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May 2022 : Peculiar Evolution of the Monkeypox Virus Genomes

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KEYWORDS

Monkeypox virus, Biomathematics, Master code, Evolution, Genomics, Proteomics.

ABSTRACT

We compare the evolution of 14 monkeypox virus genomes til that of May 2022 that is currently spreading across humans in numerous countries outside Africa. Our aim was to discover mutations or other viral evolutions (recombination) that may explain the sudden impact of this very low-level circulating epidemic or alert on a potential peculiar pathogenic character.

We have evidenced the presence of a large number of T bases in succession, at the level of the polymerase, between the DNA-dependent RNA polymerase subunit rpo132 and the cowpox A-type inclusion protein, progressively rising from the absence of a characteristically long pattern of T-bases in succession (≤ 10) in the early genomes of 1971, up to 19 T-bases in the Israel 2018 strain of reference, and 30 T bases thereafter in the 2022 strains. We find a complementary match for this long T bases sequence only in the simian hemorrhagic encephalitis virus, at the very 3' end of the genome after the stop codon, with a long succession of 28 A bases. More strikingly, we find that the corresponding 10 phenyl-alanine aa chain is reported as matching uniquely ($E \leq 0.001$) a hypothetical protein element in *Plasmodium falciparum*, *Yersinia pestis*, *Escherichia coli* and *Penicillium nordicum*. We wonder about the possibility that this region of the monkeypox genome may potentially code for a not yet identified polypeptides with a functional role situated right upstream this long T-repeat.

INTRODUCTION

Monkeypox is a zoonotic disease caused by the monkeypox virus, an orthopoxvirus closely related to variola virus, the causative agent of smallpox. Monkeypox was first discovered in 1958 in monkeys, although they are not the source of the virus. Human cases were first described in 1970. There are 2 strains of monkeypox: the West African and Central African strains.

Several cases of monkeypox have been identified in various geographically countries. In May 2022 cases were reported in Australia, Austria, Belgium, Canada, Denmark, France, Germany, Greece, Israel, Italy, the Netherlands, Portugal, Spain, Sweden, Switzerland and the U.K (NCBI, 2022), (Antwerpen M, et al, 2022), (Isidro et al, 2022).

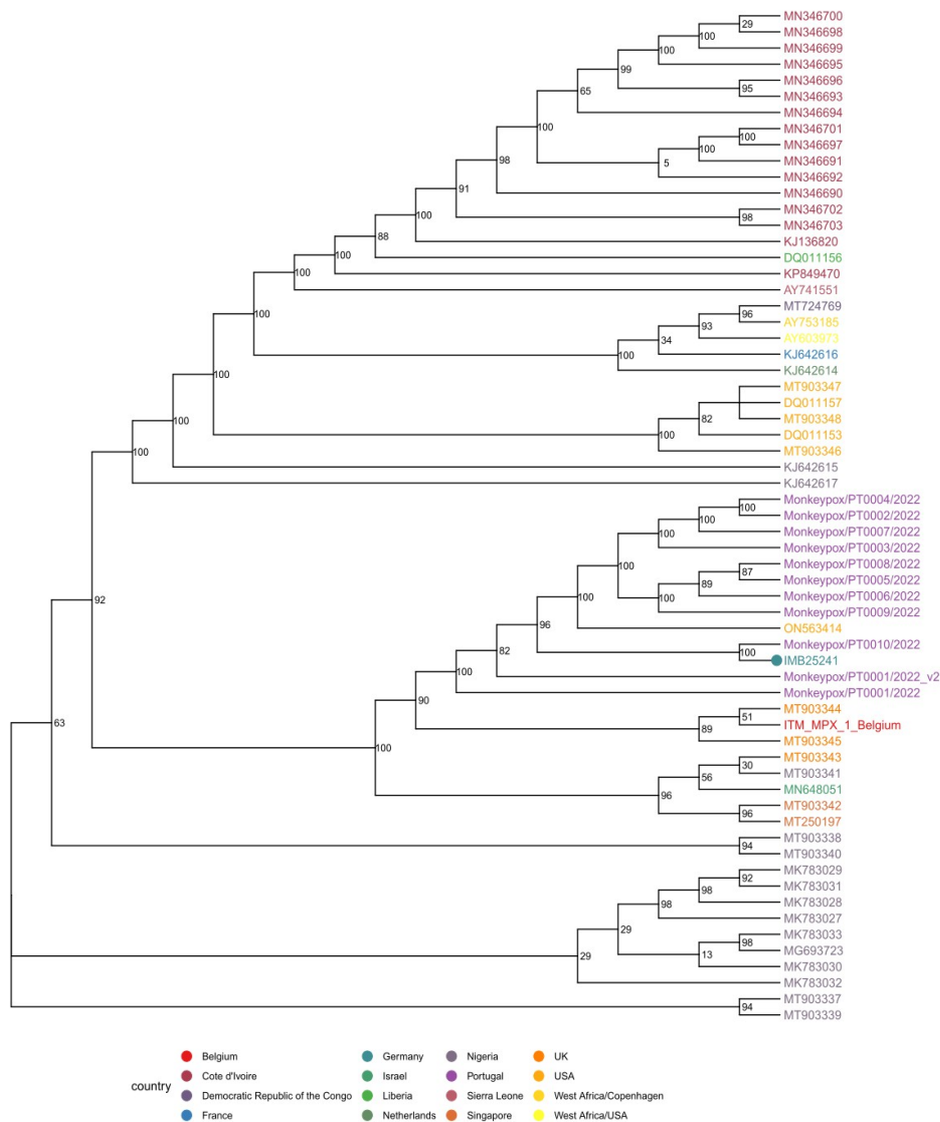


Figure1 – Monkeypox tree (from <https://virological.org/t/first-german-genome-sequence-of-monkeypox-virus-associated-to-multi-country-outbreak-in-may-2022/812>)
Nextstrain reference tree <https://nextstrain.org/monkeypox?s=03>

Monkeypox is classified as a zoonotic disease where transmission of the virus is usually due to animal-human contact. Genetically, monkeypox viruses cluster into two groups: the Congo basin and the west African clade.

Monkeypox virus

Monkeypox virus Zaire-96-l-16

This particular outbreak has been identified as due to a virus from the west African clade which is often associated with milder disease and, in this case, human-to-human spread is suspected. The first referenced human to human strain was located in Israel in 2018: a case of monkeypox in a man who returned from Nigeria to Israel in 2018 (Erez et al, 2018).

MATERIALS and METHODS

Monkeypox strains analyzed :

We analyzed 14 monkeypox whole genomes:

Gabon 1988 alias 2015 KJ642619.1

<https://www.ncbi.nlm.nih.gov/nuccore/KJ642619.1>

Cameroun 1990 alias 2015 KJ642618.1

<https://www.ncbi.nlm.nih.gov/nuccore/KJ642618.1>

Liberia 1970 DQ011156.1

<https://www.ncbi.nlm.nih.gov/nuccore/DQ011156.1>

Nigeria 1971) alias 2015 KJ642617.1

<https://www.ncbi.nlm.nih.gov/nuccore/KJ642617.1>

2018 Israel MN648051.1

<https://www.ncbi.nlm.nih.gov/nuccore/MN648051.1>

Zaire 2009 alias 2020 NC_003310.1

https://www.ncbi.nlm.nih.gov/nuccore/NC_003310.1

Rivers state 2020 MT903340.1

<https://www.ncbi.nlm.nih.gov/nuccore/MT903340.1>

UK 2020 MT903344.1

<https://www.ncbi.nlm.nih.gov/nuccore/MT903344>

USA 2022 ON563414.1

<https://www.ncbi.nlm.nih.gov/nuccore/ON563414.1?report=GenBank&s=03>

German 2022 ON568298.1

<https://www.ncbi.nlm.nih.gov/nuccore/ON568298>

Singapore 2020 MT903342.1

<https://www.ncbi.nlm.nih.gov/nuccore/MT903342.1?report=genbank>

Nigeria 2018 MG693723.1

[https://www.ncbi.nlm.nih.gov/nucleotide/MG693723.1?report=genbank&log\\$=nuclalign&blast_rank=1&RID=98T6WWFV016](https://www.ncbi.nlm.nih.gov/nucleotide/MG693723.1?report=genbank&log$=nuclalign&blast_rank=1&RID=98T6WWFV016)

UK 2020 MT903345.1

[https://www.ncbi.nlm.nih.gov/nucleotide/MT903345.1?report=genbank&log\\$=nuclalign&blast_rank=1&RID=98TT3F4E013](https://www.ncbi.nlm.nih.gov/nucleotide/MT903345.1?report=genbank&log$=nuclalign&blast_rank=1&RID=98TT3F4E013)

France 2022 ON602722.1

<https://www.ncbi.nlm.nih.gov/nuccore/ON602722.1?report=genbank>

Biomathematics methods, The Master Code analysis :

The "Master Code" method (Perez, 2009), (Perez, 2015) and (Perez&Montagnier, 2021) allows, from the atomic masses common only to DNA, RNA and amino acids numerical values, to highlight a META-CODE which would unify the 3 codes of DNA, RNA and amino acid sequences.

Particularly, the Master code coupling curves measures the level of correlation unifying the expression of 2 Genomics (DNA) and Proteomics (amino acids) for any sequence, coding for a protein, or not.

In (Perez, 2017a) we analyzed all types of Prions in the early 2000s mad cow disease (plants, yeast, humans, cows, sheep, etc.). We had then highlighted a "signature" or sort of invariant which would be common to all Prions: a typical signature of the Master code taking the characteristic form of a "W" (or even of an "M" symmetrically). We had extended this type of analysis to amyloid_s implicated in Alzheimer's disease (Perez, 2017b).

RESULTS

Table 1 – Evolution of the « T » bases contiguous region for the 14 analysed genomes.

Name	Genbank ID	Start T location	Number of T
Gabon1988 (2015)	KJ642619.1		0
Cameroun1990 (2015)	KJ642618.1		0
Liberia1970	DQ011156.1		0
ZAire2009	NC_003310.1		0
Nigeria1971 (2015)	KJ642617.1	133245	27
Israel2018	MN648051.1	133298	19
Rivers state 2020	MT903340.1	133081	25
UK2020A	MT903344.1	133081	27
Singapore2020	MT903342.1	133093	28
Nigeria2018	MG693723.1	126745	29
UK2020B	MT903345.1	133100	28
France2022	ON602722.1	132972	19
USA2022	ON563414.1	133094	30
Germany2022	ON568298.1	133201	30

The last 3 cases analyzed date from May 2022. It is of note that the 2022 French genome is limited to a succession of 19 T. But in fact this sequence may also accept C bases substituted for T as both ttt and ttc codons are translated in phenyl-alanine residue. In that respect the length of the French sequence is actually equivalent to 21T. Sequencing errors are possible but not to that extent over 8 nucleotides. So the difference of the French sequence raises some question as it is obviously not the same as the other strains in that respect. It is also the case for the Italian sequence (ON622721 from <https://www.ncbi.nlm.nih.gov/nuccore/ON622721.1/>).

DISCUSSION

This is by chance that we have discovered the presence of a 30-T long sequence in the middle of the USA2022 monkeypox genome, between the DNA-dependent RNA polymerase subunit rpo132 and the cowpox A-type inclusion protein, before a gene complement region that may become coding under circumstances that need to be specified by expert in the field.

For instance, if we look at the monkeypox strain Gabon-1988 we can identify in this region a sequence of nucleotide coding straightforwardly for a 42-aa long polypeptide that may constitute a small protein.

Number of codons : 42

MGYLRSFYKRFHVPDHVQPSYVSPSLYRVYQSSSLEGD RTP .

Monkeypox virus strain Gabon-1988, complete genome

GenBank: KJ642619.1

gene 131162..134656 /product="DNA-dependent RNA polymerase subunit rpo132"

```

      tatt gtttagtaga tactcatcaa gattatcaag ataagctaat
134701 tcacttaaaca tattatcggg ttcgggtattg ttactcgaga atagagttcg ttatgctcct
134761 gatattcggg aatctgtgga gtttcagggt ttggtggaag tgtaactgct acttgggtggg
134821 atactgaagg atatttcaga gagttgtgga tgttcgggtt cgacatccac cgatggtgtc
134881 acgccacttaa tcggttcggt aacgtctgtg gatgaggtg ctacttctac agaacctgta
134941 gcctcagttg tcaacggaga tacatcttca atgcgcgaa atgtataaatt tggtaatggt
135001 ttctcatgtg gatcttaaga agaagaggta agatatctac gaaagatacc gatcacgttc
135061 tagttctctt ttgtagaact ttaacttttt ctttctcagc atcttagttga tattccgacc
135121 tcttcacggt tcacatgggt tacctccgca gttttttacaa gcgatttcac gttccagatc
135181 acgttcagcc ttcatacgtc tctccctctc tctatcgagt ttatcagagc agtctttctg
135241 aaggcgatcg aactcctaa atttctccaa cgctttgaatt gtttccatag atttccgaag
135301 tttagcttct aggacggcga ttcttttttc tttcgaattc acgggttaca accgtttcca
135361 ttaccaccat ctctacgttt cttttcttaga tcggcaatct ttctcaacat ttcattccca
135421 tgccttttca ttctcagagt ctatcgtcgt cgaaatatcg ttccagctcc ttttcgacct
135481 caataaatt agcacgttgt ctcacaaagc tctctcttgt agtactatca tttttatctg
135541 attccctggc acgtttaaga tcttatgta attgagtcag ctcttgacac aatctcttaa
135601 ctaaacttct ctcttgcttc ttcgtcatag tacttacaat cactatggga tccattgta
135661 ccacgtctgt actcggcgag ctcacgttta agagattcaa tttccagttt gtacattgat
135721 ttcattatta cgtccgcagt cgttcaactg tatttcaaga tcttgagattc tagattgtaa
135781 tctctgtagc atttccacgg cattcactca gttgtctttc aagatcttgag attcttagatt
135841 ggagctctgct aatctctgta agatttcctc ctccgctctc gatgcagtcg gtcaacttat
135901 tctcttagttc tcttaaacgt gaacgcagtg catcaacttc ttgtgtgtct tcttgattgc
135961 gtgtgcattc atcgagtcta gattcgagat ctctaactg tctcgttct tcttcaagtt

```

gene complement(135770..137860) /product="cowpox A-type inclusion protein"

Figure 2a – Genome sequence extract of monkeypox strain Gabon-1988 potentially coding for a small protein after the DNA-dependent RNA polymerase subunit rpo132 and before the gene complement.

Number of codons : 42

MGYLRSFYKRFHVPDHVQPSYVSPSLYRVYQSSSLEGD RTP .

Monkeypox virus isolate MPXV_USA_2022_MA001, complete genome

GenBank: ON563414.3

[Gene](#) 128941..132435 /note="A25R RNA polymerase subunit (RPO132) (Cop-A24R) RNA polymerase, 132 kDa subunit similar to [Vaccinia virus strain Copenhagen A24R](#)"

[gene complement](#) (133217..133444) /note="A-type inclusion protein (Cop-A25L); A26L" "MDPIVIVSTMTKKQERKLVKRLRQELTQLHEDLKRVRSDKNDSTTRESLMKQRAKVIEVEKELERYFDDNRLEE"

```

      attg tttagtagat actcatcaag ataagctaat tcactaaaca
132481 tattatcgga ttcgggtattg ttactcgaga atagagttcg ttatgctcct gatattcgga
132541 aatctgtgga gtttcaggtt ttggtggaag tgtaactgct acttgggtggg atactgaagg
132601 atatttcaga gagttgtgga tgttcgggtt cgacatccac cgatgggtgtc acgccactaa
132661 tcggttcgg aacgtctgtg gatggaggtg ctacttctac agaacctgta gcctcagttg
132721 tcaacggaga tacatattca atgcgcgaa atgtataatt tggtaatggt ttctcatgtg
132781 gatcttaaga agaagaggta agatatctac gaaagatacc gatcacggtt ctagttctct
132841 ttgttagaac tttaactttt tctttctcag catcttagttg atattccgac ctcttcacgt
132901 ttcgatggg ttacctcgc agtttttaca agcgatttca cgttccagat cacgttcagc
132961 cttcatacgt ctctccctct ctctatcgag tttatcagag cagtctttct gaaggcgatc
133021 gaactccata aatttctcca acgctt tgat tgtttccata gatttccgaa gtttagcttc
133081 taggacggcg attcttttt tttttttttt tttttttttt ttcgaattca cggggtacaa
133141 ccgtttccat taccaccatc tctatgtttc ttttc tagat cggcaatcct tctcaacatt
133201 tcatcccat accttttca
      agt
```

```

gene complement →      agtt tcctcgagtc tattgtcgtc gaaatatcgt tccagctcct
133261 tttcgacctc aataacttta gcacgttgtt tcatcaagct ctctcttgta gtactatcat
133321 ttttatctga ttccctgaca cgtttaagat cttcatgtaa ttgagtcagc tcttgacgca
133381 atctcttaac taacttcctc tcttgcttct tcgcatagt acttacaatc actatgggat
133441 ccat
```

Figure 2b – Genome sequence extract of monkeypox strain USA2022 potentially coding for a small protein after the DNA-dependent RNA polymerase subunit rpo132 and before the gene complement.

Number of codons : 42

MGYLRFSFYKRFHVPDHSVPSYVSPSLYRVYQSSLSEGDRTP.

This growing pattern of T-bases in succession follows a conserved nucleotide sequence that is conserved and may code for a small protein. The functional role of this pattern at the viral genome level is unknown to us.

While it long repeat are common finding at the terminaison of a genome, as for instance at the end of the monkey encephalitis virus, it is almost never encountered fully inside a sequence.

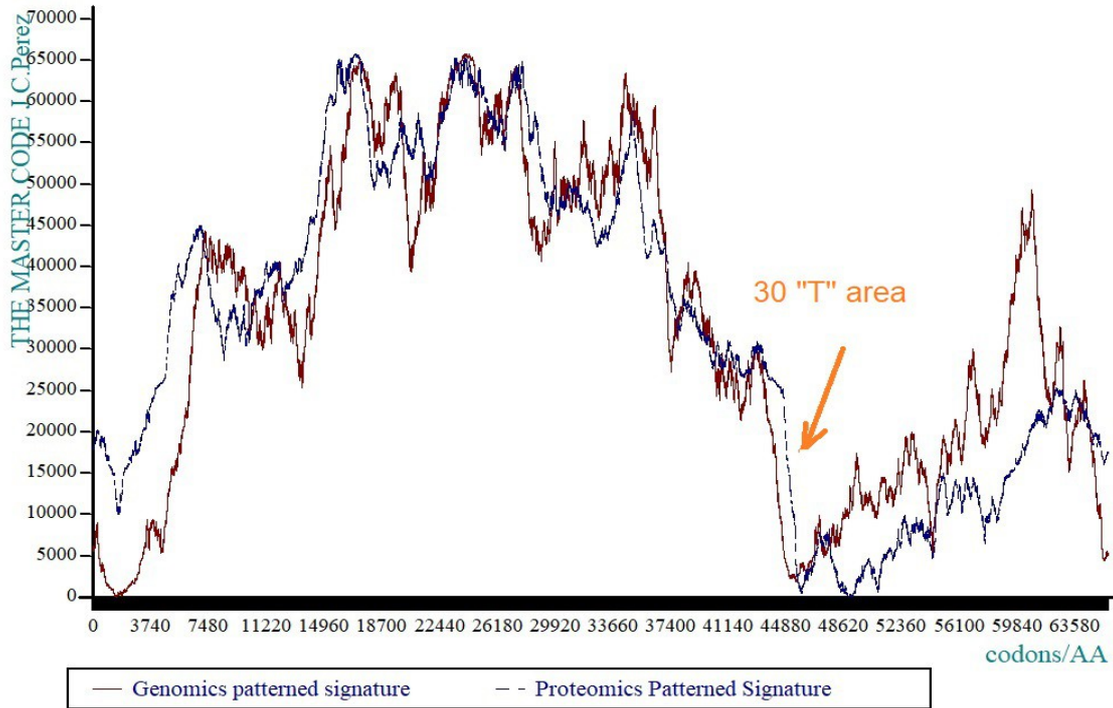


Figure3a – Master code analysis of the whole USA2022 Monkeypox genome. The region 44000 amino acids where there is the 30 T bases insert.appears to be highly functional.

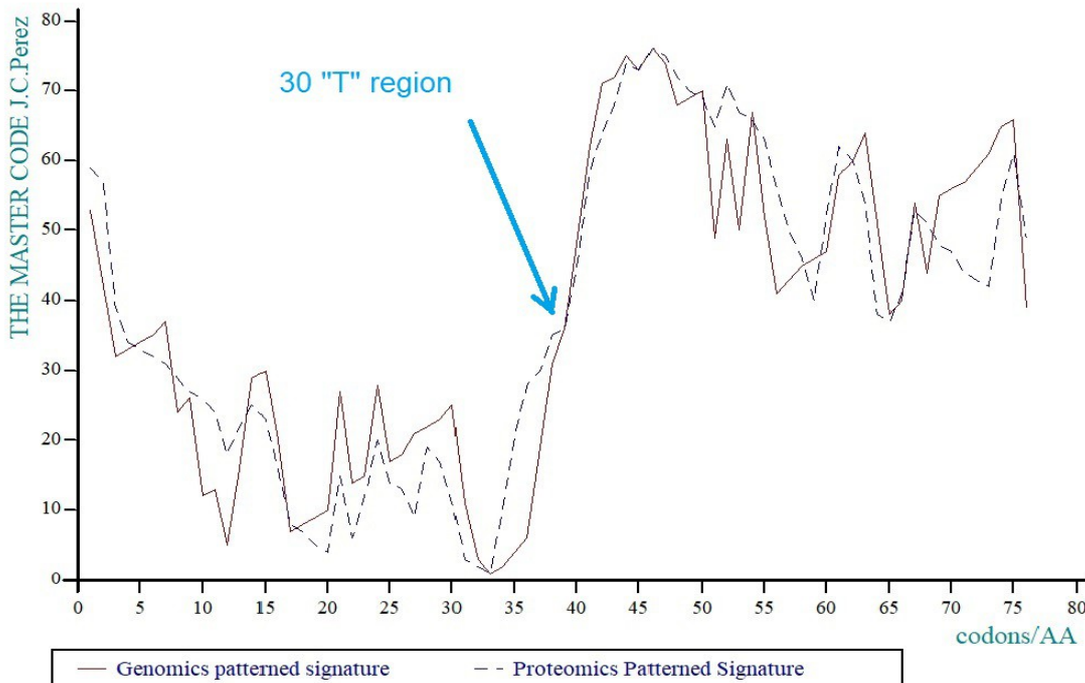


Figure3b – 100 bases upload and download the 30 T bases region in USA2022.

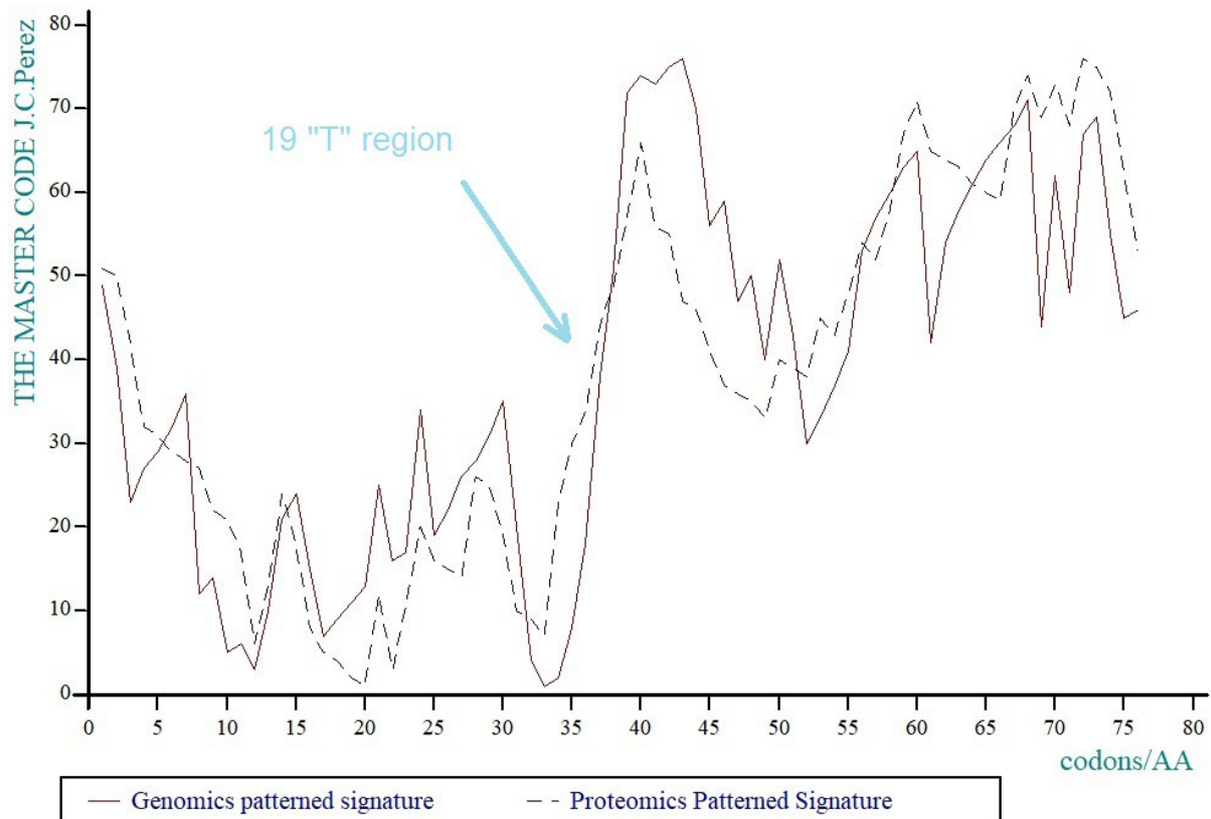


Figure4 - 100 bases upload and download the 19 T bases region in FRANCE2022.

CONCLUSIONS

The objective was here to present a genome characteristic that may partly explain the sudden propagation of the monkeypox virus in the form we observe in May 2022 in quite a number of countries.

The role of the peculiar 30-T base long sequence right in the middle of the virus genome is still to be determined.

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