functions of cellular automata 123CA are listed below with their decimal equivalents.

Linear Boolean Function

Decimal Equivalent

- 0
- $(\xi_{i-1}\xi_i\xi_{i+1})$
- $(\xi_{i-1}\xi_i\xi_{i+1})$
- $(\xi_{i-1}\xi_i)$
- $(\xi_{i-1}\xi_i\xi_{i+1})$
- $(\xi_{i-1}\xi_{i+1})$
- $(\xi_{i-1}\xi_i\xi_{i+1}) + (\xi_{i-1}\xi_i\xi_{i+1})$
- $(\xi_{i-1}\xi_{i+1}) + (\xi_{i-1}\xi_i)$
- $(\xi_{i-1}\xi_i\xi_{i+1})$
- $(\xi_{i-1}\xi_i\xi_{i+1}) + (\xi_{i-1}\xi_i\xi_{i+1})$
- $(\xi_{i-1}\xi_{i+1})$

- 0
- 1
- 2
- 3
- 4
- 5
- 6
- 7
- 8
- 9
- 10

- $(\xi_{i-1}\xi_i) + (\xi_{i-1}\xi_{i+1})$ 11
- $(\xi_{i-1}\xi_i)$ 12
- $(\xi_{i-1}\xi_{i+1}) + (\xi_i\xi_{i+1})$ 13
- $(\xi_{i-1}\xi_i) + (\xi_{i-1}\xi_{i+1})$ 14
- (ξ_{i-1}) 15
- $(\xi_{i-1}\xi_i\xi_{i+1})$ 16
- $(\xi_i\xi_{i+1})$ 17
- $(\xi_{i-1}\xi_i\xi_{i+1}) + (\xi_{i-1}\xi_i\xi_{i+1})$ 18
- $(\xi_i\xi_{i+1}) + (\xi_{i-1}\xi_i)$ 19
- $(\xi_{i-1}\xi_i\xi_{i+1}) + (\xi_{i-1}\xi_i\xi_{i+1})$ 20
- $(\xi_i\xi_{i+1}) + (\xi_{i-1}\xi_{i+1})$ 21
- $(\xi_{i-1}\xi_i\xi_{i+1}) + (\xi_{i-1}\xi_i\xi_{i+1}) + (\xi_{i-1}\xi_i\xi_{i+1})$ 22
- $(\xi_i\xi_{i+1}) + (\xi_{i-1}\xi_{i+1}) + (\xi_{i-1}\xi_i)$ 23
- $(\xi_{i-1}\xi_i\xi_{i+1}) + (\xi_{i-1}\xi_i\xi_{i+1})$ 24
- $(\xi_{i-1}\xi_i\xi_{i+1}) + (\xi_i\xi_{i+1})$ 25
- $(\xi_{i-1}\xi_i\xi_{i+1}) + (\xi_{i-1}\xi_{i+1})$ 26
- $(\xi_i\xi_{i+1}) + (\xi_{i-1}\xi_{i+1})$ 27
- $(\xi_{i-1}\xi_i\xi_{i+1}) + (\xi_{i-1}\xi_i)$ 28
- $(\xi_i\xi_{i+1}) + (\xi_{i-1}\xi_i)$ 29
- $(\xi_{i-1}\xi_i\xi_{i+1}) + (\xi_{i-1}\xi_i) + (\xi_{i-1}\xi_{i+1})$ 30

IV. CELLULAR AUTOMATA EVOLUTIONS OF GENOME ADJOINTS

The genome sequence of Brucella Suis 1330 is considered here for a case study. Due to space limitations, a part of the genome sequence and its adjoints are shown below. As defined already, adjoint of genome sequence concerning a particular nucleotide is the binary string obtained by marking a '1' in the place of that particular nucleotide and by marking a '0' in the places of other nucleotides. A segment consisting of 60 nucleotides of Brucella Suis 1330 is shown below.

Row Number	Column 1	Column 2	Column 3	Column 4	Column 5	Column 6
1.	GAATGATATAC	CAAGGCCAAG	CTCAAGCTCT	CTCTCTTGG	GCCTGAGCTTT	TGCTTTTGCA
2.	ATTCGTGATCT	TTTTCCTGT	CACGTCTGAC	GAAAGACCA	GCCAGCTGGG	CCCAAGTCGT
3.	CTGTCTGCT	GAGCTCTAG	TCCAGTCTG	AGCAGCTCT	GATCTCTGAC	TCTTTCCT
4.	CTGAGCTCT	TCGCTCTGAC	CTCTGACCT	CTGCTCTG	CTCATGCTCT	TCCTTTCCT
5.	GCCTCTCT	AGGCTCTCT	TGCTCTGAC	GCTCTCTG	CTTGTCTCT	GTCTCTGAC
6.	GTCTCTCT	CTGCTCTG	GCTCTCTG	AGTCTCTG	GCTCTCTG	TCCTCTCT
7.	TCGCTCTG	GCTCTCTG	CTCTCTG	CTCTCTG	CTCTCTG	CTCTCTG
8.	CTCTCTG	GCTCTCTG	CTCTCTG	CTCTCTG	CTCTCTG	CTCTCTG
9.	TGCTCTCT	CTCTCTG	CTCTCTG	CTCTCTG	CTCTCTG	CTCTCTG
10.	TCCTCTCT	CTCTCTG	CTCTCTG	CTCTCTG	CTCTCTG	CTCTCTG
11.	TGCTCTCT	CTCTCTG	CTCTCTG	CTCTCTG	CTCTCTG	CTCTCTG
12.	CTCTCTCT	CTCTCTG	CTCTCTG	CTCTCTG	CTCTCTG	CTCTCTG
13.	AGCTCTCT	CTCTCTG	CTCTCTG	CTCTCTG	CTCTCTG	CTCTCTG
14.	GACTCTCT	CTCTCTG	CTCTCTG	CTCTCTG	CTCTCTG	CTCTCTG
15.	CTCTCTCT	CTCTCTG	CTCTCTG	CTCTCTG	CTCTCTG	CTCTCTG
16.	GCTCTCTG	CTCTCTG	CTCTCTG	CTCTCTG	CTCTCTG	CTCTCTG
17.	GCTCTCTG	CTCTCTG	CTCTCTG	CTCTCTG	CTCTCTG	CTCTCTG
18.	CACTCTCT	CTCTCTG	CTCTCTG	CTCTCTG	CTCTCTG	CTCTCTG
19.	CTCTCTCT	CTCTCTG	CTCTCTG	CTCTCTG	CTCTCTG	CTCTCTG
20.	AGTCTCTG	CTCTCTG	CTCTCTG	CTCTCTG	CTCTCTG	CTCTCTG

The adjoints of the genome sequence segment are given below.

Adjoint A(n)

Row Number	Column 1	Column 2	Column 3	Column 4	Column 5	Column 6
1.	0110010010	0110000110	0001100000	0000000000	0000100000	0000000001
2.	1000000100	0000000000	0100000110	0111101000	0001100000	0001100000
3.	0000000000	0100000110	0001100000	1001100000	0100000000	0100000000
4.	0001000000	0000000100	0000000000	0000000000	0001100000	0000000000
5.	0000000000	1000010000	0000000000	0000000000	0000000000	0000000001
6.	0000000110	0000000001	0000100000	1011000011	0000000000	0000100001
7.	0001000000	0010000000	0001100001	0000000000	0000000001	0100000000
8.	0000011001	0010000000	0000000000	1000000011	0000000000	0000100000
9.	0000000110	0010000000	0001000001	0000000000	0000000100	0100000001
10.	0010010000	0000100011	0000100000	0000000010	0000000010	1001001100
11.	0000000000	0000000000	0000000000	0000000000	0100000000	1101100000
12.	0000001000	0000000000	0000001010	0000000000	0000000010	0000000000
13.	1010001100	1101011001	0000000000	0000100011	0000000000	0001000000
14.	0100001000	0001011000	0000100000	0000010000	0000010000	0000000000
15.	0000000001	0000000011	0001010001	0100000110	1000001000	0100001000
16.	0000000101	0000000000	0000000000	0001100000	0000001001	0100100000
17.	0000000001	0010010000	0100000000	0000010000	0000010000	1111000010
18.	0110101000	0000000101	0000000000	0100100111	0010000100	0000011000
19.	0000100101	0000000001	0001101000	0001000100	0000100000	0000001000
20.	1001000000	0010000100	0000001011	1000111001	1000000011	0001010000

Adjoint T(n)

Row Number	Column 1	Column 2	Column 3	Column 4	Column 5	Column 6
1.	0001001100	0000000000	0100000101	0011001100	0010000111	0000111000
2.	0110010011	1111110101	0001001000	0000000000	0000000000	0000001001
3.	0101010001	0000011000	1000001000	0000000001	0010010000	0001110001
4.	0000001011	0001000001	0101000001	0100010100	0101000000	0101011001
5.	0000001000	0000000010	1010000000	0000000000	0010010111	0110100000
6.	0010000000	0010000000	0100001110	0000001000	0010110010	0000001000
7.	1000010001	0000010000	0000000000	0000001000	0010010000	0000000000
8.	0001000000	0000110000	0000000000	0000000000	0000000010	0100000000
9.	1000000001	0000000010	1000000000	0001000000	0010000010	0001110000
10.	1000000000	0110001000	0000000000	0100000000	0100000000	0100000000
11.	1000100000	0001001000	1100000101	1101000100	0001101001	0000001011
12.	0000000000	1011000110	1000000000	0000000010	0000111000	1100000000
13.	0000000001	0000000000	0000100000	1000010000	0000101110	0000001100
14.	0000000000	0100000100	0010000010	0111000001	0011000011	0001000000
15.	0000010100	0010010000	0000000000	0000010000	0100110101	0010010000
16.	0000100000	0100000000	0010100000	1100000000	0001000000	0000000101
17.	0001000100	0000000011	0000001000	0001000111	0011000100	0000100000
18.	0000000000	0011010000	0000000000	0000001000	0000110010	0010000000
19.	0010000000	0000010000	0000000000	0000100000	0011000100	0111010111
20.	0010000011	1000000000	0000100000	0000000000	0100000000	0100000100

Adjoint G(n)

Row Number	Column 1	Column 2	Column 3	Column 4	Column 5	Column 6
1.	1000100000	0001100001	0000010000	0000000011	1001010000	0100001000
2.	0000100000	0000000010	0000100100	1000100000	1000100111	0000010010
3.	0010001000	1010000101	0000100010	0100010000	1001000110	0000000000
4.	0010101000	0011000010	0000001000	0010000001	0000001000	0000000000
5.	1000000000	0110000000	0100100010	1100000110	0001010000	1001011000
6.	1000101001	0101011000	1010100001	0100011010	1101000001	0000001000
7.	0010100000	1100101111	0000001000	0001010111	0000100000	0001000101
8.	0010100100	1101000001	0010111000	0011000000	1100101001	0010011001
9.	0101010000	0001000001	0000011110	0000100111	0001100000	0010001000
10.	0000001110	0000000100	1001011101	1001010000	1000000100	0000100011
11.	0101010001	0000100111	0000010010	0010010000	0000010100	0010000000
12.	1001100010	0000001100	0000010000	0000010101	1001000000	0000111000
13.	0100000000	0010100010	0110010010	0111000100	0000000000	0000000000
14.	1000001110	0000000011	1101010001	0000001110	1100100000	1100000001
15.	0000100010	1000000010	0100001010	1011101001	0010000010	1001100011
16.	1101011000	0010010011	0001000001	0000001000	1100000000	0000010010
17.	1000000010	1001101000	0010001010	1000001000	0000000000	0000001101
18.	0000000101	0000000010	1110010011	0011000000	0101000000	0100100110
19.	1000001000	1010000010	0000000110	0000001011	0100010010	1000100000
20.	0100010000	0101111101	1110000100	0010000100	0010000000	1000000011

Adjoint C(n)

Row Number	Column 1	Column 2	Column 3	Column 4	Column 5	Column 6
1.	0000000001	1000011000	1010001010	1100010000	0100001000	0011000010
2.	0001001000	0000001000	1010010001	0000001110	0110011000	1110000100
3.	1000100110	0001100000	0110000001	0010011110	0000010001	0110000110
4.	1100010010	0100001001	1011000110	1000111000	1010010111	0101000110
5.	0111101111	0000010101	0000100001	0011101000	1100000000	0000000000
6.	0101010100	1000100110	0000000000	0000000000	0000001100	0111000010
7.	0100001110	0001000000	1110000110	1110000000	1100001001	0010100100
8.	1100000010	0000001110	1001000111	0100010100	0011010100	1001000110
9.	0001010100	1100110100	0110100000	1100000100	1100001001	1000000010
10.	0110100001	1001000000	0110000010	0010101100	0011110001	0010010000
11.	0010001110	1110010000	0011010000	0000101011	1010000100	0000010100
12.	0110010101	0100000001	0110101001	1111010000	0110000001	0010000011
13.	0000110010	0000000100	1001000101	0000000100	1111010000	1110110011
14.	0011110001	1010100000	0000010000	0000010000	0000001100	0010111110
15.	1111001000	0101010000	1010100100	0000000100	0001000000	0000000000
16.	0010000010	1001100100	1100011010	0000011111	0010101100	1011000000
17.	0110110000	0100000100	1001110001	0110100000	1101010001	0000000000
18.	1001010010	1100010000	0001010000	1000010000	1000000100	0010000001
19.	0101001010	0110110000	1110010001	1110000000	1000000000	0000000000
20.	0000101100	0000000000	0000101000	0101000010	0000111100	0010101000

The size of the images shown in Fig. 2 is 500x500, though the actual size is 5806x500. The first 500 columns of the actual images are clipped and presented here for visual clarity. From Fig. 2, it is clear that the evolution pattern of each adjoint is different. One can observe that there are certain fractal patterns in the evolutions and such fractals are distributed in the images very differently. For instance, the zoomed in versions of the evolution patterns of A(n), T(n), G(n) and C(n) using rule 137 are shown in Figs. 3, 4, 5 and 6 respectively.

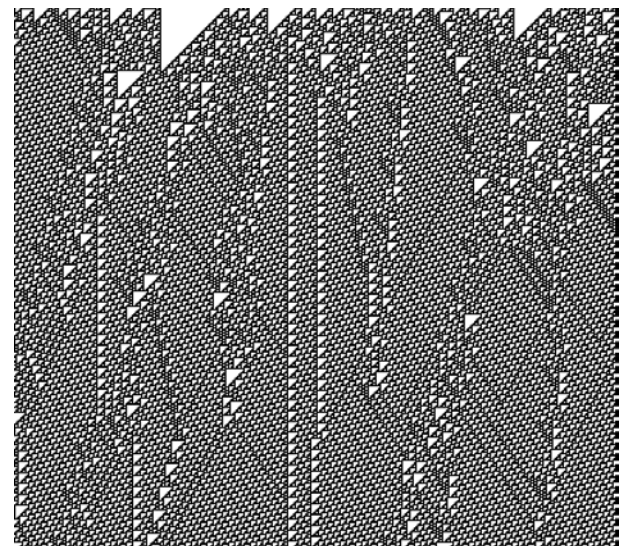


Fig. 3: Zoomed in version of evolution pattern of A(n)

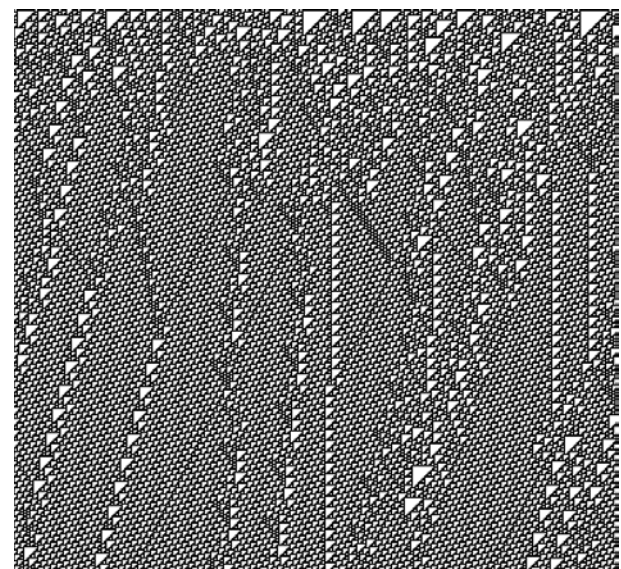


Fig. 4: Zoomed in version of evolution pattern of T(n)

Cellular automata evolutions of adjoints of a genome are carried out using 256 rules of 123CA. As an example, rule number 137 of 123CA, that is, $(\xi_{i-1}\xi_i\xi_{i+1}) + (\xi_i\xi_{i+1})$ is applied to adjoints of Brucella Suis 1330 genome and results shown below in Fig. 2.

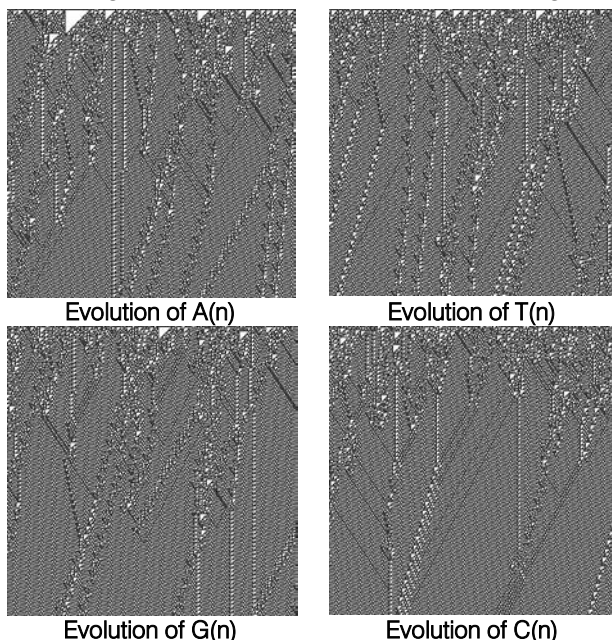


Fig. 2: Evolution of adjoints using rule 137 of 123CA

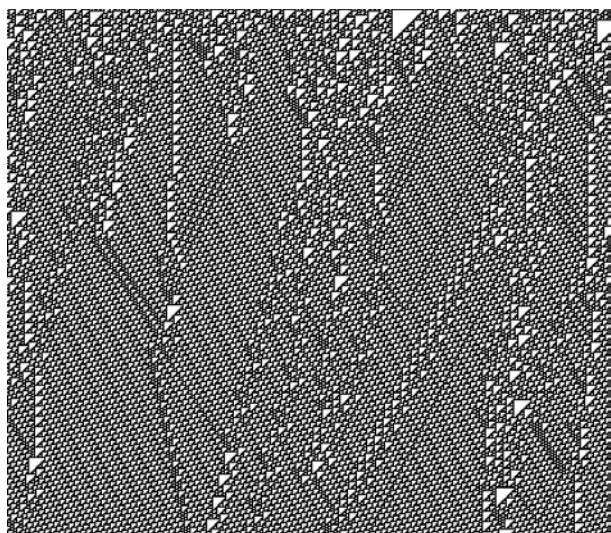


Fig. 5: Zoomed in version of evolution pattern of G(n)

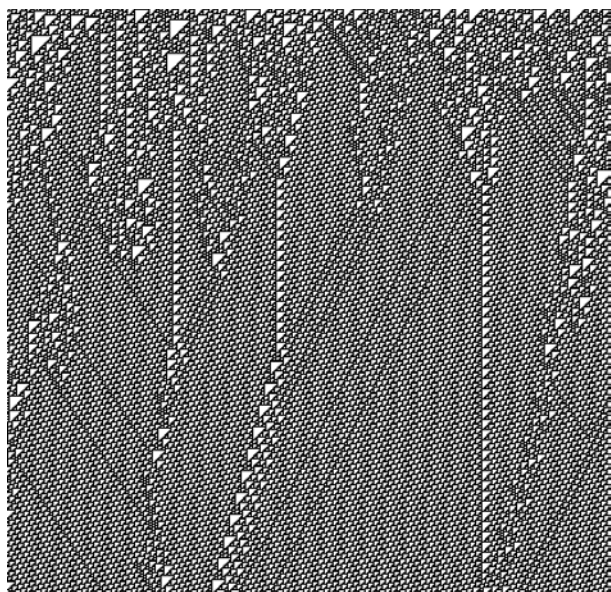
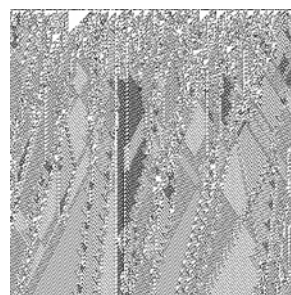


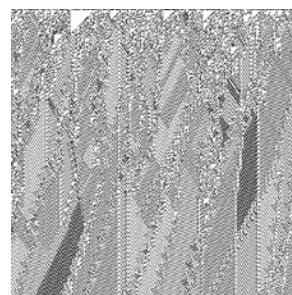
Fig. 6: Zoomed in version of evolution pattern of C(n)

V. DYADIC OPERATIONS BETWEEN CELLULAR AUTOMATA EVOLUTIONS OF GENOME ADJOINTS

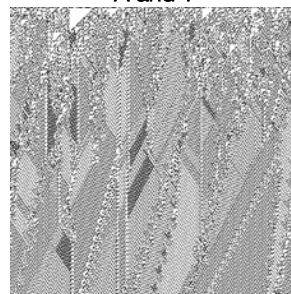
It is a matter of an empirical study to examine the conjoint behavior of various evolution patterns of adjoints and it could be carried out by combining evolution patterns of adjoints dyadically. The various dyadic operations are (i) Boolean addition, (ii) Boolean subtraction, (iii) Boolean multiplication, (iv) Boolean division, (v) Dyadic relation of maximum and (vi) Dyadic relation of minimum. Out of these six different dyadic operations and relations, the Boolean operation of binary addition is considered here for the intended study.



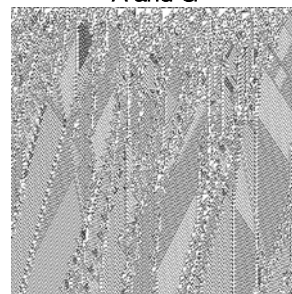
Addition of patterns of A and T



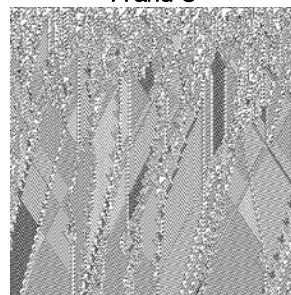
Addition of patterns of A and G



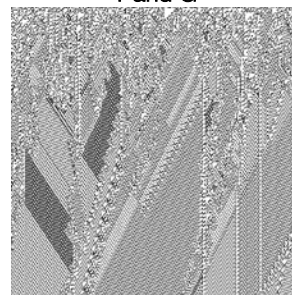
Addition of patterns of A and C



Addition of patterns of T and G



Addition of patterns of T and C



Addition of patterns of G and C

Fig. 7: Boolean addition of evolution patterns of adjoints

The zoomed in versions of the Boolean additions of evolution patterns of A(n), T(n), G(n) and C(n) using rule 137 are shown in Figs. 7, 8, 9 and 10 respectively.

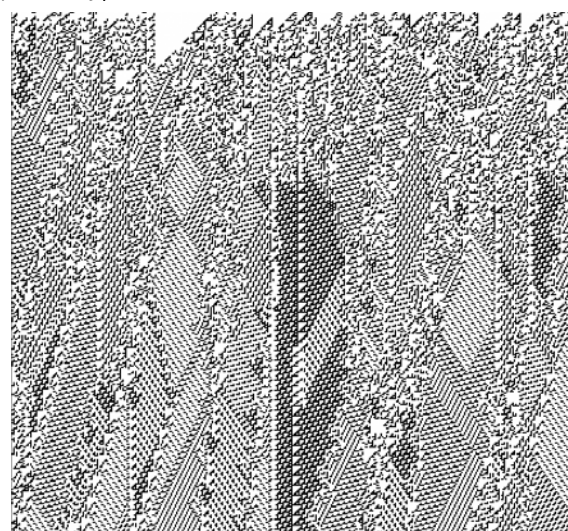


Fig. 8: Zoomed in version of addition of patterns of A and T

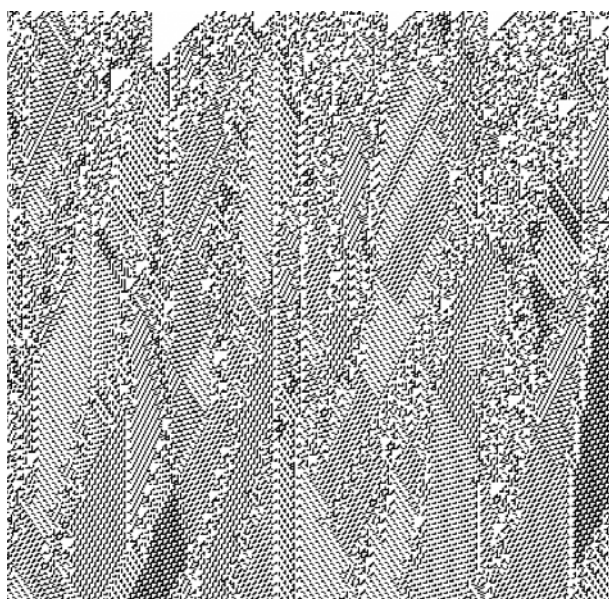


Fig. 9: Zoomed in version of addition of patterns of A and G

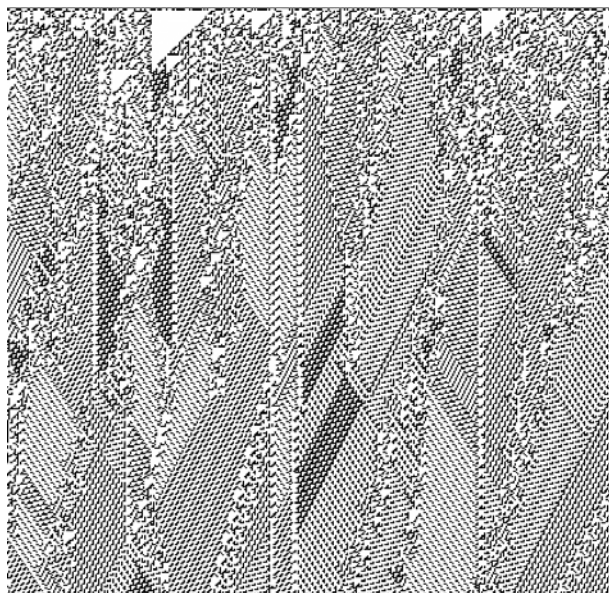


Fig. 10: Zoomed in version of addition of patterns of A and C

VI. OBSERVATIONS AND CONCLUSIONS

From the above empirical study, it is observed that cellular automata modeling and simulation of evolutions of adjoints of a given genome sequence and the inter-pattern operations and relations exhibit distinct patterns of fractals and fractal distributions. The novel technique and results presented in this paper are outcome of prolonged research carried out in the mathematical modeling of genomes and their evolutions. It is evident that one can as well look into the possibilities of genome editing using such cellular automata tools.

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