



Science

## WUHAN COVID-19 SYNTHETIC ORIGINS AND EVOLUTION

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### Abstract

The main result of this updated release is the formal proof that 2019-nCoV coronavirus is partially a SYNTHETIC genome. We proof the CONCENTRATION in a small région of wuhan New genome (300bp) of 3 different régions from HIV1 ENVELOPPE gene and 3 others from HIV2 and SIV (ENV and POL RT). All this is remarkable and bears the mark of a desire for organization of a human nature: LOGIC, SYMETRIES.

In this article, we demonstrate also that there is a kind of global human hosts adaptation strategy of SARS viruses as well as a strategy of global evolution of the genomes of the different strains of SARS which have emerged, mainly in China, between years 2003 first SARS genomes and the last 2019 COVID-19 Wuhan seafood market pneumonia virus isolate Wuhan-Hu-1, complete genome.

This global strategy, this temporal link, is materialized in our demonstration by highlighting stationary numerical waves controlling the entire sequence of their genomes.

Curiously, these digital waves characterizing the 9 SARS genomes studied here are characteristic whole numbers: the "Fibonacci numbers", omnipresent in the forms of Nature, and which our research for several decades has shown strong links with the proportions of nucleotides in DNA.

Here we demonstrate that the complexity and fractal multiplicity of these Fibonacci numerical waves increases over the years of the emergence of new SARS strains.

We suggest that this increase in the overall organization of the SARS genomes over the years reflects a better adaptation of SARS genomes to the human host.

The question of a link with pathogenicity remains open.

However, we believe that this overall strategy for the evolution of the SARS genomes ensures greater unity, consistency and integrity of the genome.

Finally, we ask ourselves the question of a possible artificial origin of this genome, in particular because of the presence of fragments of HIV1, HIV2 and SIV retroviruses.

**Keywords:** SARS; Wuhan COVID-19; Fibonacci Numbers; Fractal Genome; Numerical Stationary Periodic Waves; HIV1; HIV2; SIV; Synthetic Genomes.

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### 1. Introduction

*“Where there is matter, there is geometry.”*

- Johannes Kepler

Since the SARS coronavirus emergence 18 years ago, a large number of severe acute respiratory syndrome related coronaviruses (SARSr-CoV) have been discovered in their natural host, bats. Some of those bat SARSr-CoVs have the potential to infect humans. In January 2020, Wuhan China megacity was the origin of a Novel SARS disease entitled by OMS « COVID-19 » [6]. This novel coronavirus (COVID-19) caused an epidemic of respiratory syndrome in humans, in Wuhan, China then in other World countries (Iran South Korea, Italia...). As show in the following figures, we analyse in this article 9 whole SARS genomes with the goal to discover a possible strategy of SARS GENOMES EVOLUTION from 2003 original viruses to the 2020 WUHAN virus.

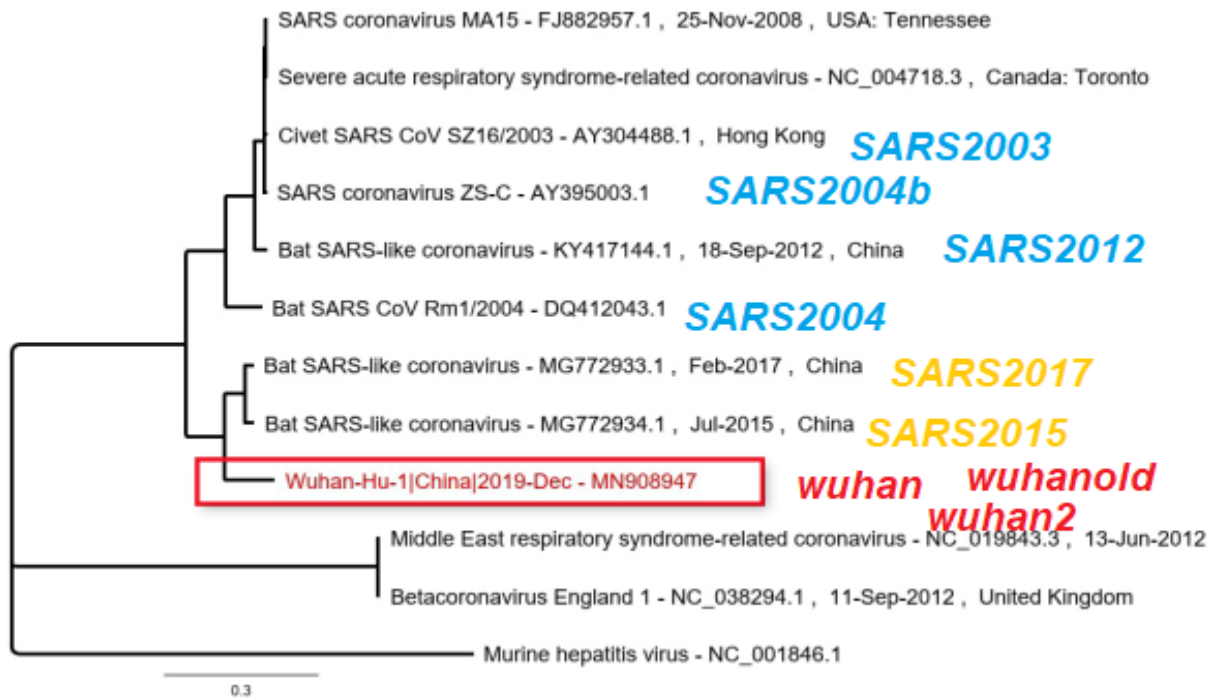


Figure 1: From Ncbiinsights 2020. Coloured References are our 9 Analysed SARS/Wuhan Genomes.

Secondly, for about 30 years, we have been looking for possible global, even digital, structures that would organize DNA, genes, chromosomes, and even whole genomes [27, 28, 30, 50, 51].

however, it is only by deepening the notion of "fractal periodicity", outlined in [7, 38], and we will highlight here that we have re-discovered the major role of **Fibonacci fractal stationary waves** at both scales of each whole individual chromosomes and whole genome. We then demonstrate a sort of "hierarchical classification" of the 24 chromosomes. In this hierarchy, the chromosome4 seems to play a major and privileged role.

By comparing chromosome, the 3 reference genomes of Neanderthal, Sapiens BUILD34 of 2003 and Sapiens HG38 of 2013, we demonstrate the evidence of « fractal periods » and « Resonance periods» characterizing each of the 24 human chromosomes [47]. As illustrated in Figure1 below, these resonances make it possible to differentiate the respective genomes of Neanderthal and Sapiens on the global scale of the chromosome (here chromosome 4). Here, a resonance of 34 nucleotides is common to both chromosomes 4 of Sapiens and Neanderthal, however, the respective forms of these resonance curves are radically different.

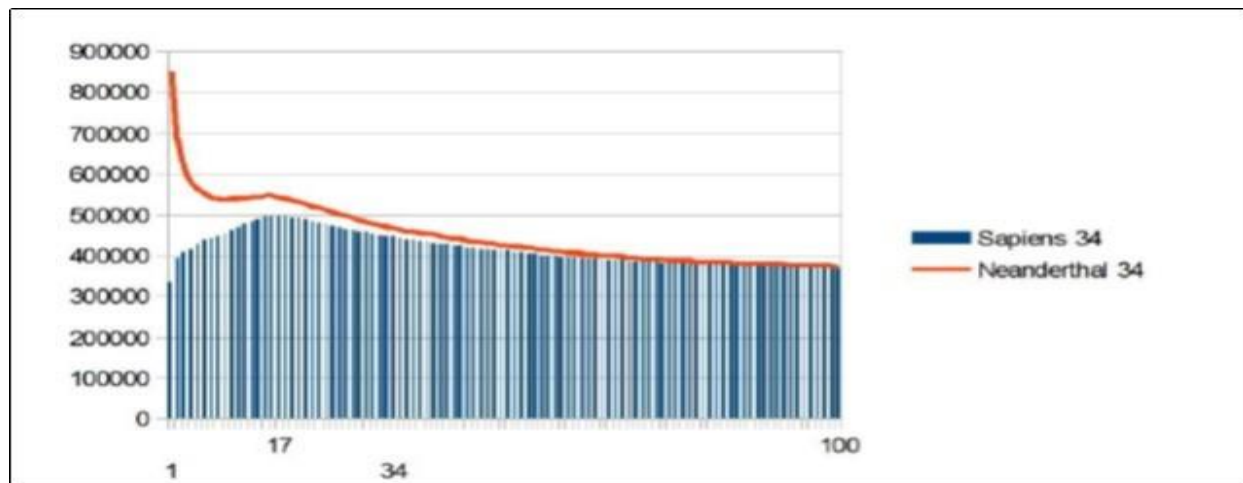


Figure 2: As will be demonstrated here, the 2 respective Chromosomes 4 of Neanderthal and Sapiens HG38 share a "resonance" of 34 bp, however, these two radically different resonance curves illustrate a major differentiation of the 2 human species at the GLOBAL scale of chromosome 4.

## 2. Methods

### The 9 Analysed Genomes:

- SARS2003 SARS Coronavirus SZ16, Complete Genome  
<https://www.ncbi.nlm.nih.gov/nuccore/Ay304488>
- SARS2004 Bat SARS Coronavirus Rm1, Complete Genome  
<https://www.ncbi.nlm.nih.gov/nuccore/DQ412043>
- SARS2004b SARS Coronavirus ZS-C, Complete Genome  
<https://www.ncbi.nlm.nih.gov/nuccore/AY395003>
- SARS2012 Bat SARS-Like Coronavirus Isolate Rs4084, Complete Genome - Nucleotide - NCBI. Ky417144.1  
<https://www.ncbi.nlm.nih.gov/nuccore/KY417144.1>
- SARS2015 Bat SARS-Like Coronavirus Isolate Bat-SL-Covzxc21, Complete Genome  
<https://www.ncbi.nlm.nih.gov/nuccore/Mg772934>

- SARS2017 Bat Sars-Like Coronavirus Isolate Bat-SI-Covzc45, Complete Genome  
<https://www.ncbi.nlm.nih.gov/nuccore/Mg772933>
- WUHANOLD (first genome sequenced 12 January 2020) Wuhan Seafood Market Pneumonia Virus Isolate Wuhan-Hu-1, Complete Genome  
<https://www.ncbi.nlm.nih.gov/nuccore/MN908947.1>
- WUHAN2 (second improved sequenced genome 14 January 2020) Wuhan Seafood Market Pneumonia Virus Isolate Wuhan-Hu-1, Complete Genome Genbank: MN908947.2  
<https://ncbiinsights.ncbi.nlm.nih.gov/2020/01/13/novel-coronavirus/>  
<https://www.ncbi.nlm.nih.gov/nuccore/MN908947.3?report=fasta>
- WUHAN (23 January 2020) Wuhan Seafood Market Pneumonia Virus Isolate Wuhan-Hu-1, Complete Genome GenBank: MN908947.3  
<https://www.ncbi.nlm.nih.gov/nuccore/MN908947.3>

### Computing Fractal Periods and Resonances Summary

The complete description of this method can be found in [47], Six Fractal Codes of Biological Life: perspectives in Exobiology, Cancers Basic Research and Artificial Intelligence Biomimetism Decisions Making. Preprints 2018, 2018090139 (doi: 10.20944/preprints201809.0139.v1).  
<https://www.preprints.org/manuscript/201809.0139/v1>

We introduce here a method of global analysis of the roughness or fractal texture of the DNA sequences at the chromosome scale. To do this, we generalize the method of numerical analysis of the "Master Code of Biology" [39-42, 44]. Thus, we restructure the sequence into different generic sequences based on "meta codons" no longer triplets of 3 nucleotides but values ranging from 17 to 377 nucleotides, i.e 360 simulations. This method of analysis will then reveal, in most cases, discrete waves or interferences, most often dissonances. However, sometimes there will emerge kinds of resonances where all scales of analysis appear to be in symbiosis.

**Function:** The Genomics master code (-II-) is generalized to meta-codons that no longer have 3 nucleotides as a codon, but 4, 5, ... 377 nucleotides. Then we analyze the textures by the undulatory code (-IV-). It then appears dissonances and resonances that will reveal periods of discrete waves, resonances, and standing waves. The Genomics Binary code analysis (-III-) confirms these periods using a complementary independent method.

**Inputs:** Double strand DNA sequence Pi-mass grouped by meta-codons (each Pi-mass is = -1 times number of « G » bases in meta-codon double strand or also = -1 times number of « C+G » bases in single strand meta-codon).

**Outputs:** Period and resonance standing wave computed by two complementary methods.

**Summary:** We introduce here a method of global analysis of the roughness or fractal texture of the DNA sequences at the chromosome scale. To do this, we generalize the method of numerical analysis of the "Master Code" [47 -II- ]. Thus, we restructure the sequence into different generic sequences based on "meta codons", no longer triplets of 3 nucleotides, but values ranging from 17 to 377 nucleotides, ie 360 simulations. This method of analysis will then reveal, in most cases, discrete waves or interferences, most often dissonances (based on Genomics Undulatory waves described here in [47 -II- ]. However, sometimes there will emerge kinds of resonances where all scales of analysis appear to be in symbiosis.

**Process:** The discrete interferences fields resulting from the analysis of an entire chromosome are therefore a three- dimensional space: Dim y (vertical) restructuring in meta codons of lengths 17 to 377 nucleotides (or in this Coronavirus article meta codons 1bp to 100bp) Dim x (horizontal) Leibnitz differentiations such that primary 1/2 secondary 1/3... 1/4 ... 1 / n Dim z cumulated populations from the "Master code" operators. The + 1 / -1 derivatives will be of type increase, ie +1 if derivative increasing and will be of type decrease, ie -1 if derived decreasing. In this context we will explore these 3D spaces in 2 forms:

- Horizontally [47 -IV-], meta codons dimension: curves for a given meta codon dimension, see in the example "resonances" below (see **Figure 3** and **Figure 4**).
- Vertically [47 -III-], spectral differentiation: discrete series d2-d1 is +1 if increase and -1 if decrease (see **Figure 5**). We represent in top the +1 and in low the -1, (see **Figure 5**).

Table 1: Computing the periodic standing waves and resonances for various metacodons  
 Genomics Master code.

Dim y	Dim x	d1	d2	.../... d100		
	0	1	2	3	4	5
	17	1298833	1181005	1133041	1103633	1087486
	18	1029171	1074033	960839	1000920	1028712
	19	1091521	982429	937709	912626	975473
	20	878537	903906	914801	922094	927631
	21	933380	834734	893561	848319	885361
	22	761233	774174	779102	783714	786854
	23	809977	837877	764596	791545	755377
	24	852758	779786	750287	735631	726226
	25	710190	727911	736109	742027	699579
	.../... 377					

Horizontal scan: exp. meta codons of 22 bases: 22 761233 774174 779102 783714 786854 .../... (see **Figure 3**)

Vertical scan example derivations of first order: 1 if d2>d1 and -1 if d2<d1 then: -1 1 -1 1 -1 1 1 -1 1 -1 1 1 .../... (see **Figure 4** and **Figure 5**).

### HG38 Human Reference Chromosome21

Computing PERIODS by Fractal "Increase/Decrease" Textures

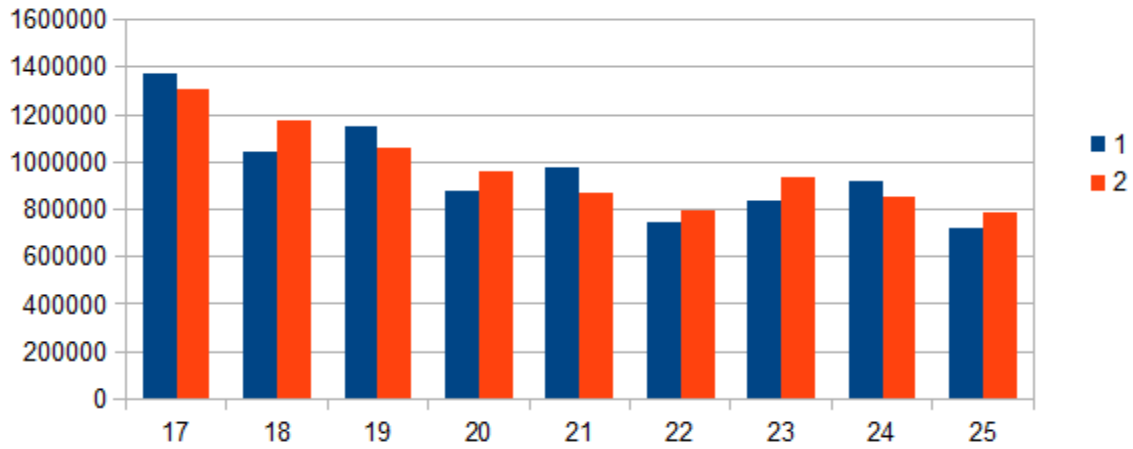


Figure 3: Zoom on Vertical Scan Method Revealing PERIOD = 22 From HG38 Reference Chromosome21.

These two independent methods lead in all the cases analyzed to the same period value: here, for example, the period "horizontal scan" is a resonance of 22bp (**Figure 4**) and the period "vertical scan" is a period of repeatability of 22bp also (**Figure 5**).

### CHR21 HG38 reference chromosome

resonance 22bp

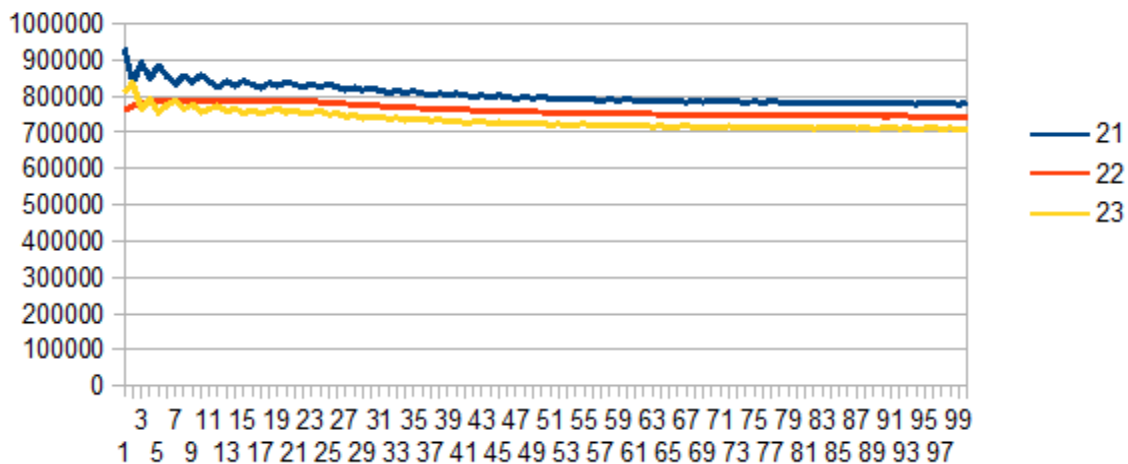


Figure 4: Evidence of a resonance of 22bp period in the whole HG38 human reference chromosome21.

(see **Figure 4** and **Figure 5**), [47 -IV-].

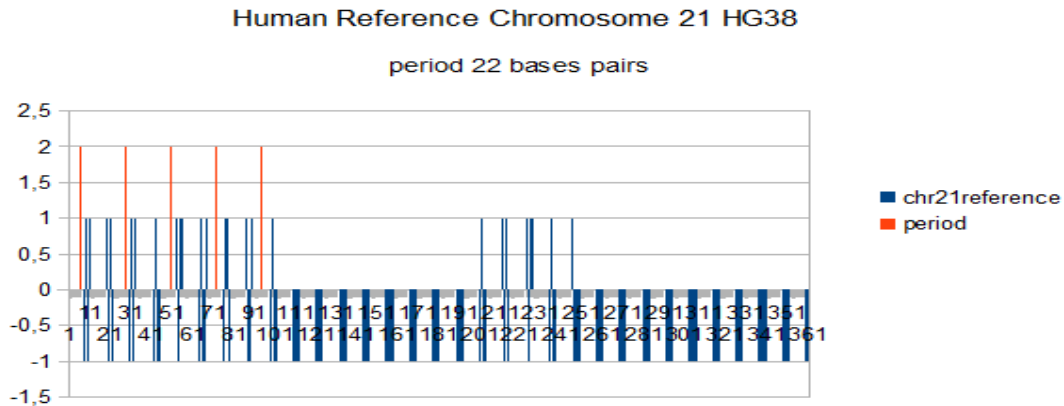


Figure 5: Confirmation of a 22bp period in the whole HG38 human reference chromosome21 [47-IV-].

A third complementary method is presented here: knowing the period determined and confirmed by the two previous methods, we segment the complete sequence of the chromosome by consecutive segments according to this period, for example here for the chromosome21, we will "cut" the entire sequence of the chromosome in successive sections of 22 bases, the length of the period discovered. Then we record for each segment the C + G populations on the one hand and T + A on the other hand. We then represent the cumulative distribution curve of these different CG and TA populations throughout the chromosome sequence.

We then represent the cumulative distribution curve of these different CG and TA populations throughout the chromosome sequence (Table2).

Table 2: This table shows a C+G top for 8 bases value within 22 bases segments distribution.

7	8	9
205735	230173	219804
46083	75340	106183

The **Figure 6** shows a C+G top for 8 bases value within 22 bases segments distribution. segmented by 22 bases periods.

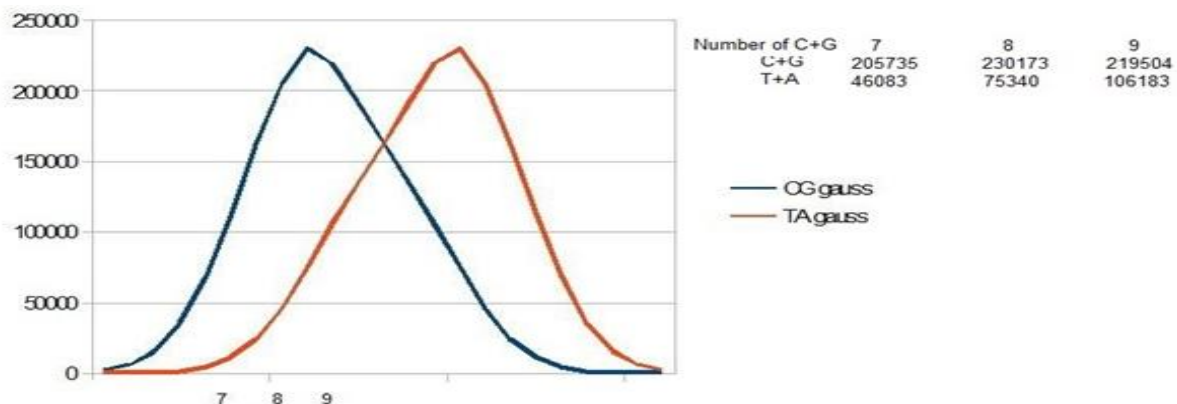


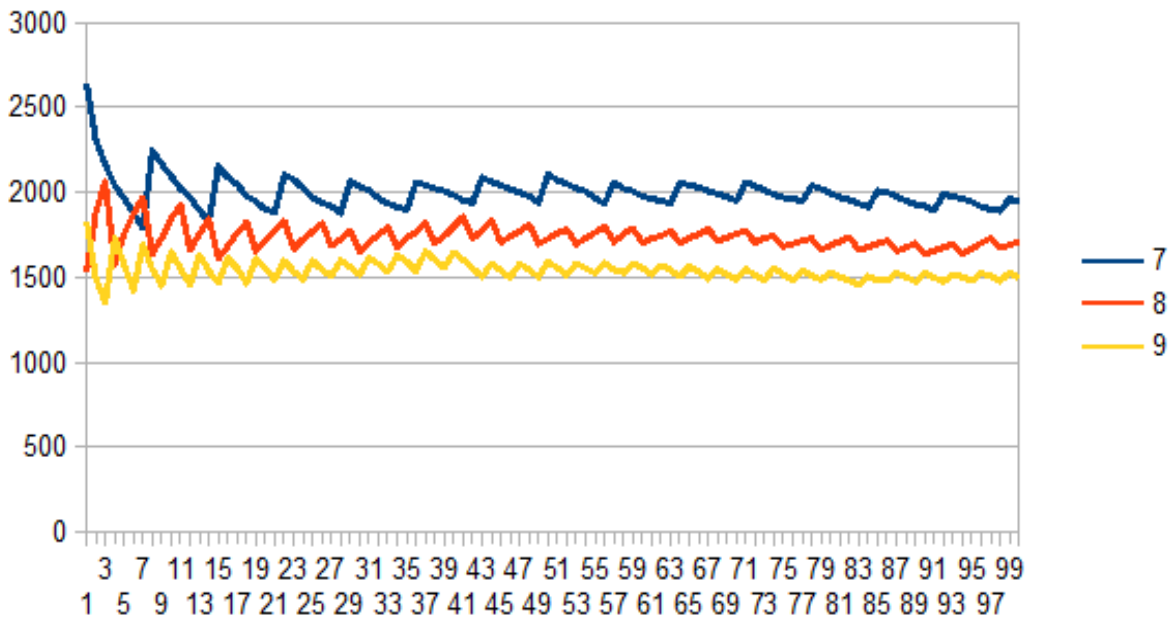
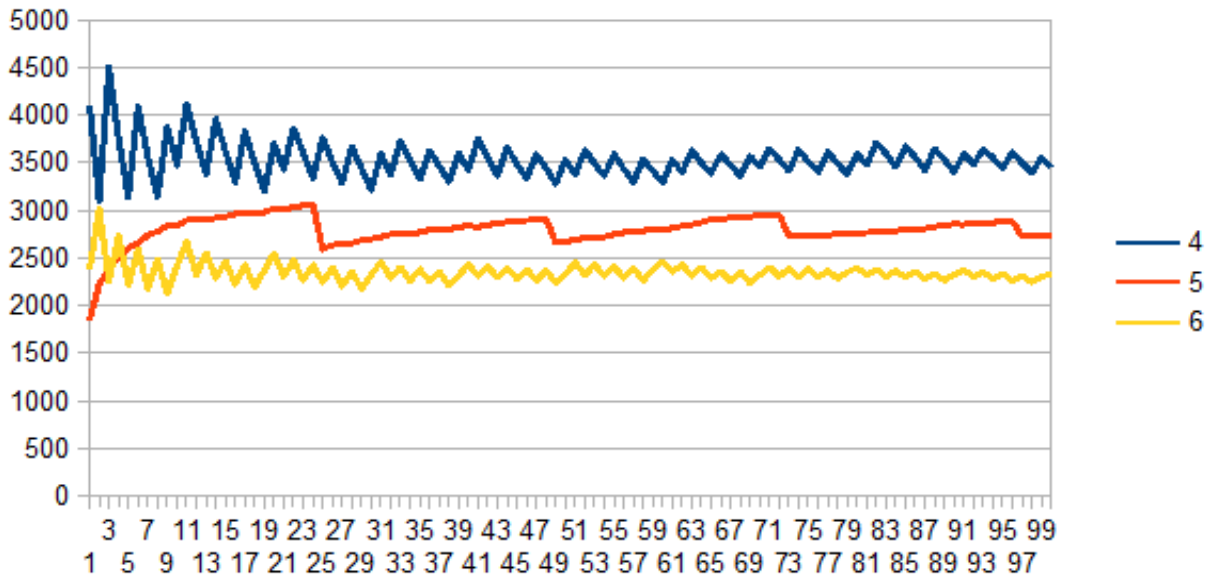
Figure 6: Gauss like CG / TA distribution within the whole human HG38 chromosome21.

### 3. Results and Discussion

- SARS2003

#### SARS Coronavirus SZ16, Complete Genome

<https://www.ncbi.nlm.nih.gov/nuccore/Ay304488>





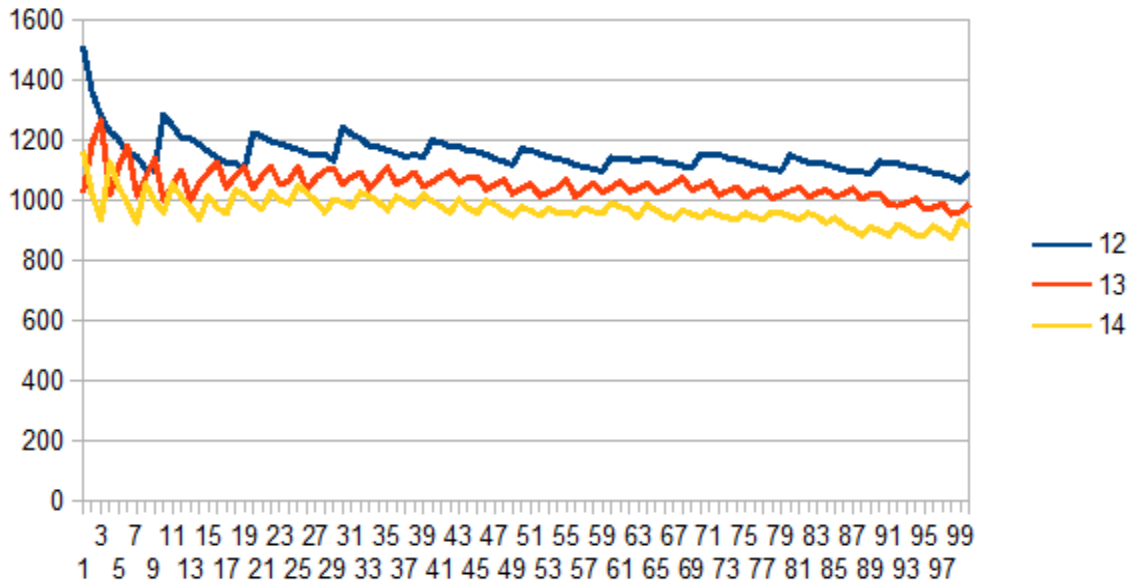


Figure 9: Stationary wave 13bp OFF

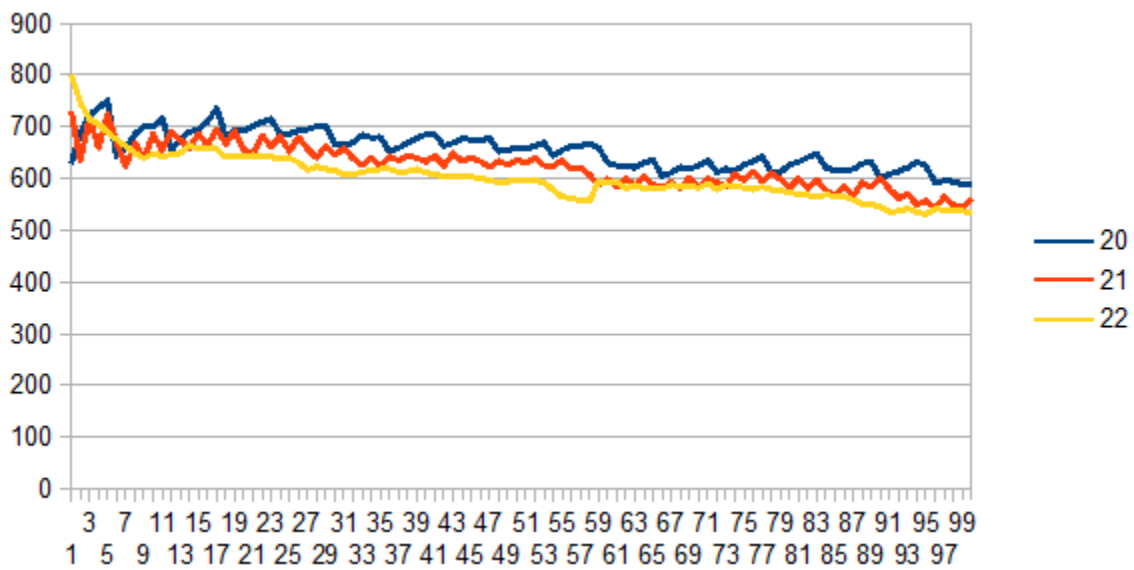


Figure 10: Stationary wave 21bp OFF

In this initial SARS2003 genome, we find only the Fibonacci stationary wave 5bp (Figure 7). All other fractal waves 8,13, 21 bp are absent.

- SARS2004

### Bat SARS coronavirus Rm1, complete genome

<https://www.ncbi.nlm.nih.gov/nuccore/DQ412043>

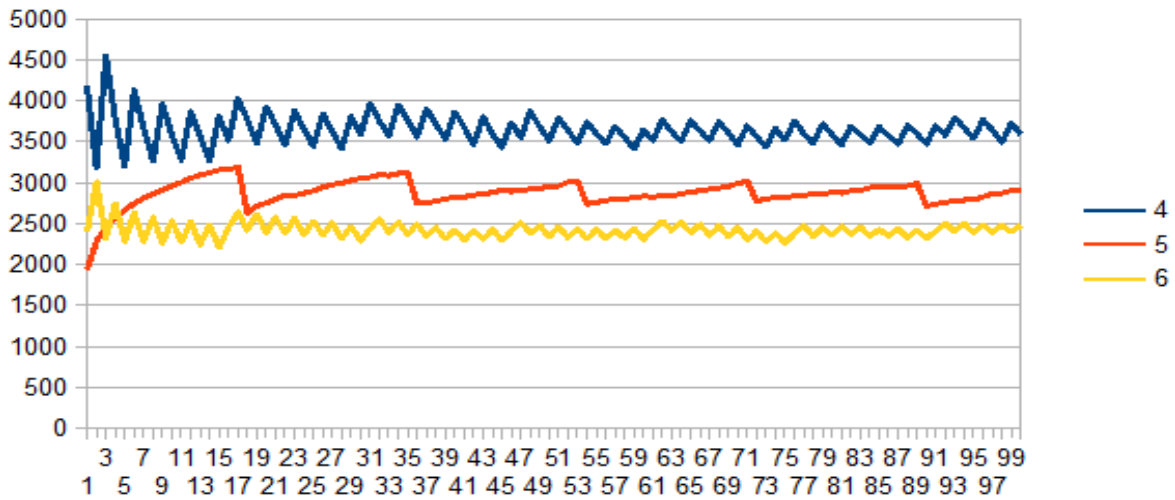


Figure 11: Stationary wave 5bp ON

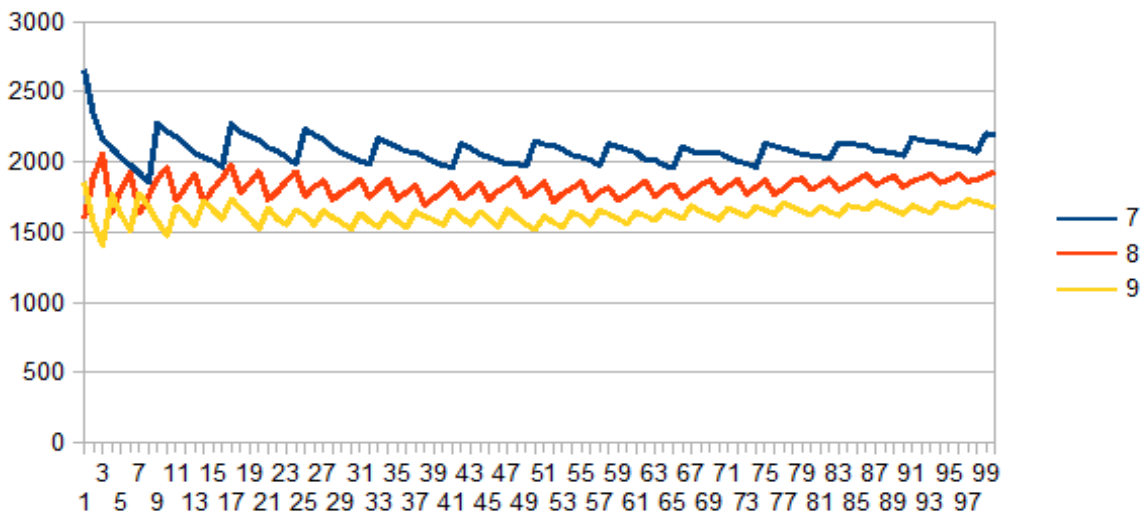


Figure12: Stationary wave 8bp OFF

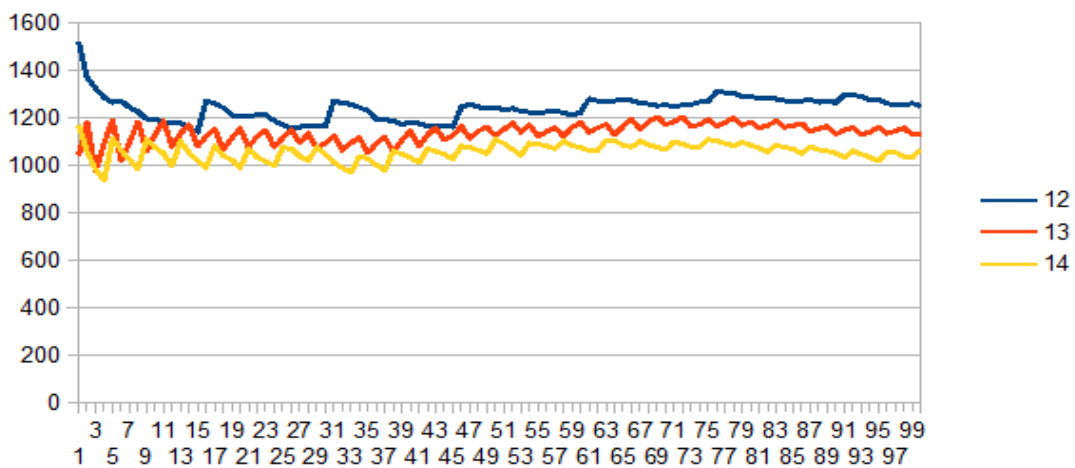


Figure 13: Stationary wave 13bp OFF

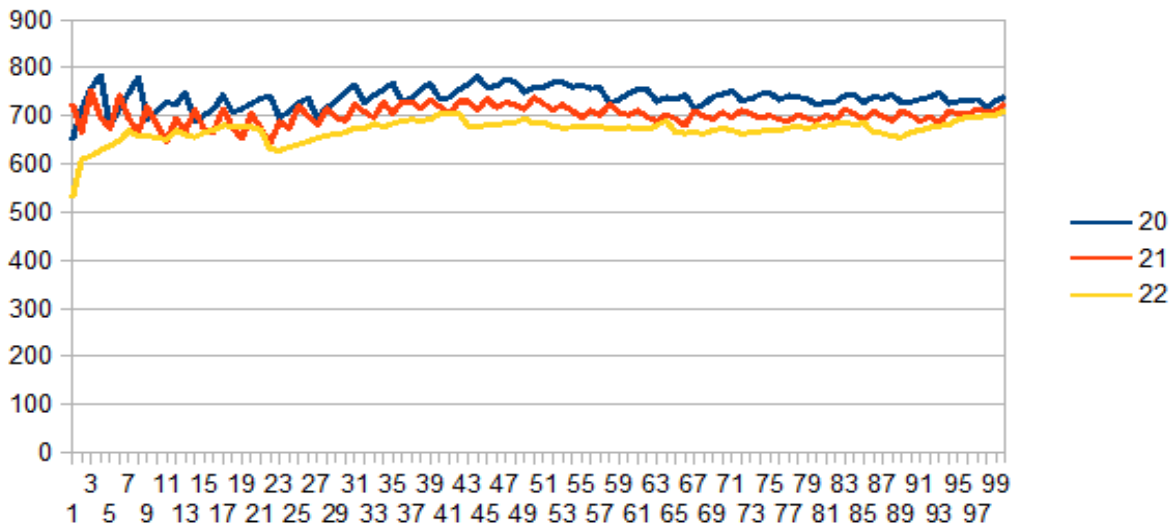


Figure 14: Stationary wave 21bp OFF

In this second SARS2004 genome, we find only the Fibonacci stationary wave 5bp (Figure 11). All other fractal waves 8,13, 21 bp are absent.

- SARS2004b

**SARS Coronavirus ZS-C, Complete Genome**

<https://www.ncbi.nlm.nih.gov/nuccore/AY395003>

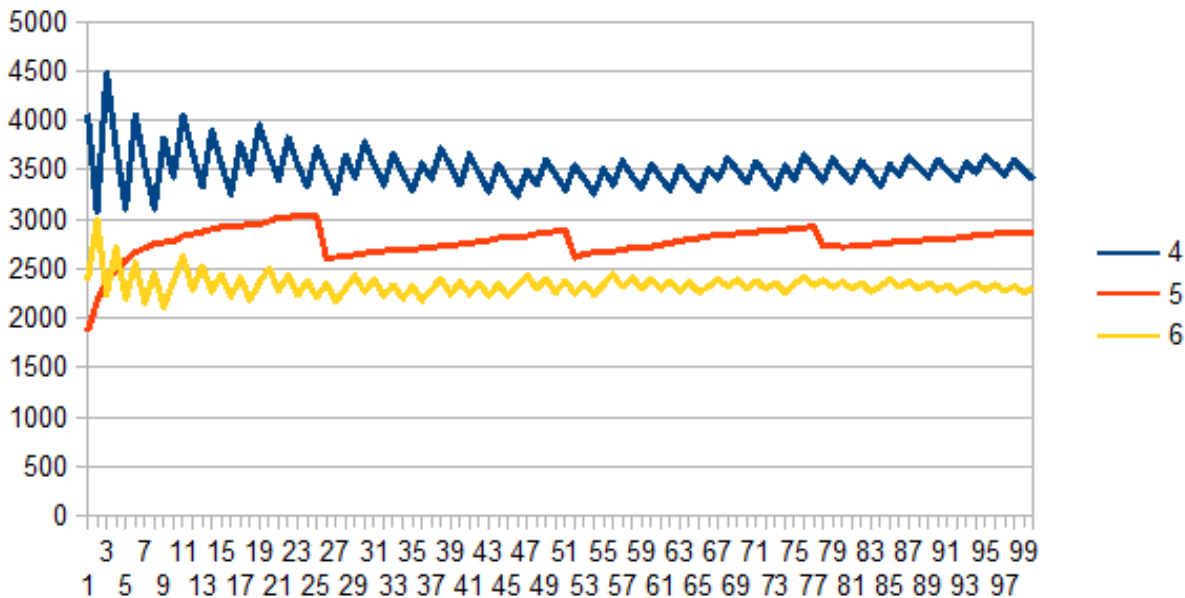


Figure 15: Stationary wave 5bp ON

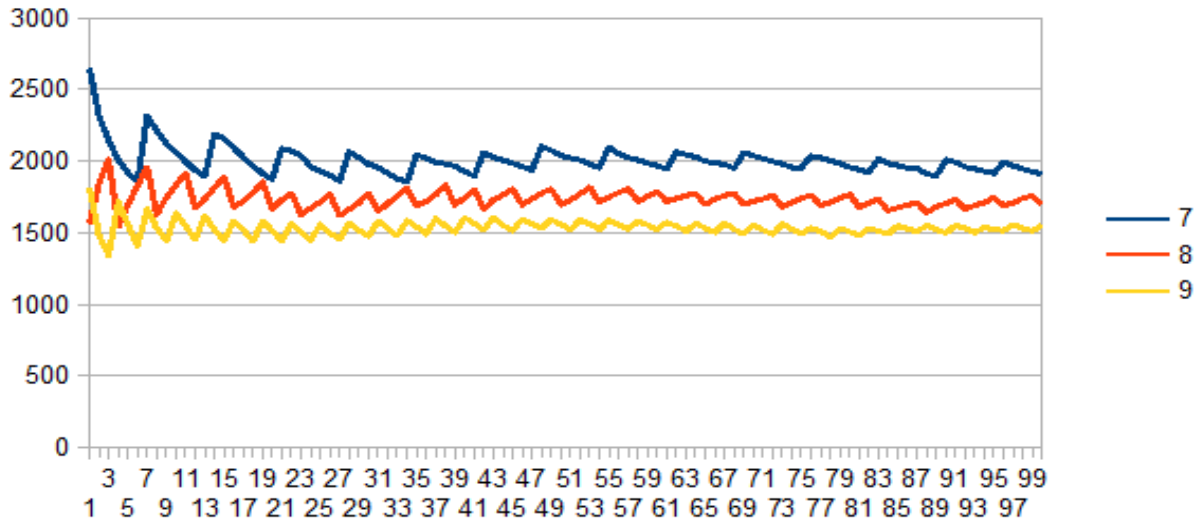


Figure 16: Stationary wave 8bp OFF

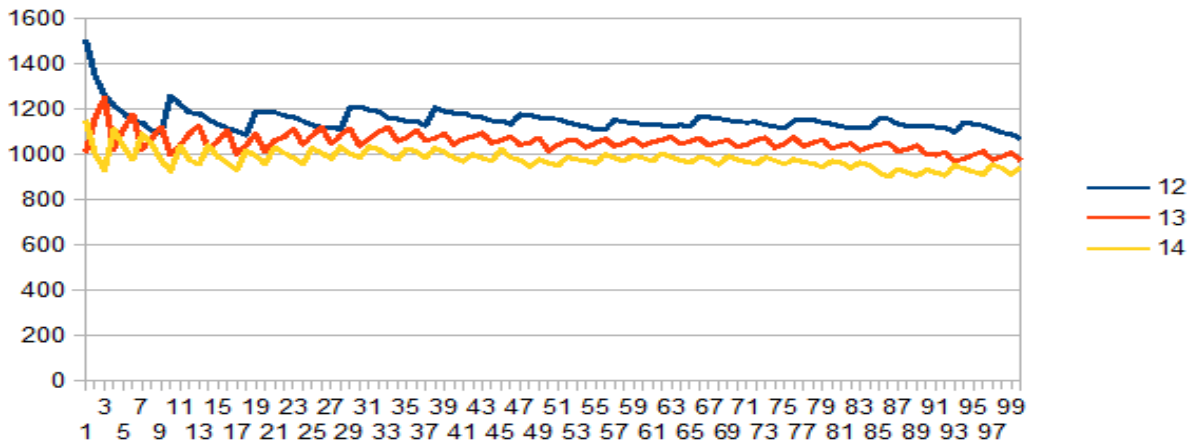


Figure 17: Stationary wave 13bp OFF

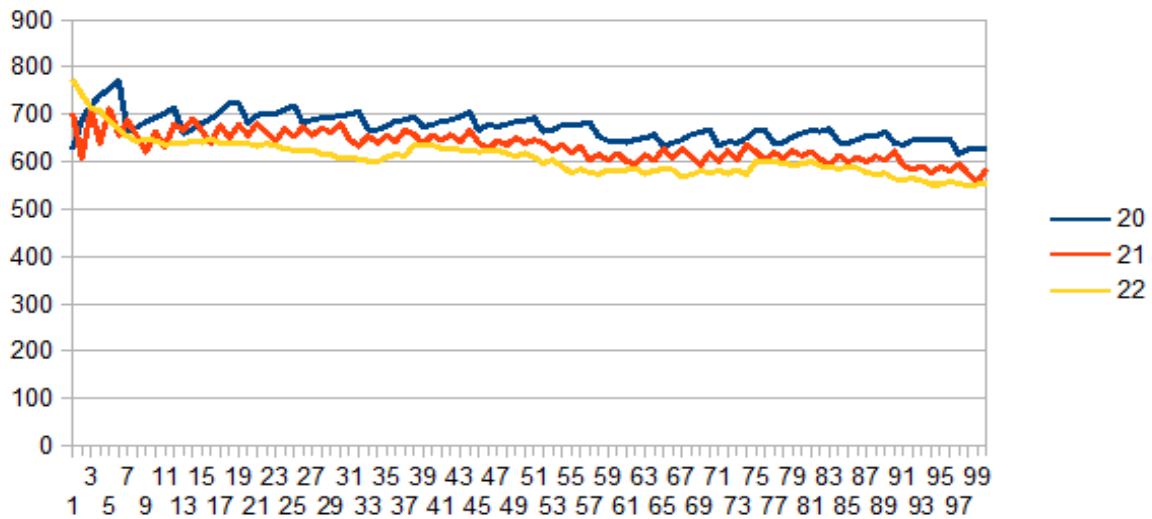


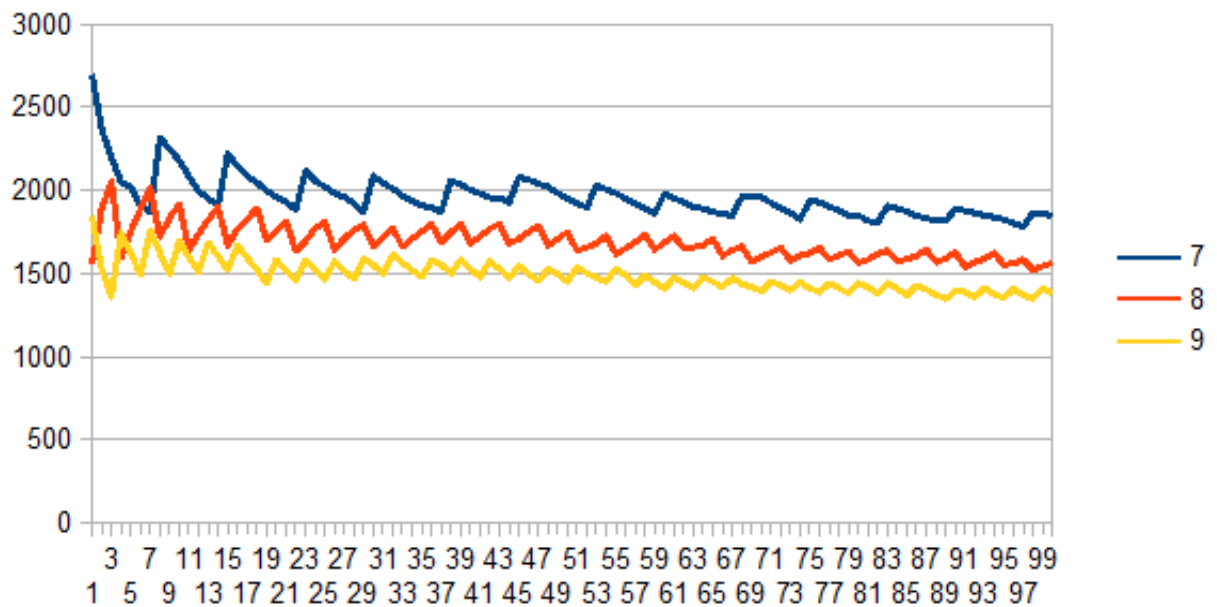
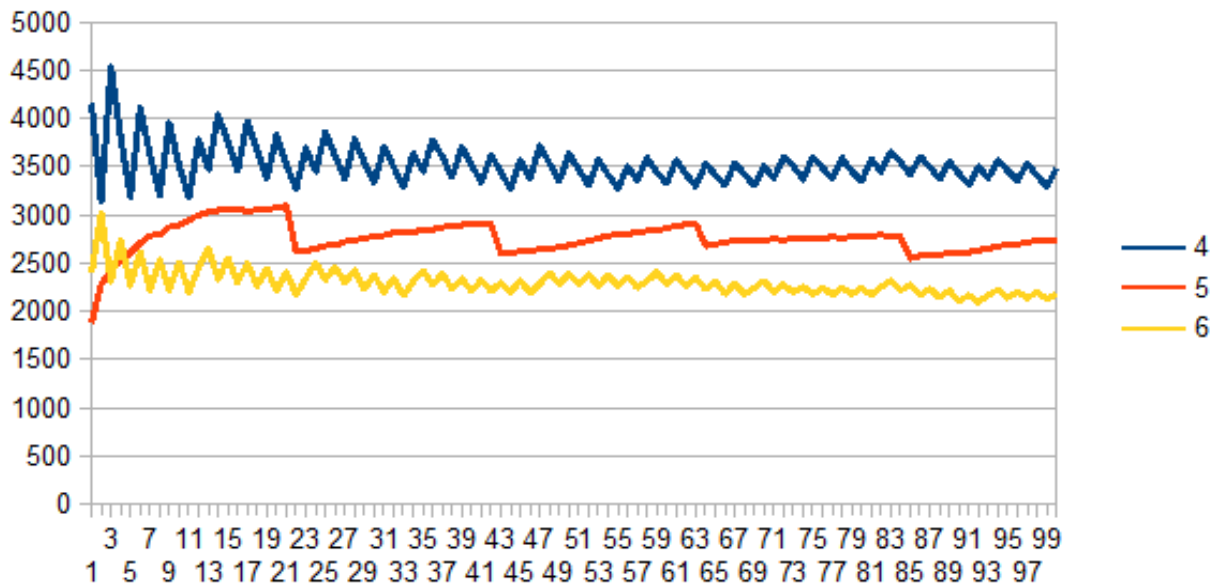
Figure 18: Stationary wave 21bp OFF

In this third SARS2004b genome, we find only the Fibonacci stationary wave 5bp (Figure 15). All other fractal waves 8,13, 21 bp are absent.

- SARS2012

**Bat SARS-Like Coronavirus Isolate Rs4084, Complete Genome - Nucleotide - NCBI. Ky417144.1**

<https://www.ncbi.nlm.nih.gov/nucore/KY417144.1>



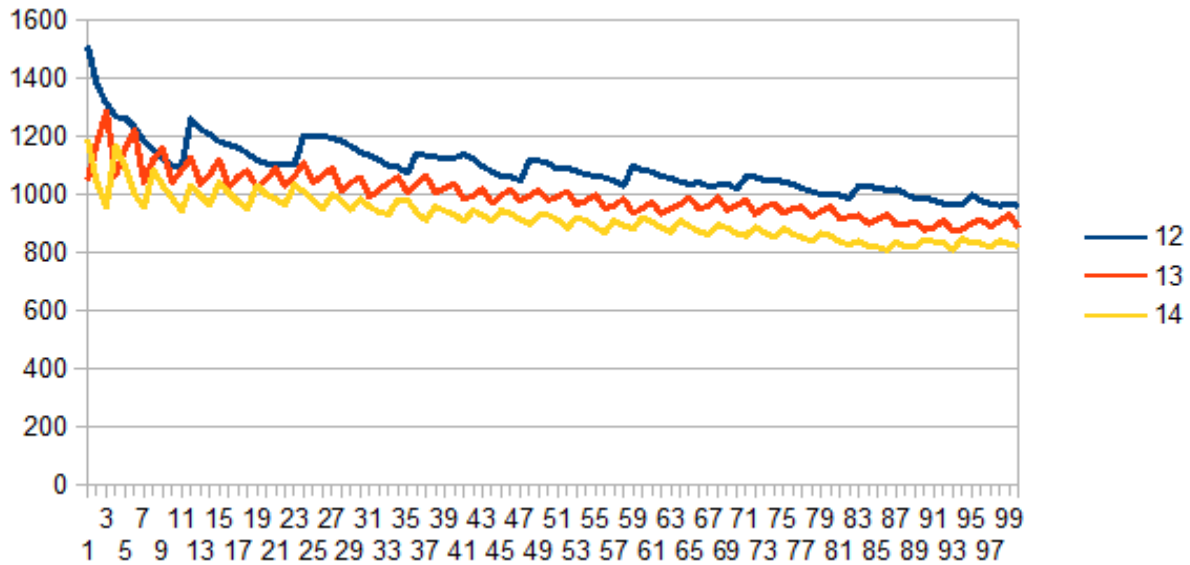


Figure 21: Stationary wave 13bp OFF

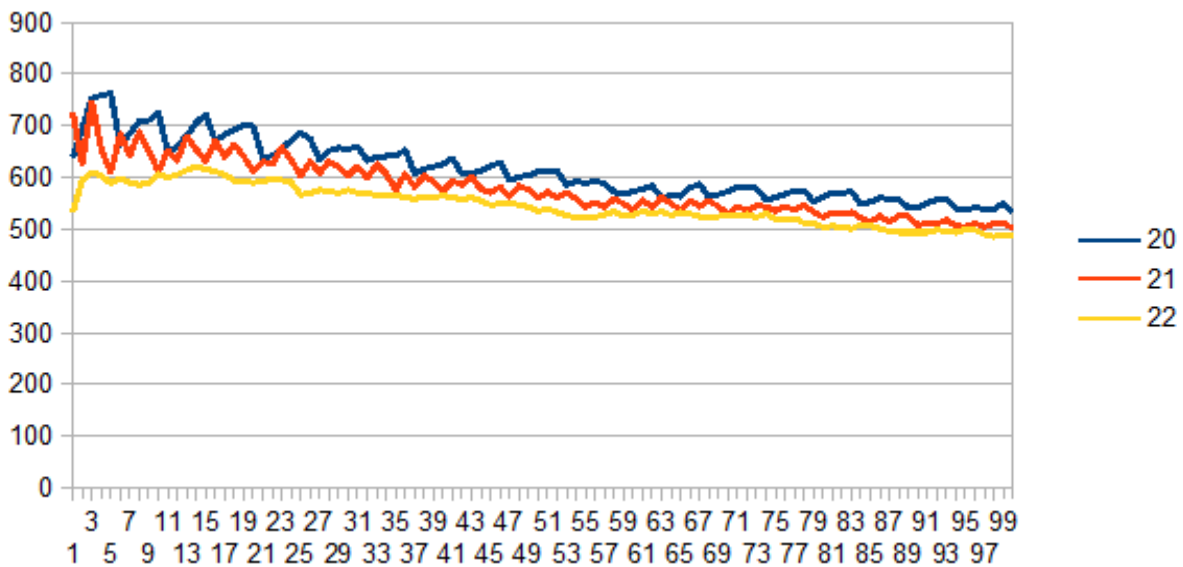


Figure 22: Stationary wave 21bp OFF

In this fourth SARS2012 genome, we find only the Fibonacci stationary wave 5bp (Figure 19). All other fractal waves 8,13, 21 bp are absent.

- SARS2015

**Bat SARS-Like Coronavirus Isolate Bat-SL-Covzxc21, Complete Genome**

<https://www.ncbi.nlm.nih.gov/nuccore/Mg772934>

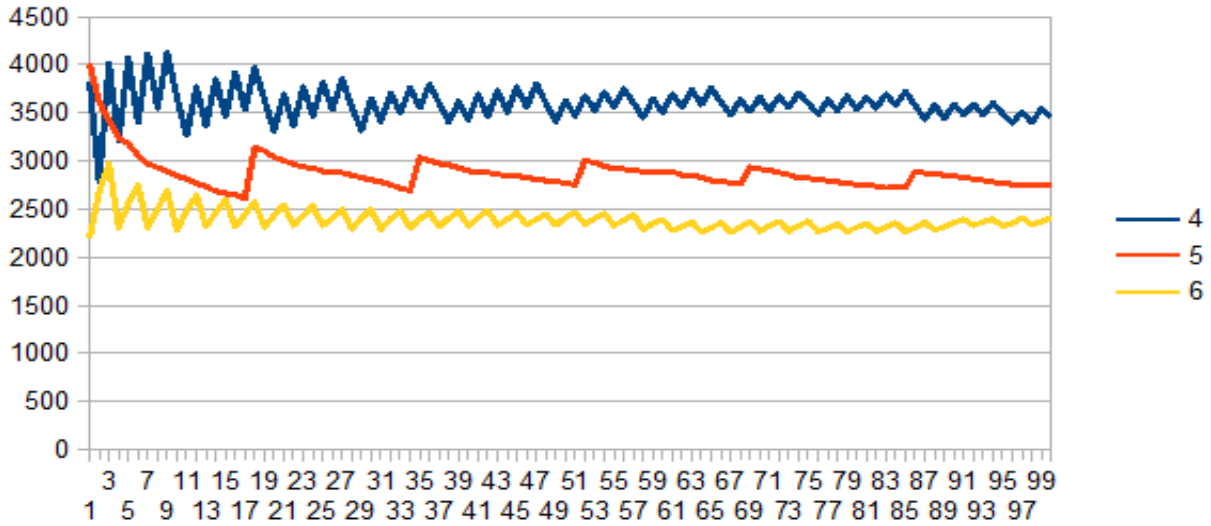


Figure 22: Stationary wave 5bp ON

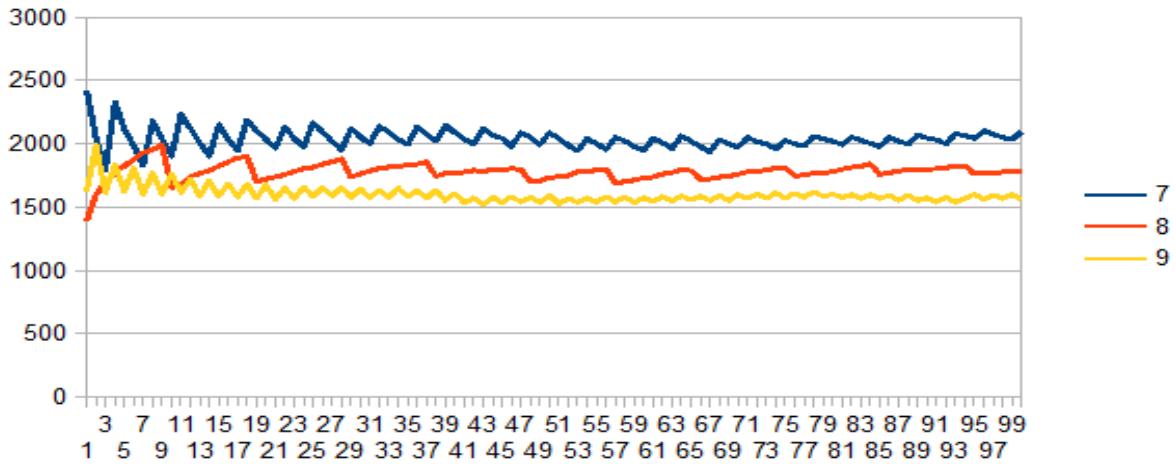


Figure 23: Stationary wave 8bp ON

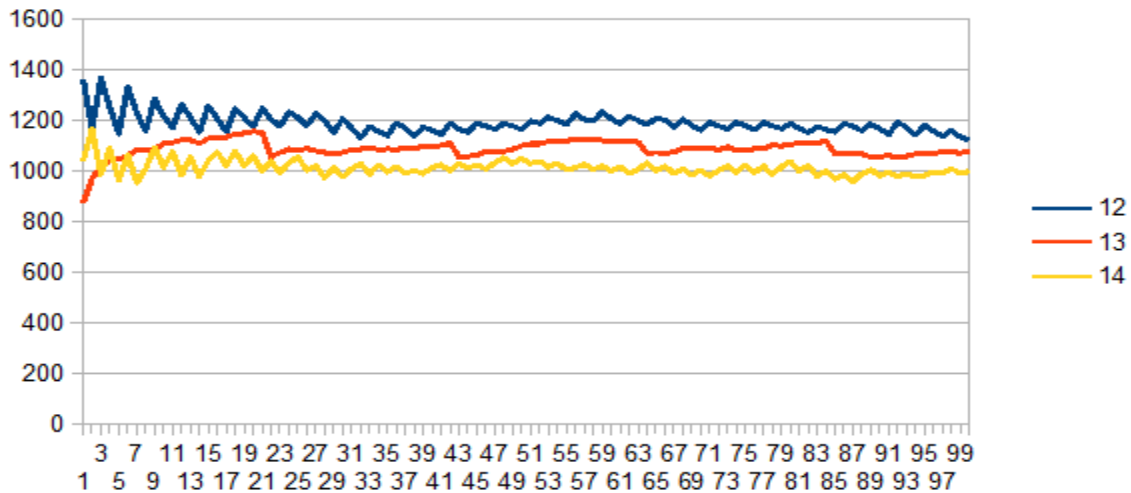


Figure 24: Stationary wave 13bp ON

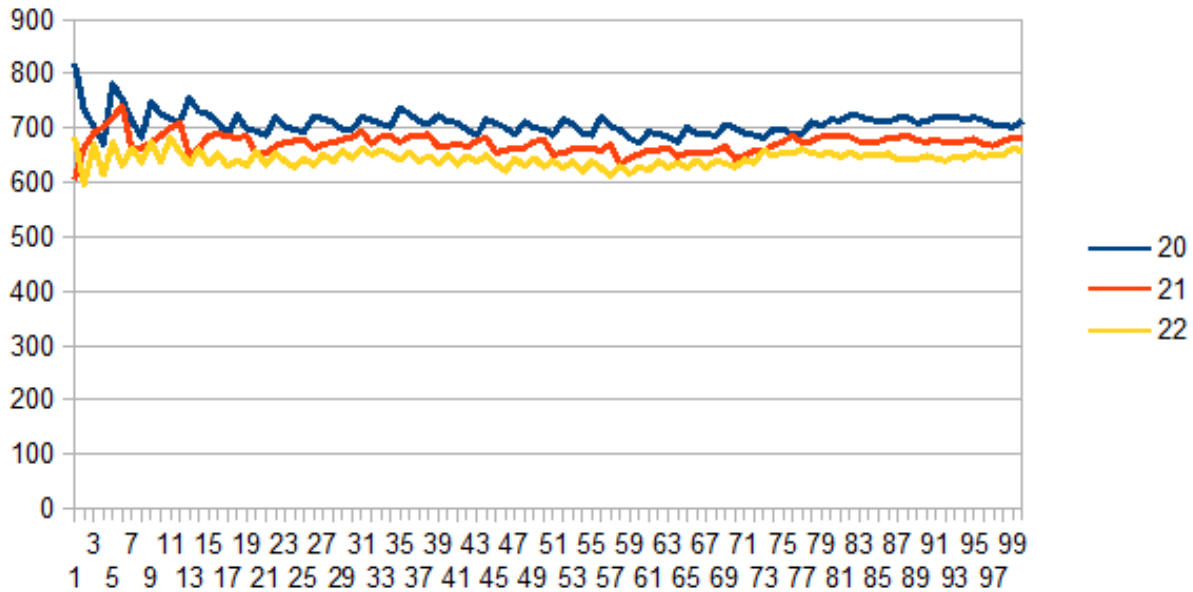


Figure 25: Stationary wave 21bp OFF

In this fifth SARS2015 genome, we find now three Fractal Fibonacci stationary wave 5bp, 8bp, and 13bp (Figures 21, 22, 23). Meanwhile the fourth other fractal wave 21bp remain absent.

- SARS2017

**Bat SARS-Like Coronavirus Isolate Bat-SL-Covzc45, Complete Genome**

<https://www.ncbi.nlm.nih.gov/nuccore/Mg772933>

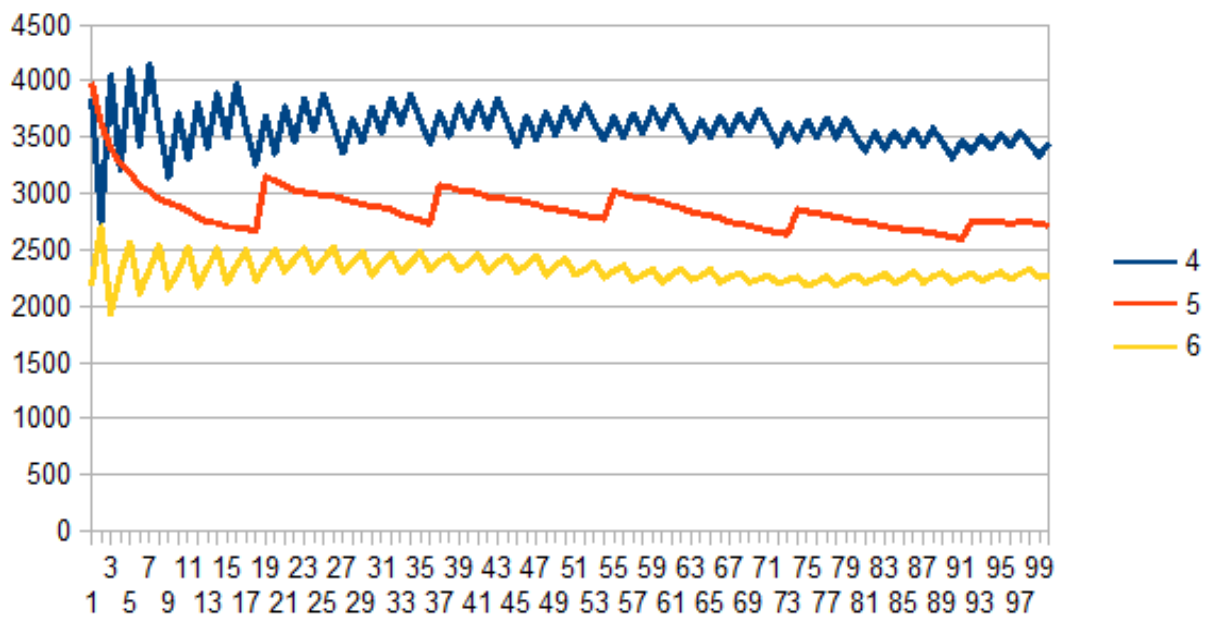


Figure 26: Stationary wave 5bp ON



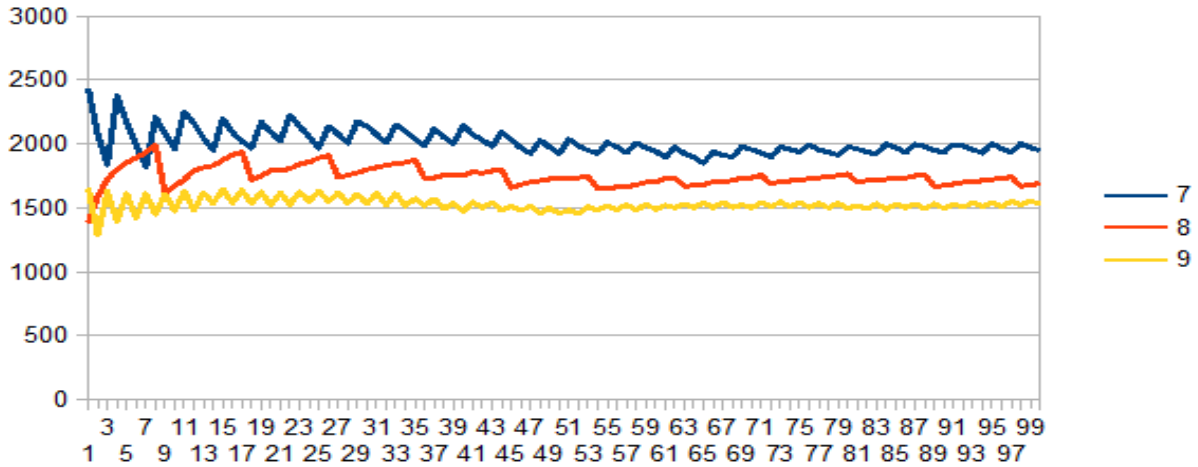


Figure 27: Stationary wave 8bp ON

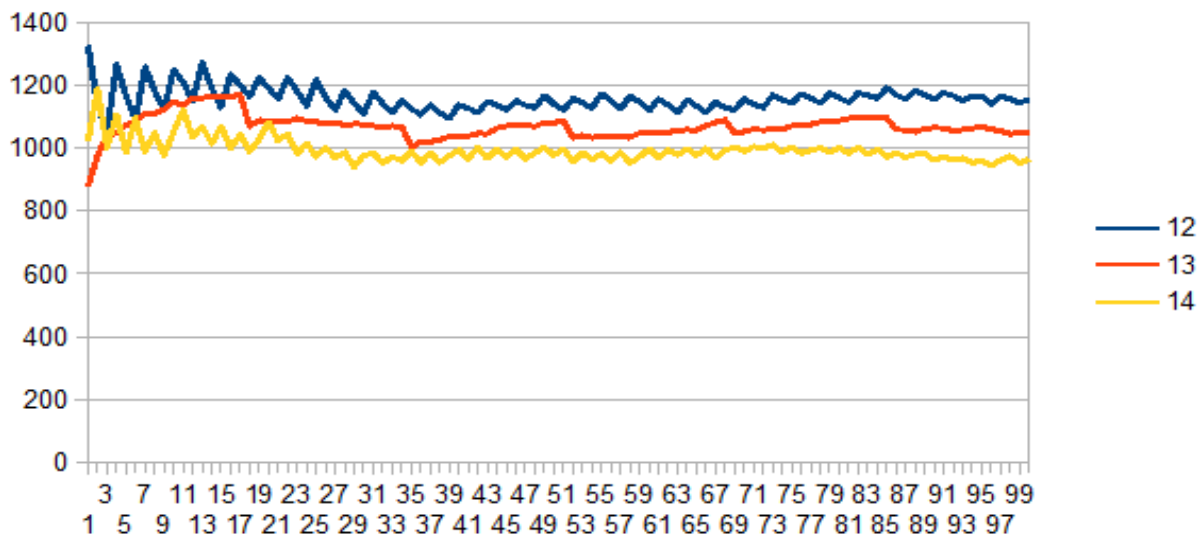


Figure 28: Stationary wave 13bp ON

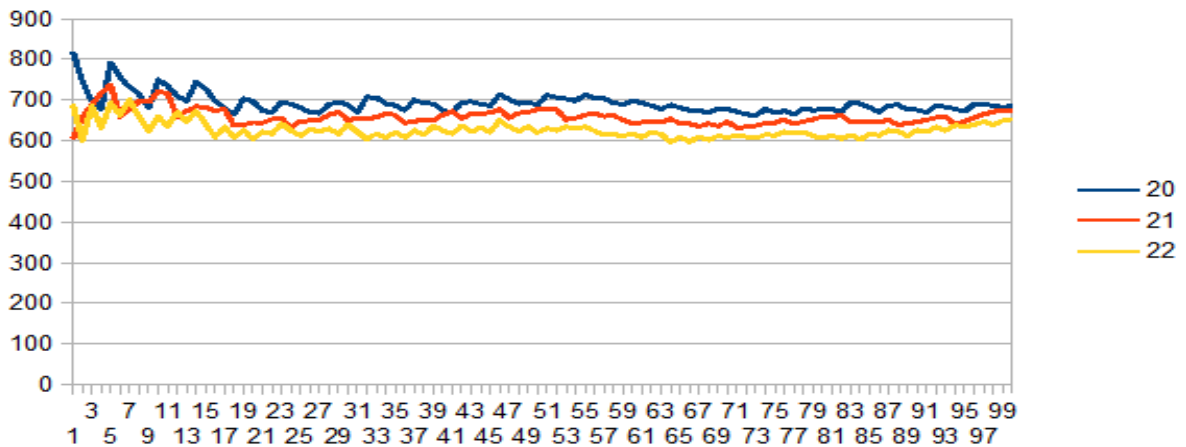


Figure 29: Stationary wave 21bp OFF

In this sixth SARS2017 genome, we find also three Fractal Fibonacci stationary wave 5bp, 8bp, and 13bp (Figures 26, 27, 28). Meanwhile the fourth other fractal wave 21 bp remain absent.

**WUHANOLD (first genome sequenced 12 january 2020) Wuhan seafood market pneumonia virus isolate Wuhan-Hu-1, complete genome MN908947.1**

<https://www.ncbi.nlm.nih.gov/nucore/MN908947.1>

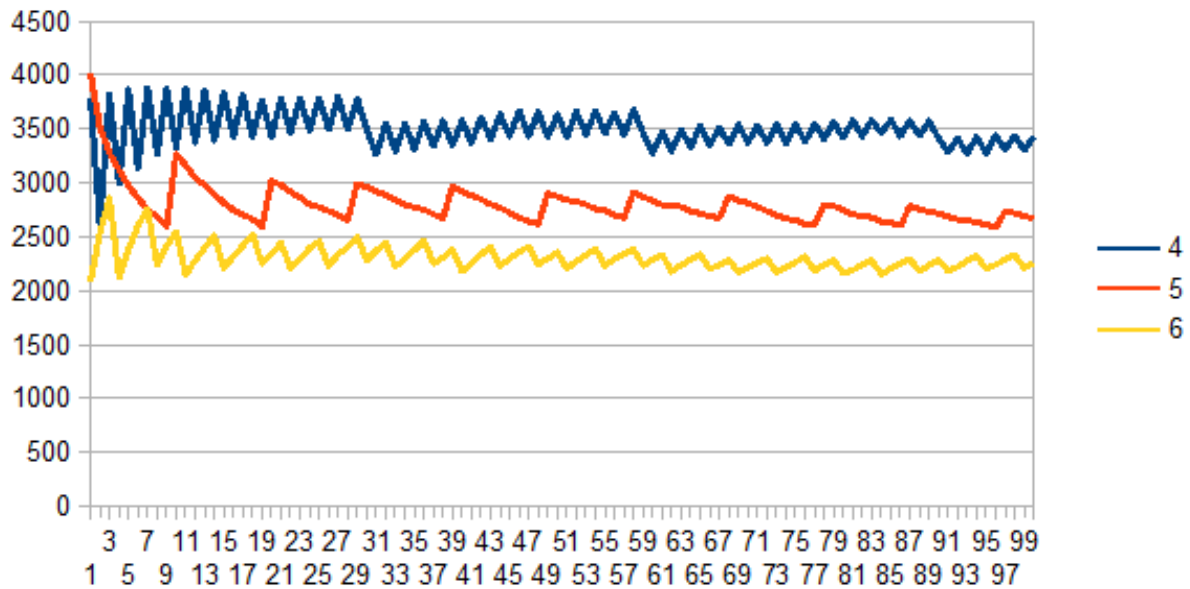


Figure 30: Stationary wave 5bp ON

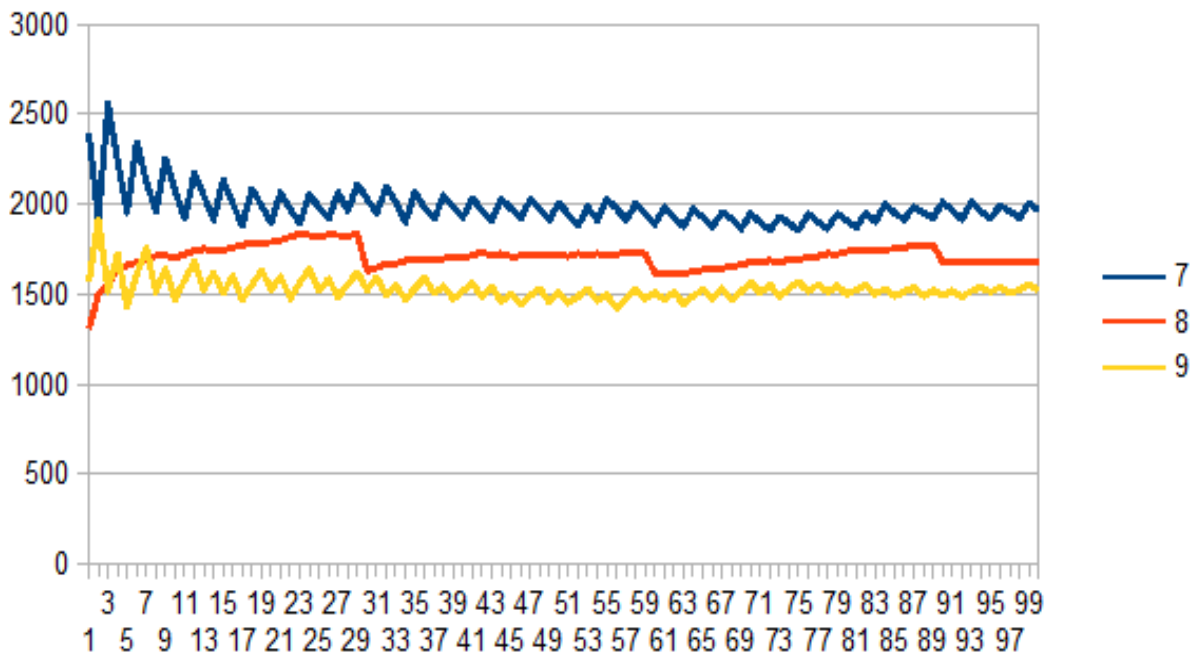


Figure 31: Stationary wave 8bp ON

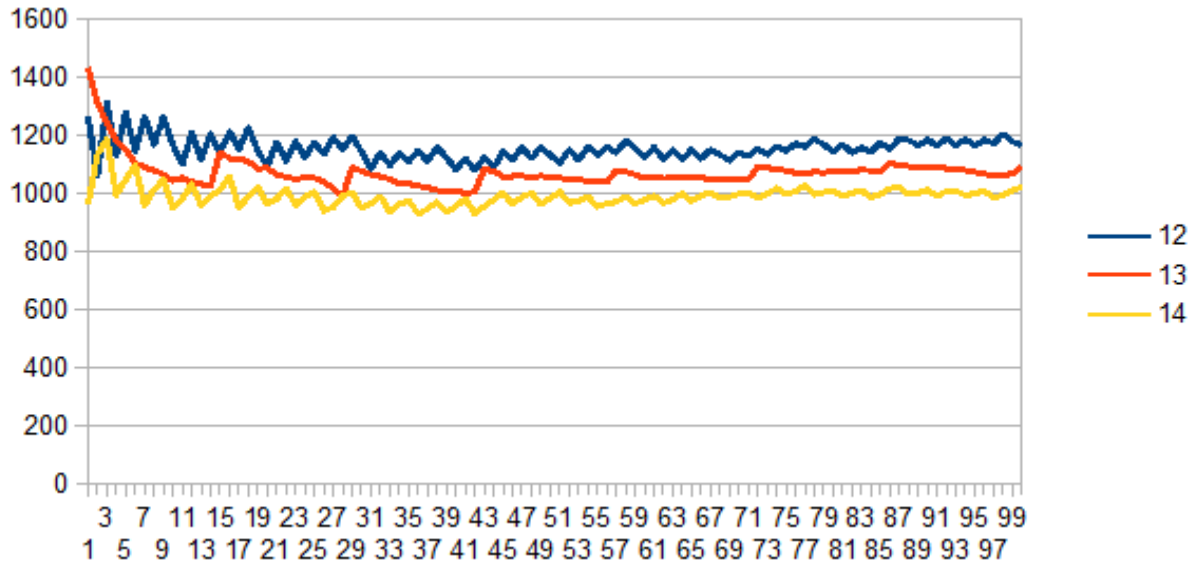


Figure 32: Stationary wave 13bp ON

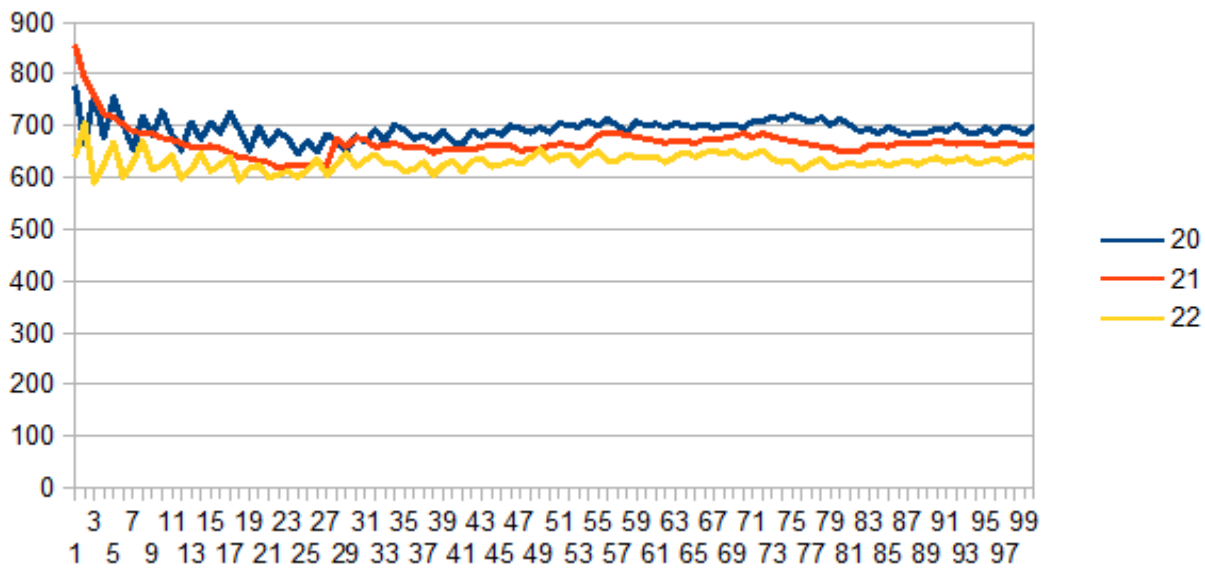


Figure 33: Stationary wave 21bp ON

In this seventh WUHANOLD 12 January 2020 genome, we find also three Fractal Fibonacci stationary wave 5bp, 8bp, and 13bp (Figures 29, 30, 31). Now the fourth other fractal wave 21 bp is also present (Figure 33).

**WUHAN2 (second improved sequenced genome 14 January 2020) Wuhan seafood market pneumonia virus isolate Wuhan-Hu-1, complete genome GenBank: MN908947.2**

<https://ncbiinsights.ncbi.nlm.nih.gov/2020/01/13/novel-coronavirus/>

<https://www.ncbi.nlm.nih.gov/nucore/MN908947.3?report=fasta>

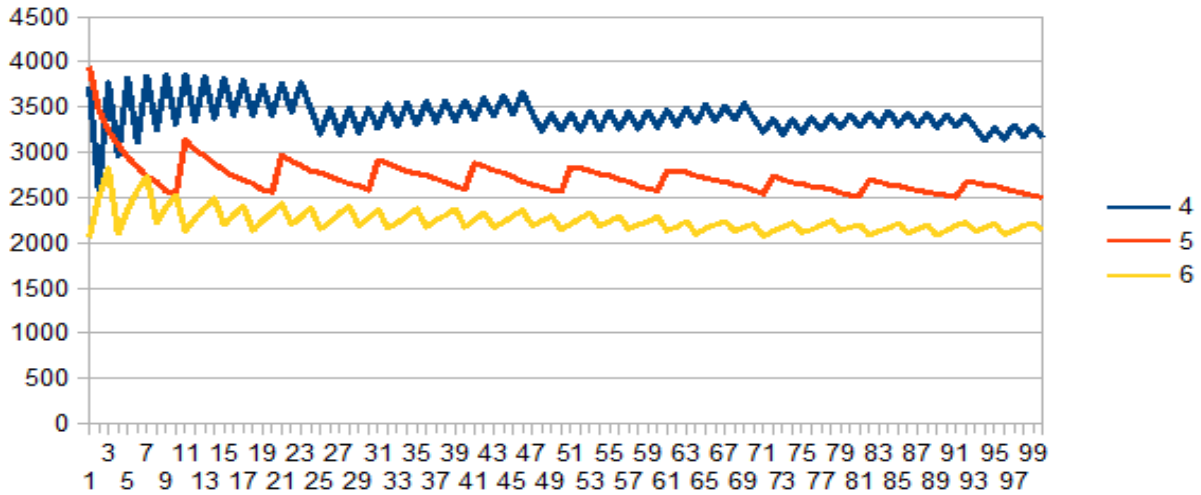


Figure 34: Stationary wave 5bp ON

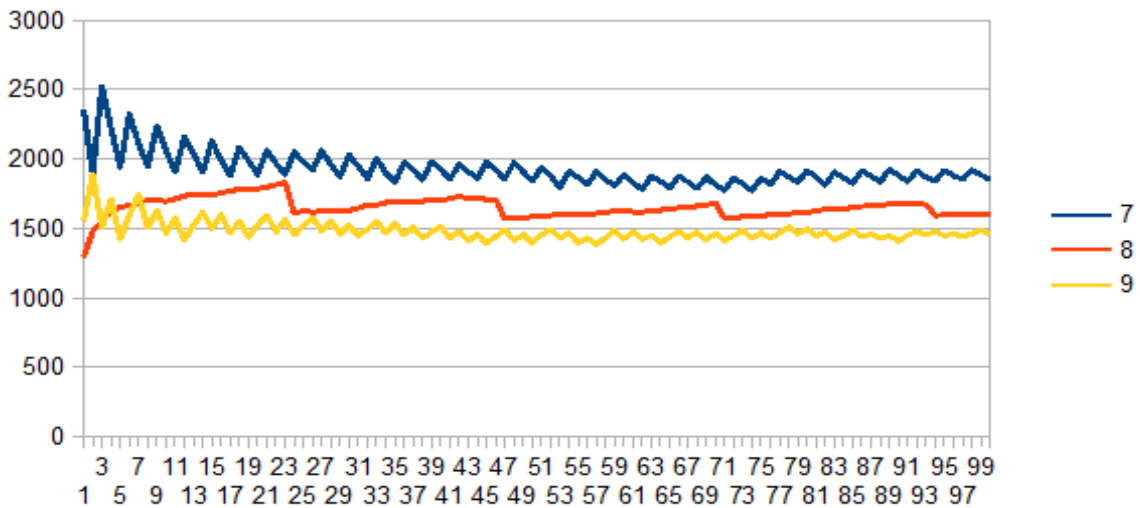


Figure 35: Stationary wave 8bp ON

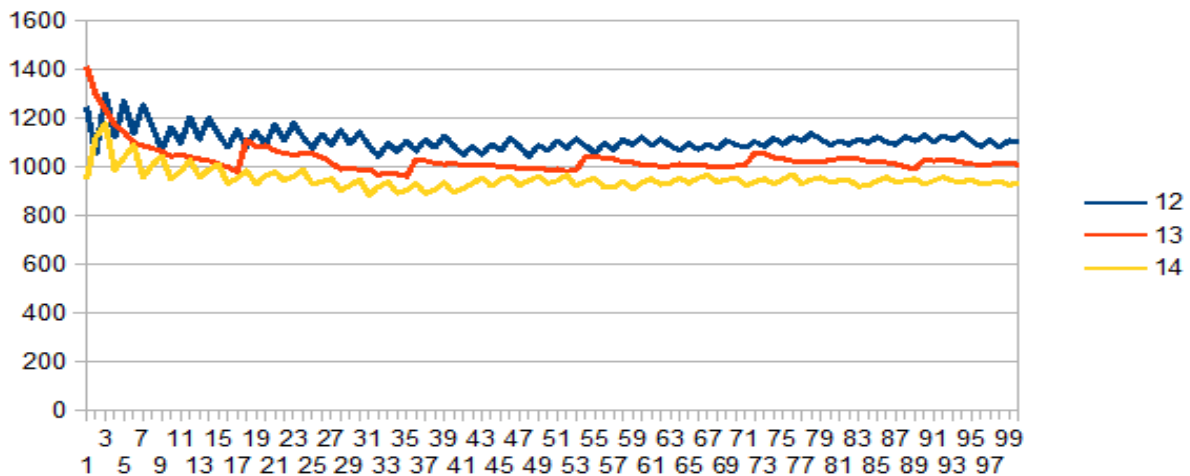


Figure 36: Stationary wave 13bp ON

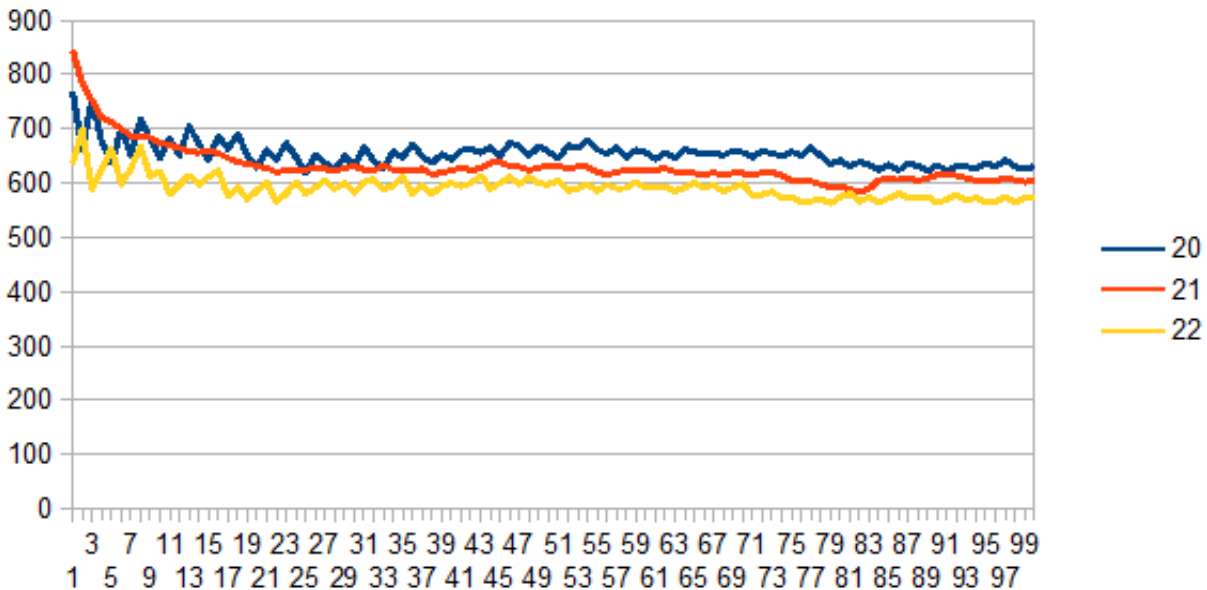


Figure 37: Stationary wave 21bp ON

In this eighth WUHAN2 14 January 2020 genome, we find also three Fractal Fibonacci stationary wave 5bp, 8bp, and 13bp (Figures 33, 34, 35). Now the fourth other fractal wave 21 bp is also present (Figure 36).

**WUHAN (23 January 2020) Wuhan seafood market pneumonia virus isolate Wuhan-Hu-1, complete genome GenBank: MN908947.3**

<https://www.ncbi.nlm.nih.gov/nuccore/MN908947.3>

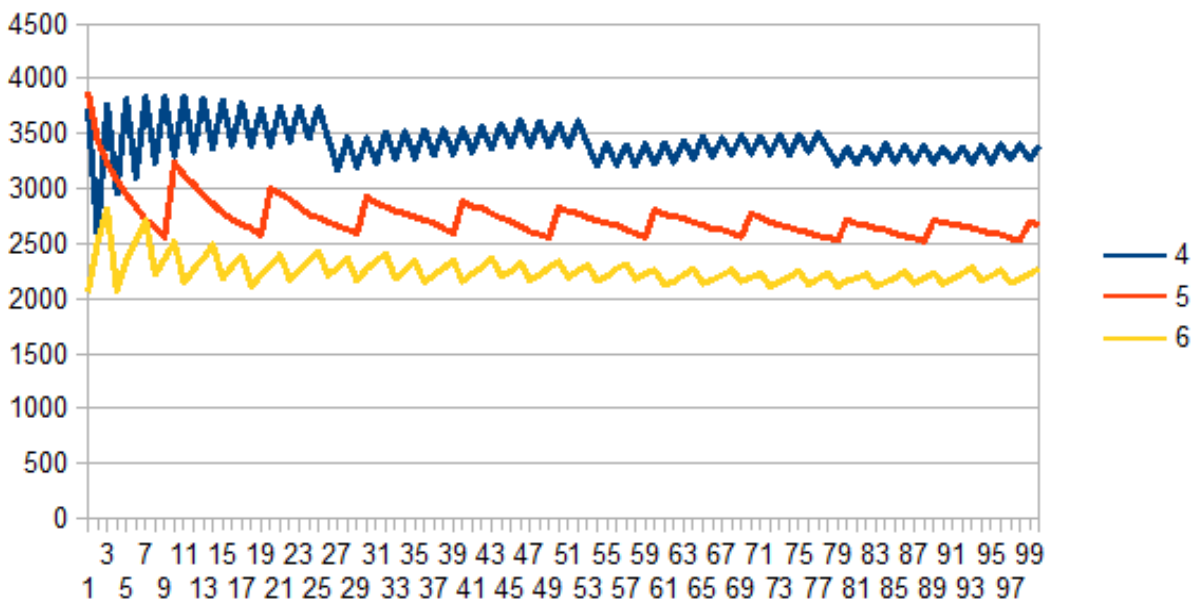


Figure 39: Stationary wave 5bp ON

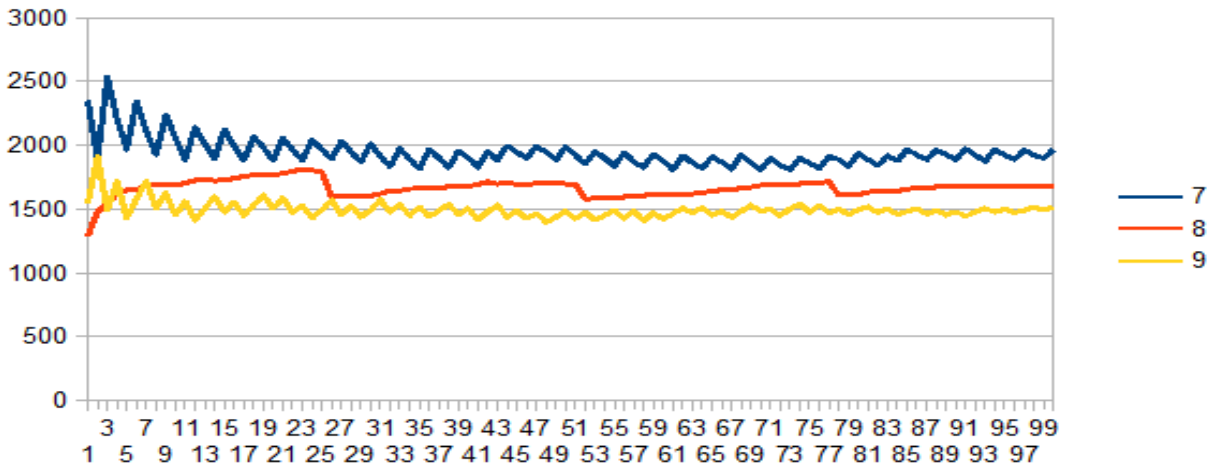


Figure 40: Stationary wave 8bp ON

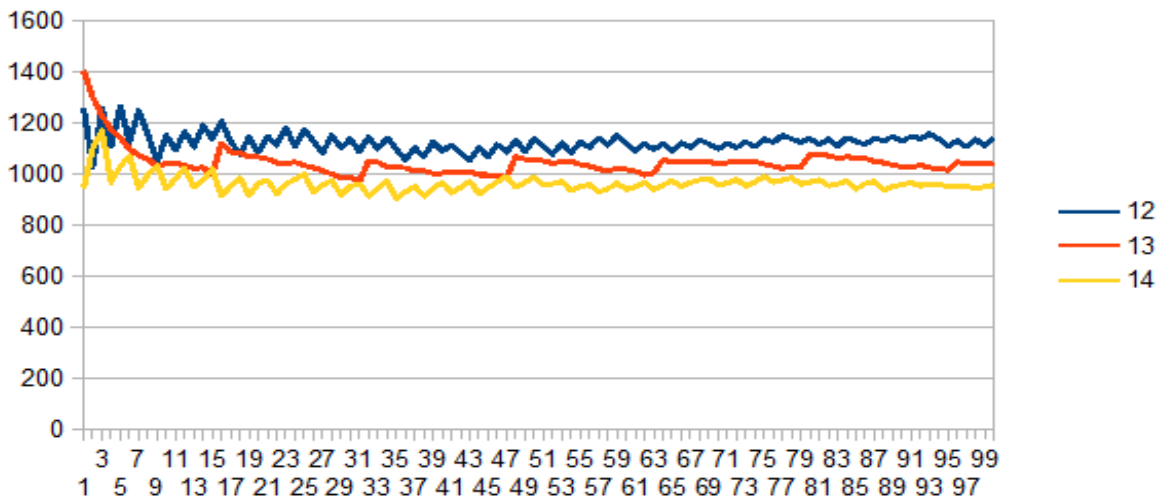


Figure 41: Stationary wave 13bp ON

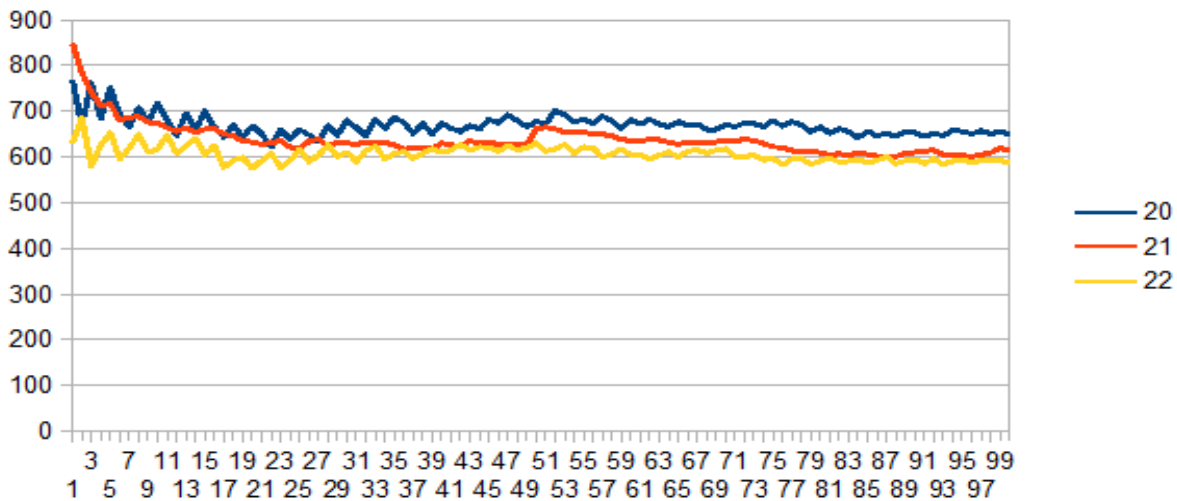


Figure 42: Stationary wave 21bp ON

In this last and ninth reference WUHAN 23 January 2020 genome, we find also three Fractal Fibonacci stationary wave 5bp, 8bp, and 13bp (Figures 39, 40, 41). Now the fourth other fractal wave 21 bp is also present (Figure 42).

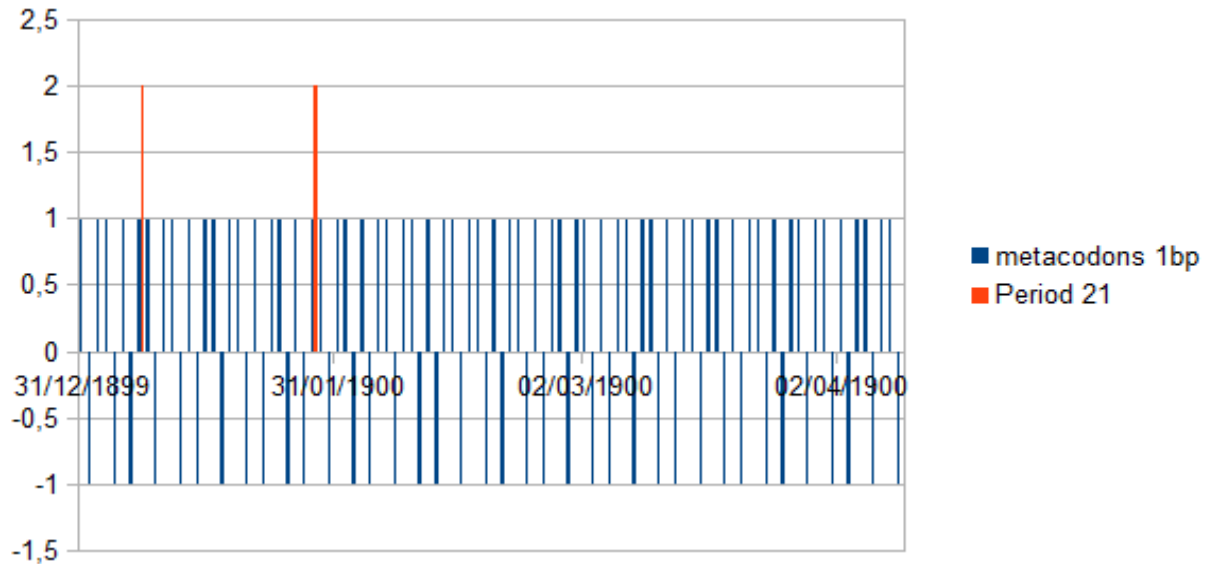


Figure 43: A second method of highlighting the standing wave of period 21bp.

#### 4. Conclusions and Recommendations

Table 3 below clearly demonstrates a kind of evolution of SARS genomes between 2003 and 2020. If it remains difficult to associate this evolution with an increase in pathogenicity for humans, data already published by us [44-46] rather would suggest a greater ADAPTABILITY of these genomes to the human host genome by the same coherence and united via standing waves of Fibonacci [37].

Table3: High level of correlation between the years of emergence of the SARS virus and the presence of Fibonacci fractal standing waves.

Genome reference	NCBI reference	Length bp	Fibonacci numbers FRACTAL embedded stationary waves periods				
			5bp	8bp	13bp	21bp	34bp
SARS2003	AY304488	29731	Yes				
SARS2004	DQ412043	29749	Yes				
SARS2004b	AY395003	29647	Yes				
SARS2012	KY417144.1	29770	Yes				
SARS2015	MG772934	29732	Yes	Yes	Yes		
SARS2017	MG772933	29802	Yes	Yes	Yes		
WUHANOLD	MN908943.1	30473	Yes	Yes	Yes	Yes	
WUHAN2	MN908943.2	29875	Yes	Yes	Yes	Yes	
WUHAN	MN908943.3	29903	Yes	Yes	Yes	Yes	

Hypothetical Evolution genome (see fig 45)	Deleting region in MN908943.3	29153	Yes	Yes	Yes	Yes	Yes
--	-------------------------------	-------	-----	-----	-----	-----	-----

In figure 44 we have superimposed the standing waves of 21bp corresponding to the 3 published versions of the wuhan genome of 2020. There appears a very high sensitivity, which suggests the fact that this new genome is in full phase of evolution in its adaptation to the human host. the 3 figures 33, 37, 42 show this evolution better: WUHANOLD: 3.5 waves, WUHAN2: one and epsilon wave, WUHAN: 2 waves.

**3 genomes releases of NcOv-2019 Wuhan published in 2020**  
 stationary waves 21bp variability periods

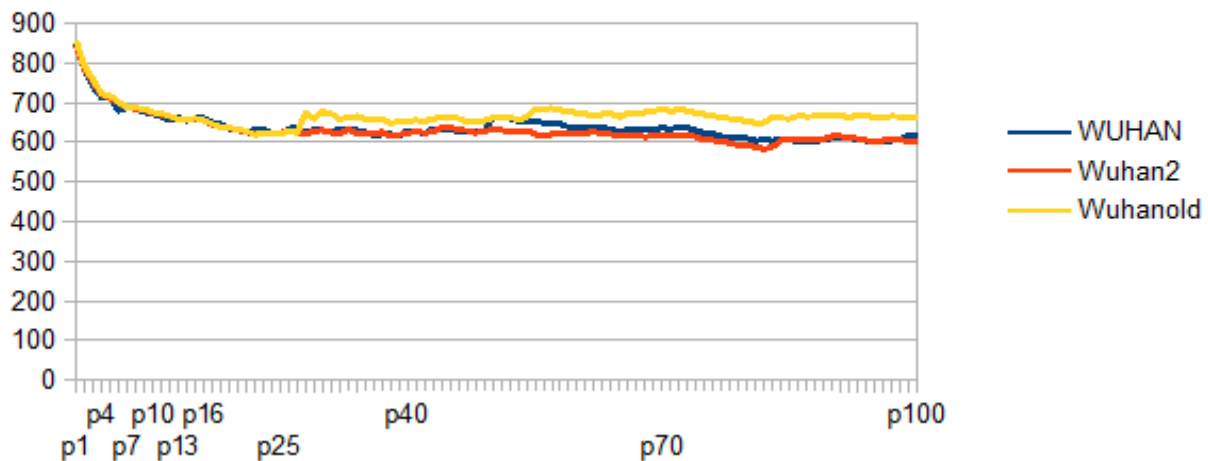


Figure 44: High level of sensitivity for 3 releases of Wuhan CoV-2019 genomes of almost identical lengths (30473, 29875 and 29903bp).

**How Could Tomorrow 2019-Ncov Evolve?**

In [2] and [5], authors show that there is no doubt that 2019-nCoV is a novel unknown sequence. In an unformal draft, Dr Lyons-weiler [4] even suggests that a region between the bases 21600-22350 bp would be completely new. Considering that this region could be "foreign" to the family of coronaviruses we tried to test how its absence could have had an impact on the waves that we reveal here. We then construct a hypothetical genome which would no longer have this insertion between the bases 21600-22350 bp. It then appears (Figure 45), for the first time a Fibonacci wave of 34bp. This genome would therefore be structured by a fractal nesting of 5 Fibonacci waves, 5 8 13 21 and now 34bp. Curiously, we have only encountered such a level of organization in the entire human chromosome4 (figure 46). A new question: "Can there be some kind of affinity between the waves of a retrovirus and the human host genome in which this retrovirus could integrate?"



### 2019-nCoV Coronavirus standing wave 34bp deleting 21600-22350 bp

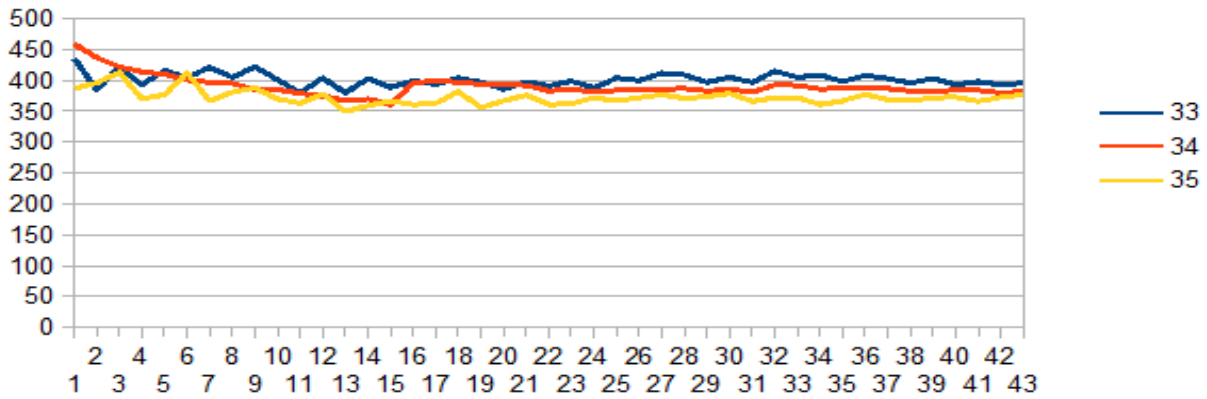


Figure 45: Standing wave 34bp structuring modified 2019-NCoV genome.

The chromosome 4 HG38 of reference:

### chr4 HG38 (2013) resonance 34

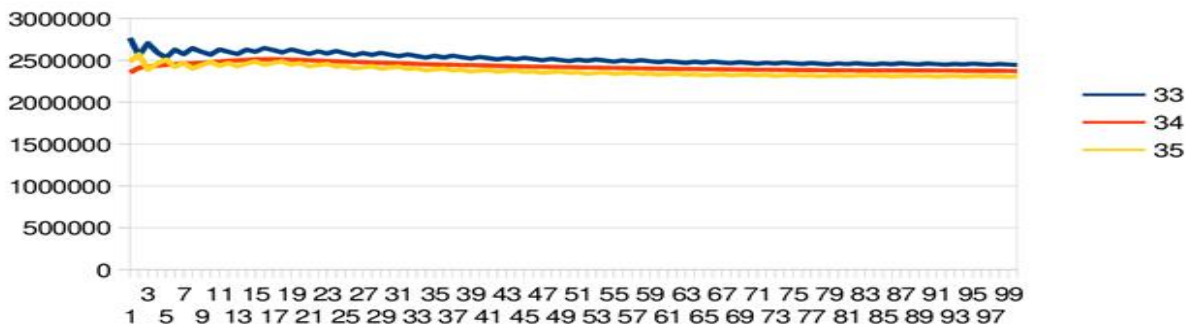


Figure 18 : The main resonance of 34bp characterizing the HG38 reference Chromosome 4.

### Sapiens HG38 Reference Chromosome4 Period 34



Figure 46: Standing wave 34bp overlapping the whole human chromosome4 (from [43]).

### On A Possible "Harmonic Standing Wave Agreement" Between the Human Host Chromosome and The Coronavirus Retrovirus:

This last observation opens here an interesting theoretical track on 2 possible strategies of integration of retroviruses in eucharyotic chromosomes:

The chromosome 4 HG38 of reference:

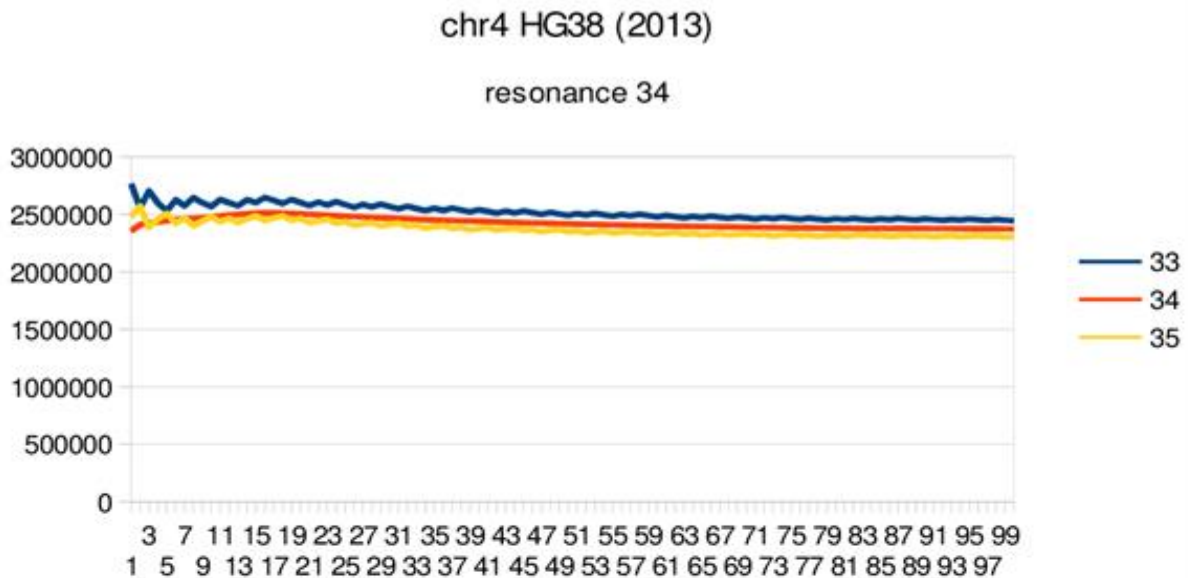
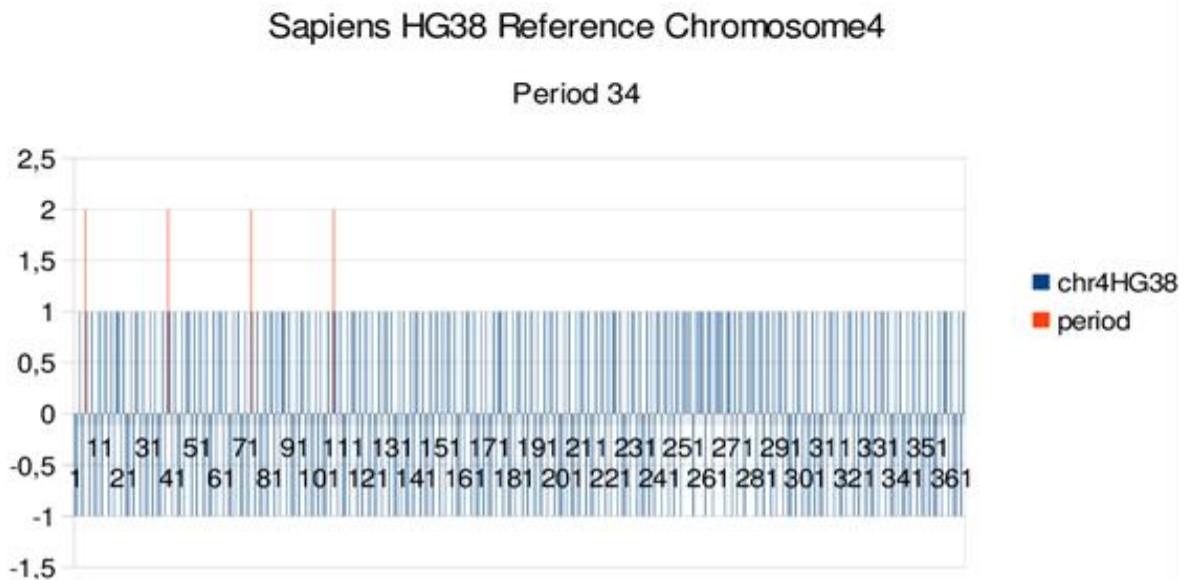


Figure 18 : The main resonance of 34bp characterizing the HG38 reference Chromosome 4.



1 / symbiosis strategy by complementarity: example of HIV more frequently integrating chromosomes 20 16 17 22 19. These chromosomes have a poor HGO (Human Genome Optimum) ratio, it will be slightly improved by the integration of the virus.

Indeed, in [8, 46], we demonstrated why the permeability to the integration of retrovirus in each of our chromosomes is correlated with this classification HGO (HUMAN GENOME OPTIMUM) of this article <http://www.imedpub.com/abstract/towards-a-universal-law-controlling-all-human-cancer-chromosome-loh-deletions-perspectives-in-prostate-and-breast-cancers-screening-20846.html>

For example, chromosomes 20. 16. 17. 22 19 are very permeable: this table shows that they are located at the bottom of this table of classification of HGOs of 24 human chromosomes.

Table 3: HGO human chromosomes classification.

Table 1: CG values, TA values and CG/TA ratio for each Sapiens HG38 individual chromosome.

Chromosome	C + G	T + A	(C + G)/(T + A)
Chromosomes UPSTREAM HGO point = (3-Phi)+2 = 0.6909830056			
4	72568001	1.17E+08	0.619262
13	37772797	60210328	0.627347
5	71611274	1.1E+08	0.653065
X	61221521	93671508	0.653577
6	67360020	1.03E+08	0.655773
3	78577742	1.2E+08	0.657431
18	31856106	48233499	0.660456
Y	10572683	15842360	0.667368
8	58133960	86634176	0.671028
2	96769083	1.44E+08	0.67304
7	64696843	94273288	0.686269
12	54275482	78862334	0.688231
14	36982791	53585358	0.690166
Chromosomes DOWNSTREAM HGO point = (3-Phi)+2 = 0.6909830056			
21	16411625	23676994	0.693146
9	50270473	71520077	0.702886
11	55885058	78648684	0.710566
10	55359481	77903481	0.710616
1	96166571	1.34E+08	0.715981
15	35578844	49062481	0.725174
20	28010605	35933652	0.779509
16	36472718	45333225	0.804547
17	37575444	45344760	0.828661
22	18406838	20752939	0.886951
19	28015712	30425046	0.920811

<https://juniperpublishers.com/ctoij/images/CTOIJ.MS.ID.555756.T001.png>

We also see in figure 47 from Figure 4 (Wang 2007), that these chromosomes are very permeable to the HIV retrovirus (Dark green bands in the following image from [3]).

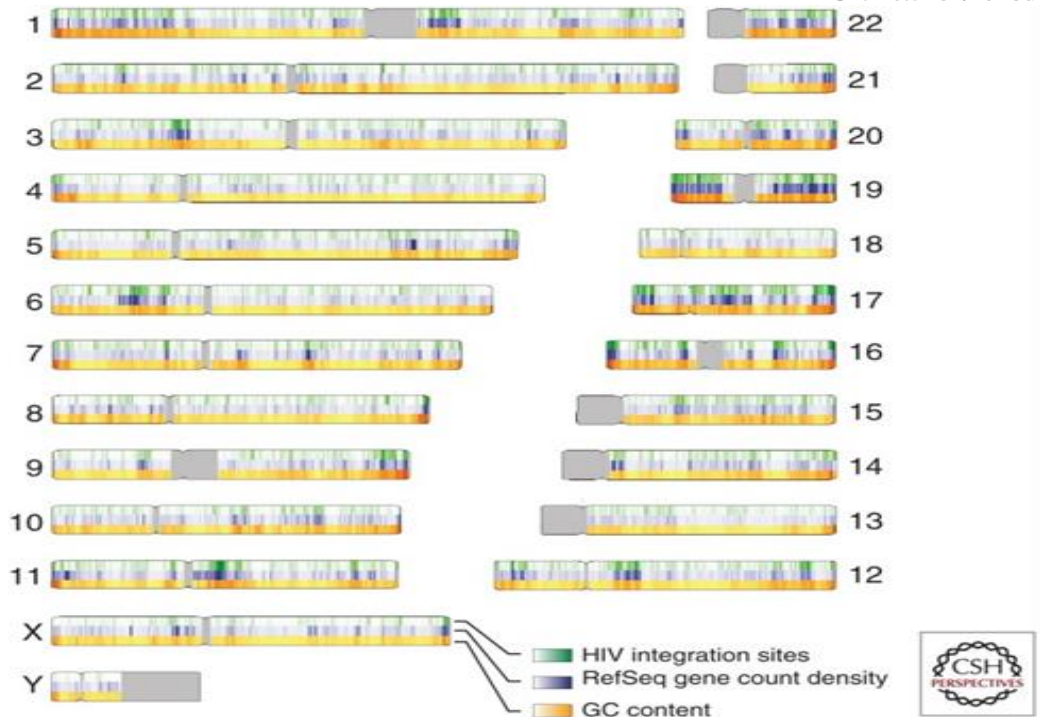


Figure 47: from [3] Green regions are retroviruses integration region,

<https://www.ncbi.nlm.nih.gov/pmc/articles/PMC3385939/figure/A006890F4/>

2 / symbiosis strategy by wave agreement, resemblance: this could be the case of the coronavirus which would integrate by a kind of harmonic agreement to chromosomes 4 or 13, located at the beginning of the HGO table, which have like it Fibonacci waves . This would hardly affect the resulting HGO ratio, or even strengthen it: This is surely the case with RETROTRANSPOSONS.

Finally, we have recognized here 2 laws of symbiosis of nature: agreement because very DIFFERENT, and agreement because very SIMILAR.  
 Strategies that we find up to the affinities between humans ...!!!

If the genome of the 2019-nCoV retrovirus has adopted this second strategy of integration into the human genome, this could explain the fact observed at the beginning of 2020: a moderately pathogenic virus but which spreads very quickly ... Because its retrovirus would be judiciously adapted to its (supposed) host chromosome4 of the human genome?

**On A Possible Origin Of 2019-Ncov Genome:**

It is very likely that there was HUMAN INTERVENTION in this LYONS's region [4] of wuhan genome:

Analysis of this region in all coronaviruses shows a 100% jump in homology for the Wuhan genomes and 70 to 80% for the closest SARS. Although there is already a trace of ENV HIV1 in the genome that we have referenced here SARS2003.

While there is NO TRACE of HIV1 ENV in the region (Lyons-weiler 20020) in all other SARS Coronavirus genomes, Indeed, we find on a mini region (243-86bp = ~ 160bp) 3 partial regions of ENV HIV1 which are all 3 "DIFFERENT" and from 3 different HIV1 strains:

- COVID-19 (86-113) ==> in ENV HIV1: 1201 <==> 1228.
- COVID-19 (213-244) ==> in ENV HIV1: 424 <==> 394.
- COVID-19 (243-281) ==> in ENV HIV1: 1064 <==> 1029.

in ENV HIV1 Here, these 3 regions, while hyper compressed in Lyons wuhan (<200bp), they are more widely spaced in hiv1 env (394 <==> 1228), ie> 830bp.??? HOW TO EXPLAIN THIS CONCENTRATION OF 3 SEQUENCES ENV HIV1???? IF NOT BY HUMAN ACTION????

Then, in a second time, we discovered 3 other regions from HIV2 and SIV...

See details and Blast proves here:

Location of the 300 first bp in [4] from wuhan COVID-19 reference genome (starting from bp 21672).

**Wuhan seafood market pneumonia virus genome assembly, chromosome: whole\_genome  
 Sequence ID: LR757998.1 Length: 29866Number of Matches: 1**

Range 1: 21672 to 21971GenBankGraphicsNext Match Previous Match

Alignment statistics for match #1

Score	Expect	Identities	Gaps	Strand
555 bits(300)	2e-158	300/300(100%)	0/300(0%)	Plus/Plus

Query 1

```

CTCAGTTTTACATTCAACTCAGGACTTGTTCTTACCTTTCTTTTCCAATGTTACTTGGTT 60
      |||
Sbjct 21672 CTCAGTTTTACATTCAACTCAGGACTTGTTCTTACCTTTCTTTTCCAATGTTACTTGGTT 21731

Query 61  CCATGCTATACATGTCTCTGGGACCAATGGTACTAAGAGGTTTGATAACCCTGTCCTACC 120
      |||
Sbjct 21732 CCATGCTATACATGTCTCTGGGACCAATGGTACTAAGAGGTTTGATAACCCTGTCCTACC 21791

Query 121 ATTTAATGATGGTGTTTATTTTGCTTCCACTGAGAAGTCTAACATAATAAGAGGCTGGAT 180
      |||
Sbjct 21792 ATTTAATGATGGTGTTTATTTTGCTTCCACTGAGAAGTCTAACATAATAAGAGGCTGGAT 21851

Query 181 TTTTGGTACTACTTTAGATTCGAAGACCCAGTCCCTACTTATTGTTAATAACGCTACTAA 240
      |||
Sbjct 21852 TTTTGGTACTACTTTAGATTCGAAGACCCAGTCCCTACTTATTGTTAATAACGCTACTAA 21911

Query 241 TGTTGTTATTAAAGTCTGTGAATTTCAATTTTGTAAATGATCCATTTTGGGTGTTTATTA 300
      |||
Sbjct 21912 TGTTGTTATTAAAGTCTGTGAATTTCAATTTTGTAAATGATCCATTTTGGGTGTTTATTA 21971
    
```

**Details:**

**Region HIV1a 86-113:**

*HIV-1 isolate 19663.24H9 from Netherlands envelope glycoprotein (env) gene, complete cds*

**Sequence ID: GU455503.1Length: 2598Number of Matches: 1**

Range 1: 1201 to 1228 GenBankGraphics Next Match Previous Match

Alignment statistics for match #1

Score	Expect	Identities	Gaps	Strand
38.3 bits(41)	1.9	25/28(89%)	0/28(0%)	Plus/Plus

```

Query 86  AATGGTACTAAGAGGTTTGATAACCCTG 113
          |||||
Sbjct 1201 AATGGTACTAAGAGGGTAGATAACACTG 1228
    
```

**Region HIV1b 213-244:**

*HIV-1 isolate 4045\_Plasma\_Visit1\_amplicon5a from Malawi envelope glycoprotein (env) gene, complete cds*

**Sequence ID: KC187063.1Length: 2547Number of Matches: 1**

Range 1: 394 to 424GenBankGraphics Next Match Previous Match

Alignment statistics for match #1

Score	Expect	Identities	Gaps	Strand
37.4 bits(40)	7.1	28/32(88%)	1/32(3%)	Plus/Minus

```

Query 213 CCCTACTTATTGTTAATAACGCTACTAATGTT 244
          |||||
Sbjct 424 CCCTACTAAT-GTTACTAACCCTACTAATGTT 394
    
```

**Region HIV1c 243-281:**

*HIV-1 isolate 07.RU.SP-R497.VI.G3 from Russia envelope glycoprotein (env) gene, complete cds*

**Sequence ID: GU481453.1Length: 2580Number of Matches: 1**

Range 1: 1029 to 1064GenBankGraphicsNext Match Previous Match

Alignment statistics for match #1

Score	Expect	Identities	Gaps	Strand
38.3 bits(41)	7.1	32/39(82%)	3/39(7%)	Plus/Minus

```

Query 243 TTGTTATTAAAGTCTGTGAATTTCAATTTTGTAAATGATC 281
          ||| | | | | | | | | | | | | | | | | | | | | |
Sbjct 1064 TTGTTATTAAAGTATTT--TTTCAATTTGTACTTATC 1029
    
```

**Region HIV2a 24-43:**

*HIV-2 isolate 106CP\_RT from Cote d'Ivoire reverse transcriptase gene, partial cds*

**Sequence ID: KJ131112.1Length: 924Number of Matches: 1**

Range 1: 66 to 85GenBankGraphicsNext MatchPrevious Match

Alignment statistics for match #1

Score	Expect	Identities	Gaps	Strand
32.8 bits(35)	0.38	19/20(95%)	0/20(0%)	Plus/Minus

```

Query 24 ACTTGTTCTTACCTTTCTTT 43
          ||| | | | | | | | | | | |
Sbjct 85 ACTTGTTCTTATCTTTCTTT 66
    
```

**Region HIV2b 133-158:**

*Human immunodeficiency virus type 2 complete genome from strain HIV-2UC1*

**Sequence ID: L07625.1Length: 10271Number of Matches: 1**

Range 1: 6701 to 6726GenBankGraphics Next Match Previous Match





## **To Conclude:**

Figure 48: Double level of correlation between the years of emergence of the SARS virus, the presence of standing Fibonacci fractal waves, and the Fractal number of nested and embedded Fibonacci layers.

One track that will need to be deepened is that of integrating the SARS coronavirus genome into human chromosomes [9, 57].

Indeed, if we show that each of the 24 human chromosomes is characterized by one or more specific standing waves, some of our chromosomes (chromosomes 4 and 13 for example) have standing waves which are Fibonacci numbers [7]. There could then be a kind of harmonic agreement between the genome of the SARS virus and its human host chromosome, both of which have standing Fibonacci waves, which will facilitate and strengthen the integration and persistence of these viruses in humans.

Finally, That is a formal proof of an evolution increasing global structure of SARS whole genomes, probably linked with genome integrity and coherence and, human génome adaptation [8, 38] , perhaps pathogenicity.

## **Evidence of A Kind of "Intelligent Will" ...**

Please, now, see figures 49 then 50.

Both figures proves evidence that the 6 HIV/SIV inserts are not the result of natural evolution and mutations.

Particularly,

- Firstly, the 6 inserts are highly contiguous: 169bp within 275bp regions.
- Secondly, HIV/SIV inserted strains are very homogeneous: Russia, Cote d'ivoire, Netherlands, Malawi origins.
- Thirdly, the functions of inserts are also various: 2 from POL/RT, 4 from ENVELOPPE functional genes.

Not reported in this article, wz found also within the COVID-19 genome 3 others HIV/SIV regions: one from ENVELOPPE, one from RT, and one from INTEGRASE.

**POL/RT, INTEGRASE, and ENVELOPPE: we have here 3 majors functional pieces of genes necessary to build a retrovirus...**

In other hand,

1 / SYMMETRY: strategy of realism (numbers of inserts by increasing frequencies and pathogenicity):

1 SIV

2 HIV2  
3 HIV1

2 / SYMMETRY: economy strategy (sense and antisense in an insert of minimum length):

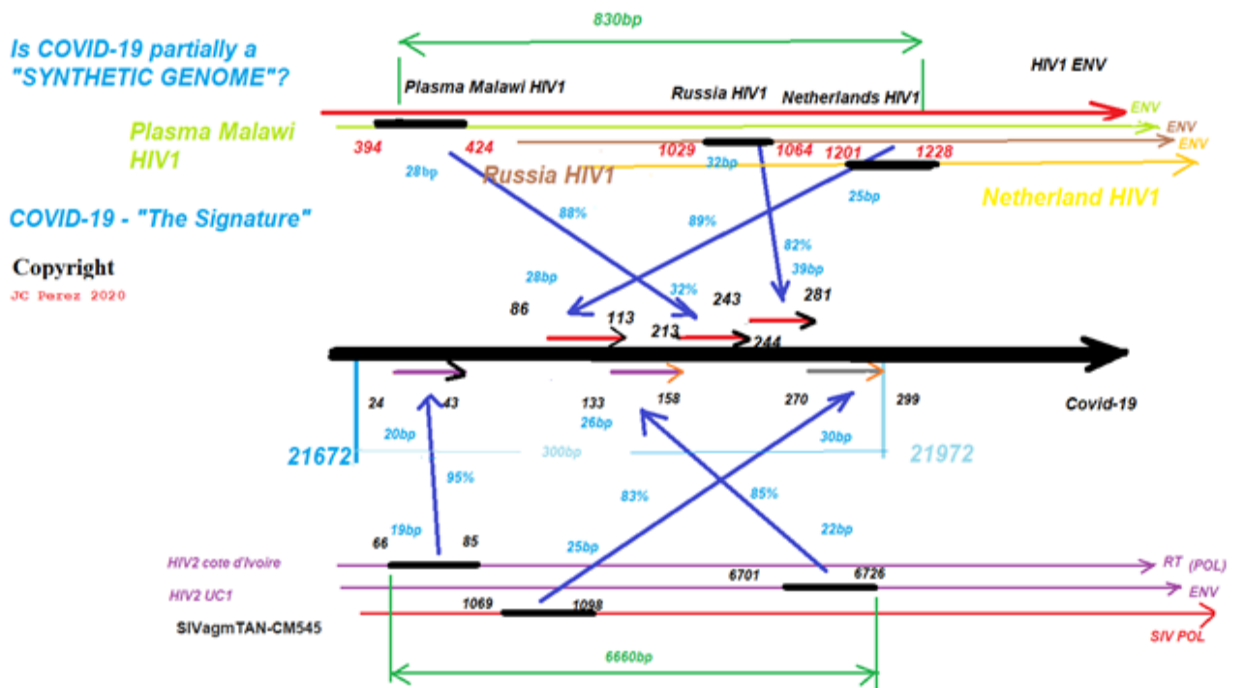
Hiv2a. <---  
Hiv1a. --->  
Hiv2b --->  
Hiv1b <---  
HIV1c <---  
Siva --->

3 / SYMETRY: Symmetry Pol / ENV

The 6 inserts are positioned in this order in Covid-19  
RT(POL) ENV ENV ENV ENV POL

**All this is remarkable and bears the mark of a desire for organization of a human nature:  
LOGIC, SYMETRIES!!!!**

This conclusion is summarized by Figure 49 below.



JC Perez 2020

Figure 49: Is COVID-19 partially a “SYNTHETIC GENOME”?



**We must conclude this remarkable fact:**

To adapt to the disorder of its DNA resulting from the insertion of the 6 HIV / SIVs, the covid-19 genome has most certainly increased its level of global organization by adaptation mutations.

And we will now have to fear that this genome will continue to mutate in order to optimize its overall level of organization ...

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**References**

- [1] (Prashant Pradhan 2020) Prashant Pradhan et al, Uncanny similarity of unique inserts in the 2019-nCoV spike protein to HIV-1 gp120 and Gag, <https://www.biorxiv.org/content/10.1101/2020.01.30.927871v1>
- [2] Lu, R et al., 2020. Genomic characterisation and epidemiology of 2019 novel coronavirus: implications for virus origins and receptor binding *The Lancet*. <https://www.thelancet.com/journals/lancet/article/PIIS0140-6736%2820%2930251-8/fulltext>
- [3] Wang GP, Ciuffi A, Leipzig J, Berry CC, Bushman FD 2007. HIV integration site selection: Analysis by massively parallel pyrosequencing reveals association with epigenetic modifications. *Genome Res* 17: 1186–1194 [PMC free article] [PubMed] [Google Scholar]
- [4] Lyons Weiler J., 2020, <https://jameslyonsweiler.com/2020/01/30/on-the-origins-of-the-2019-ncov-virus-wuhan-china/>
- [5] Wei Ji, et al, Homologous recombination within the spike glycoprotein of the newly identified coronavirus 2019-nCoV may boost cross-species transmission from snake to human, <https://onlinelibrary.wiley.com/doi/pdfdirect/10.1002/jmv.25682>
- [6] Peng Zhou et al, Discovery of a novel coronavirus associated with the recent pneumonia outbreak in humans and its potential bat origin, *BioRxiv*, January 2020, <https://doi.org/10.1101/2020.01.22.914952>, [https://disq.us/url?impression=3e338628-41df-11ea-b46a8a92c9d430bc&product=email&object\\_type=thread&variant=active&forum\\_id=2634513&utm\\_source=digest&adjective=thread\\_title&thread=7832955149&verb=click&event=activity&threadrank=1&user\\_id=80626806&adverb=internal&zone=digest&url=https%3A%2F%2Fwww.biorxiv.org%2Fcontent%2F10.1101%2F2020.01.22.914952v2%3ACWcfg\\_iPzYtYABql\\_6odh4yGKIM&object\\_id=&experiment=new\\_notifications](https://disq.us/url?impression=3e338628-41df-11ea-b46a8a92c9d430bc&product=email&object_type=thread&variant=active&forum_id=2634513&utm_source=digest&adjective=thread_title&thread=7832955149&verb=click&event=activity&threadrank=1&user_id=80626806&adverb=internal&zone=digest&url=https%3A%2F%2Fwww.biorxiv.org%2Fcontent%2F10.1101%2F2020.01.22.914952v2%3ACWcfg_iPzYtYABql_6odh4yGKIM&object_id=&experiment=new_notifications)

- [7] Perez 2018e, Intriguing Humans and Primates chromosomes 4, in Journal of primates, <https://openaccesspub.org/jp/article/746>
- [8] Perez2019a,Perez, J. Epigenetics Theoretical Limits of Synthetic Genomes: The Cases of Artificially Caulobacter (C. eth-2.0), Mycoplasma Mycoides (JCVI-Syn 1.0, JCVI-Syn 3.0 and JCVI\_3A), E-coli and YEAST chr XII. Preprints 2019, 2019070120 (doi:10.20944/preprints201907.0120.v1). <https://www.preprints.org/manuscript/201907.0120/v1>
- [9] Schafer 2014, Schäfer A, Baric RS, Ferris MT. Systems approaches to Coronavirus pathogenesis. *Curr Opin Virol.* 2014;6:61–69. doi:10.1016/j.coviro.2014.04.007 <https://www.ncbi.nlm.nih.gov/pmc/articles/PMC4076299/>
- [10] Suzan R. Weiss 2005, Coronavirus Pathogenesis and the Emerging Pathogen Severe Acute Respiratory Syndrome Coronavirus Susan R. Weiss, Sonia Navas-Martin *Microbiology and Molecular Biology Reviews* Dec 2005, 69 (4) 635-664; DOI: 10.1128/MMBR.69.4.635-664.2005 <https://mmbbr.asm.org/content/69/4/635>
- [11] Fleaux Rachel, « La musique des genes ». *Scientific journal Sciences et avenir* , April 1995.
- [12] Fraser CM, Gocayne JD, White O, Adams MD, Clayton RA, Fleischmann RD, et al. (1995), The minimal gene complement of mycoplasma genitalium. *Science.* 1995;270:397–403. [PubMed] [Google Scholar] <https://science.sciencemag.org/content/270/5235/397>
- [13] Fredens J. et al., Total synthesis of Escherichia coli with a recoded genome, *Nature*, May 2019, DOI <https://doi.org/10.1038/s41586-019-1192-5> <https://www.nature.com/articles/s41586-019-1192-5>
- [14] Robert Friedman and Matthew Cross, (2018), *Nature's Secret Nutrient: Golden Ratio Biomimicry for PEAK Health, Performance & Longevity*, <https://www.amazon.fr/Natures-Secret-Nutrient-Biomimicry-Performance-ebook/dp/B07N8LHDQ1>
- [15] Hutchinson III et al, Design and synthesis of a minimal bacterial genome, *Science* 25 Mar 2016: Vol. 351, Issue 6280, aad6253 DOI: 10.1126/science.aad6253 ; <https://science.sciencemag.org/node/676507.full>
- [16] Gibson DG, Glass JI, Lartigue C, Noskov VN, Chuang RY, Algire MA, et al. (2010), Creation of a bacterial cell controlled by a chemically synthesized genome. *Science.* 2010;329:52–56. [PubMed] [Google Scholar]
- [17] Jee Loon Foo, Synthetic yeast genome reveals its versatility, *Nature* 557, 647-648 (2018), doi: 10.1038/d41586-018-05164-3 <https://www.nature.com/articles/d41586-018-05164-3/>
- [18] Marcer P., *Communications: Order and Chaos in DNA — the Denis Guichard Prizewinner: Jean-Claude Pérez*, *Kybernetes*, Vol. 21, 1992, Issue: 2, pp.60 61, <https://doi.org/10.1108/eb005922>.
- [19] Orsini M. et al, Whole-Genome Sequencing of Mycoplasma mycoides subsp. mycoides Italian Strain 57/13, the Causative Agent of Contagious Bovine Pleuropneumonia. *Genome Announc.* 2015 Mar 26;3(2). pii: e00197-15. doi: 10.1128/genomeA.00197-15. [https://www.ncbi.nlm.nih.gov/pubmed?LinkName=nucore\\_pubmed&from\\_uid=731474337](https://www.ncbi.nlm.nih.gov/pubmed?LinkName=nucore_pubmed&from_uid=731474337)
- [20] Parayon Gabriel, (2011), *On musical self-similarity*, ISBN 978-952-10-6970-3 (PDF) : <https://helda.helsinki.fi/bitstream/handle/10138/26236/onmusica.pdf>
- [21] Pellionisz AJ, Graham R, Pellionisz PA , Pérez JC., *Recursive Genome Function of the Cerebellum: Geometric Unification of Neuroscience and Genomics*. In: Manto M, DL, et al. (Eds) *Handbook of the Cerebellum and Cerebellar Disorders*. 1st (Edn), Springer, USA, 2012.
- [22] Pérez J.C., *De nouvelles voies vers l'Intelligence Artificielle - 1988 - Ed. MASSON Paris - ISBN: 2-225-81326-4*
- [23] Pérez JC (1989). *De nouvelles voies vers l'Intelligence Artificielle*. Ed. Masson Paris Second edition, ISBN: 2-225-81815-0
- [24] Pérez JC (1990a). *La révolution des ordinateurs neuronaux*. Hermes Ed. Paris, ISBN: 2-86601 - 203-8. <https://www.amazon.fr/R%C3%A9volution-ordinateurs-neuronaux-Jean>

- ClaudePérez/dp/2866012038
- [25] Pérez JC (1990b). Integer neural network systems (I.N.N.S) using resonance properties of a Fibonacci chaotic Golden Neuron, Published in: 1990 IJCNN International Joint Conference on Neural Networks , INSPEC Accession Number: 3926657 DOI: 10.1109/IJCNN.1990.137678 Publisher: IEEE Conference Location: San Diego, CA, USA, USA , <https://ieeexplore.ieee.org/document/5726638/>
- [26] Pérez JC (1990c) Digital Holograms Computers: concepts and applications. International conference on Neural Networks - LYON France March 1990; Springer Verlag. ISBN2-287-00051-8 1-8
- [27] Pérez JC,(1991), Chaos DNA and neurocomputers. [https://www.researchgate.net/publication/258439719\\_JC\\_Pérez\\_1991\\_Chaos\\_DNA\\_and\\_Neurocomputers\\_A\\_Golden\\_Link\\_in\\_Speculations\\_in\\_Science\\_and\\_Technologyvol\\_14\\_no\\_4\\_ISSN\\_0155-7785](https://www.researchgate.net/publication/258439719_JC_Pérez_1991_Chaos_DNA_and_Neurocomputers_A_Golden_Link_in_Speculations_in_Science_and_Technologyvol_14_no_4_ISSN_0155-7785)
- [28] Perez j.C, (1997), « L'ADN décrypté » (Decyphered DNA), Resurgence publisher Liege, Belgium. [https://www.amazon.co.uk/LADN-deciphered-security-language-experts/dp/2872110178/ref=asap\\_bc?ie=UTF8](https://www.amazon.co.uk/LADN-deciphered-security-language-experts/dp/2872110178/ref=asap_bc?ie=UTF8).
- [29] Pérez JC (2000). From Prions and Prions-like invariants to the self-assembly thesis. International symposium on Prion Diseases and Elated Processes. Annecy, France.
- [30] Pérez JC (2009) Codex Biogenesis. Resurgence, Liege Belgium. <https://www.amazon.fr/Codex-Biogenesis13-codes-lADN/dp/2874340448>
- [31] Perez J.C., Codon populations in single-stranded whole human genome DNA Are fractal and fine-tuned by the Golden Ratio 1.618., Interdiscip Sci. 2010 Sep;2(3):228-40. doi: 10.1007/s12539-010-0022-0. Epub 2010 Jul 25. <https://www.ncbi.nlm.nih.gov/m/pubmed/20658335/>
- [32] Pérez JC (2011), BIT Life Sciences' 3rd World Congress of Vaccine, Beijing, China" Decoding non-coding Dna Codes:Human Genome MetaChromosomes Architecture " <http://fr.scribd.com/doc/57828784/jcPérezBeijing032011>
- [33] Pérez JC, Applied Mathematics, The “3 Genomic Numbers” Discovery: How Our Genome Single-Stranded DNA Sequence Is “Self-Designed” as a Numerical WholeVol.4 No.10B(2013), Article ID:37457,17 DOI:10.4236/am.2013.410A2004. <http://fr.scribd.com/doc/57828784/jcPérezBeijing032011>
- [34] Perez JC (2015). Deciphering Hidden DNA Meta-Codes -The Great Unification & Master Code of Biology. J Glycomics Lipidomics 5:131. doi: 10.4172/2153-0637.1000131 <https://www.omicsonline.org/openaccess/deciphering-hidden-dna-metacodesthe-great-unification--mastercode-ofbiology-2153-0637-1000131.php?aid=55261>
- [35] Pérez JC (2017a). Sapiens Mitochondrial DNA Genome Circular Long Range Numerical Meta Structures are Highly Correlated with Cancers and Genetic Diseases mtDNA Mutations. J Cancer Sci Ther 2017, 9:512- 527. doi: 10.4172/1948-5956.1000469, <https://www.omicsonline.org/openaccess/sapiens-mitochondrial-dna-genomecircular-long-range-numericalmetastructures-are-highly-correlated-withcancers-and-genetic-disea-1948-5956-1000469.php?aid=90737>
- [36] Pérez JC (2017b). DUF1220 Homo Sapiens and Neanderthal fractal periods architectures breakthrough(2017)SDRP Journal of Cellular and Molecular Physiology <https://www.siftdesk.org/article/details/DUF1220-Homo-Sapiens-andNeanderthal--fractal-periods-architecturesbreakthrough/184>
- [37] Pérez JC (2017c). The exceptional fractal meta organizations of whole chromosomes 4 of Sapiens, Neanderthal and superior primates, Bioinformatics and Computational BiologyLetters, 2017, 1 (1): 1-26, DOI: 10.24896/bcbl.2017112 <https://bacbl.com/index.php/bcbl/article/view/2/2>
- [38] Pérez J C (2017d). A proof of the unity, integrity and autopoietic autonomy of the whole human genome. Research in Biological Chemistry 2017, vol1,no.1,

- [https://www.rbchemistry.com/index.php/rb\\_c/article/view/2](https://www.rbchemistry.com/index.php/rb_c/article/view/2)
- [39] Pérez JC (2017e). The Master Code of Biology: from Prions and Prions-like Invariants to the Self-assembly Thesis. Biomed J Sci&Tech Res1(4)- 2017. BJSTR.MS.ID.000369. DOI: 10.26717/BJSTR.2017.01.000369 <https://biomedres.us/pdfs/BJSTR.MS.ID.000369.pdf>
- [40] Pérez JC (2017f). Decyphering “the MASTER CODE ®” Structure and Discovery of a Periodic Invariant Unifying 160 HIV1/HIV2/SIV Isolates Genomes. Biomed J Sci & Tech Res 1(2)-2017. BJSTR. MS.ID.000209. <http://biomedres.us/pdfs/BJSTR.MS.ID.000209.pdf>
- [41] Pérez JC (2017g). The “Master Code of DNA: Towards the Discovery of the SNPs Function (Single-Nucleotide Polymorphism). J Clin Epigenet. Vol. 3 No. 3:26, <http://clinicalepigenetics.imedpub.com/the-master-codeof-dna-towards-the-discovery-of-the-snpfunction-singlenucleotidepolymorphism.pdf> Pérez JC (2017e), Mannheim Germany 27 November 2017 Biomathematics conference in Mannheim Germany <https://www.cammbio.hsmannheim.de/kick-off-event.html> ... Dr. Jean-Claude Pérez, [https://www.cammbio.hsmannheim.de/fileadmin/user\\_upload/projekte/cammbio/events/20171127-kickoff/EPPérez.pdf](https://www.cammbio.hsmannheim.de/fileadmin/user_upload/projekte/cammbio/events/20171127-kickoff/EPPérez.pdf)
- [42] Pérez JC (2017h). The Master Code of Biology: Self-assembly of two identical Peptides beta A4 1-43 Amyloid In Alzheimer’s Diseases. Biomed J Sci & Tech Res 1(4)- 2017. BJSTR.MS.ID.000394. DOI: 10.26717/BJSTR.2017.01.000394, <https://biomedres.us/pdfs/BJSTR.MS.ID.000394.pdf>
- [43] Perez JC (2017i) Fractal Self-similarity, Scale Invariance and Stationary waves Codes Architecture Human Chromosomes DNA sequences. <https://www.google.com/url?sa=t&source=web&rct=j&url=https://www.semanticscholar.org/paper/Fractal-Self-similarity%252C-Scale-Invariance-and-waves-P%25C3%25A9rez/06e77ac3688dee073fac42081eab399901a12def&ved=2ahUKEwigktxi7XnAhXO34UKHcJBBK0QFjAAegQIBhAC&usq=AOvVaw0vsg7vb-e17DoKUK48ox0E>
- [44] Pérez JC (2018a). Towards a Universal Law Controlling all Human Cancer Chromosome LOH Deletions, Perspectives in Prostate and Breast Cancers screening. Canc Therapy & Oncol Int J. 2018; 9(2): 555756. DOI: 10.19080/CTOIJ.2018.09.555756. <https://juniperpublishers.com/ctoj/pdf/CTOIJ.MS.ID.555756.pdf>
- [45] Pérez JC (2018b). Cancer, Quantum Computing and TP53 Tumor Suppressor Gene Mutations Prediction. Nov Appro in Can Study. 1(2). NACS.000507.2018. <http://crimsonpublishers.com/nacs/pdf/NA CS.000507.pdf>
- [46] Pérez JC (2018c). Neuroblastoma and Glioblastoma Brain Cancers:«Human Genome Optimum» (HGO) a Global Genome Strategy controlling all Human Chromosome LOH Deletions. Nov Appro in Can Study. 1(3). NACS.000512.2018. <http://crimsonpublishers.com/nacs/pdf/NACS.000512.pdf>
- [47] Perez, J.C. (2018d) Six Fractal Codes of Biological Life:perspectives in Exobiology, Cancers Basic Research and Artificial Intelligence Biomimetism Decisions Making. Preprints 2018, 2018090139 (doi: 10.20944/preprints201809.0139.v1). <https://www.preprints.org/manuscript/201809.0139/v1>
- [48] Perez J.C. (2019b), 20219 paper to be published, « The Discovery of a "DNA SUPRACODE" Controlling Locally & Globally Humans Mitochondrial Genomes: Perspectives in Evolution and Cancers Early stage Diagnostic ».
- [49] Sergey V. Petoukhov, Zhengbing Hu, Matthew He, Advances in Artificial Systems for Medicine and Education II, DOI<https://doi.org/10.1007/978-3-030-12082-5>, Springer Nature Switzerland AG 2020, <https://link.springer.com/book/10.1007%2F978-3-030-12082-5#page=40>
- [50] Rapoport D.L., Klein Bottle Logophysics, Self-reference, Heterarchies, Genomic Topologies, Harmonics and Evolution. Part III: The Klein Bottle Logic of Genomics and its Dynamics, Quantum Information, Complexity and Palindromic Repeats in Evolution Quantum Biosystems | November 2016 | Vol 7 | Issue 1 | Page 107-174 107 ISSN 1970-223X

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- [51] Rapoport D. L. and Perez J.C. (2018) Golden ratio and Klein bottle Logophysics: the Keys of the Codes of Life and Cognition. Quantum Biosystems. 9(2) 8-76. Vol. 9 – n.2 – 2018
- [52] Sikela, J. M., The jewels of our genome: the search for the genomic changes underlying the evolutionarily unique capacities of the human brain. PLoS Genetics 2 (2006) doi.org/10.1371/journal.pgen.0020080
- [53] Sleator RD., The story of Mycoplasma mycoides JCVI-syn1.0: the forty million dollar microbe. Bioeng Bugs. 2010;1(4):229–230. doi:10.4161/bbug.1.4.12465
- [54] Strecker J, Ladha A, Gardner Z, Schmid-Burgk JL, Makarova KS, Koonin EV, Zhang F., RNA-guided DNA insertion with CRISPR-associated transposases., Science. 2019 Jun 6. doi: 10.1126/science.aax9181. https://doi.org/10.1126/science.aax9181
- [55] Venetz JE et al, Chemical synthesis rewriting of a bacterial genome to achieve design flexibility and biological functionality, PNAS 2019, doi: 10.1073/pnas.1818259116
- [56] Weiming Zhang et al., Engineering the ribosomal DNA in a megabase synthetic chromosome Science 10 Mar 2017: Vol. 355, Issue 6329, eaaf3981 DOI: 10.1126/science.aaf3981
- [57] Weiss H and Weiss V., The golden mean as clock cycle of brain waves - Chaos, Solitons & Fractals, 2003.

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