

# Anomalies in BatCoV/RaTG13 sequencing and provenance

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## ABSTRACT

To this date, the most critical piece of evidence on the purported “natural origin” theory of SARS-CoV-2, was the sequence known as RaTG13, allegedly collected from a single fecal sample from *Rhinolophus Affinis*. Understanding the provenance of RaTG13 is critical on the ongoing debate of the Origins of SARS-CoV-2. However, this sample is allegedly “used up” and therefore can no longer be accessed nor sequenced independently [1], and the only available data was the 3 related Genbank accessions: MN996532.1, SRX7724752 and SRX8357956.

We report these datasets possessed multiple significant anomalies, and the provenance of the promised claims of RaTG13 or its role in proving a “probable bat origin”[2] of SARS-CoV-2 can not be satisfied nor possibly be confirmed.

## RESULTS

### Anomalous enrichment of telomere-like repeat sequences in the dataset SRX7724752

```
>gnl|SRA|SRR11085797.3.1 3 (Biological)
```

```
CTAACCCCTAACCCCTAGCACTATCCTGTTTCCAACCCCAACCCCTAACCCCTCACCCCTAACCC  
TAACCCAGCCGTGTTTCATACCTTAACCTCGCACCTCATCGCTAACCCAGCCCTCACCCG  
ATCCTGTTTCTCTCCCGAACATAACCCCT
```

```
>gnl|SRA|SRR11085797.3.2 3 (Biological)
```

```
GGTTAGGGTTAGGGTTAGGGTTGGAACAGGATAGGGTTAGGGTTAGGGTTAGGGTTAGGGTTAGG  
GTTAGAGTTAGGGTTGGAAACAGGATAGGGTTAGGGTTAGGGCGAGGGATAGGGATAGGG  
AGGGAAACAGGATAGTGGGAGGGCTAGGGGT
```

>gnl|SRA|SRR11085797.8.1 8 (Biological)

GTTAGGGTTAGGGTTAGGGTTAGGGTTGGGTTGGATACAGGATATGGTTAGGGTTAGGG  
GTAGGGTCAGGGTTAGGATTGGAAACGAGATAGGTTACGTGATAGGGTTAGCGTTAGGGT  
TAGGTTTAGTAATCCGCAACGGCTTAGGGTT

>gnl|SRA|SRR11085797.8.2 8 (Biological)

CCTAACCCTAACCCTAACCCTAACCCTAACCCTAACCCATCCTGTTCCCAACCCTAACC  
CTAACCCTAACCCTAACCCTAACACAAAACATAACCCTAACCCCAACCCAAACCCTAACC  
CCATCTTTACTCACACCCTAACCCAAACTC

>gnl|SRA|SRR11085797.10.1 10 (Biological)

GTTAGGGTTAGGGTTAGCGTTAGGGTTAGAAACAGGATAGGGTTAGGGTTAGGGTTAGGG  
TTAGGGTTTTGGTTGGTCACAGTGTTCGCTAGGCATAGGGATAGGGTTCCGTTAGGGT  
TAGGGTTAGGATTCGGAAGAGCTAGCTAAA

>gnl|SRA|SRR11085797.10.2 10 (Biological)

GTTCCCAACCCTAACCCTAACCCTAACCCTAACCCTAACCCTTTCCTTTTTCCAACCCTA  
ACACTAACCCTAACACTAACCCTAACCCCAACCCTACCCTATACTATATCCGACTCTCA  
CGCTAACACTAACATAAGTAATCACAAATT

>gnl|SRA|SRR11085797.13.1 13 (Biological)

TAACCCTAACCCTAAGCGTAAACCTAATCCAATCCTGTTCCCAACCCTAACCCTAACCC  
GACCCTAAGCTTTTTCCCGACCCGAACGCCGACCCGATCCGCCACCCTAACCGTAACCCGT  
TCCCAACCCCTCCTACTGCTCGATCCGCCT

>gnl|SRA|SRR11085797.13.2 13 (Biological)

GGAACAGGATAGGGTTAGGGTTAGGGTTAGGGTTAGGGTTAGGGTTAGGGTTAGGGTTAGG  
GTTAGGGTTAGGGTTGGGATCAGGATAGGGATAGGGATAGGGATAGGGATAGGGTTAGGG  
TGGGGAACAGGAGAGCGTTAGGCAAGG

>gnl|SRA|SRR11085797.14.1 14 (Biological)

GGTTAGGGTTAGGGTTGGAGAAGAGGATAGGTTTAGGGTTTAGGGTTAGGGTTAGGGTTAGG  
GTTAGGGGAGAGGGTTAGCTACACGATAGGAGTAGGGTAACGATTAGGGTTAGGGTTAGGT  
TTGGAAAAAGCATAGGCTATGAGGTACGGT

>gnl|SRA|SRR11085797.14.2 14 (Biological)

CTGCTTCCAACCCTAACCCTAACCCTAACCCTAACCCTAACCCTAACCCTAACCCTAACCC  
CTATCCTGTTCCCAACCCTAACCCTAACCCTAACCCCAACCCTAACCCCAACCCAAACCC  
AACCCTAACCCCAACCCATACCCCAACCAT

>gnl|SRA|SRR11085797.15.1 15 (Biological)

TGTTCCCAACCCATAACCTAAACCTAAGCCGATCCTGTTCCCAACCCATAACCTAACCCT  
ATCCTGTAACAACCCCCACCCATAAAACATCCTCGTACAAACCCATAACCAACCCCCAT  
CCCAAACCCACATACCCGTCACGAACCCACCC

>gnl|SRA|SRR11085797.15.2 15 (Biological)

GTTGGGGTTAGGGTTGGGAACAGGATAGGGTTAGGGTTAGGGTTAGGGTTGGGGTCAGGG  
TTGGGACAAGGACGGGTAGGGGGTTAGGGTAGTGCACAGGGTAGGGAGTGGGGTAGGGTT  
GGGAACAGAGAAGGGACAGTGGGGGGAGTGG

>gnl|SRA|SRR11085797.16.1 16 (Biological)

CTAACCCATAACCCTACCCCTATCCTGTTCCCTAACCCGAACCCATAACCCATAACCC  
TAACCCTCACCTGTTCCAGACCGTAATGCTAACCTTAACACTATCCTGTGCGCTACCCCG  
ACCCTAACCCTCAGCCGACGCGTCACGCCCG

>gnl|SRA|SRR11085797.16.2 16 (Biological)

GGTAAGGGTTAGGGTTAGGATAGGGTTAGGGTTAGGGTTAGGGTTAGGGTTAGGGTTAGG  
GTTGGGAACAGGATAGGGTTAGGGTTAGGGTTAGGGTTTGAACAGGATAGGGTTAGGGA  
TAGGGGTAGAGATAGGGTGAGGTGGTGGAA

>gnl|SRA|SRR11085797.17.1 17 (Biological)

AACCCATAACCCATAACCCTAACGCTATCATGATCCCATCCCTAACCCATAACCCATAACCC  
ACCCTAATACTAACCCTACCCTTTTCATCTCCCTTACACTACCCCCAACACGCCACCCAT  
CCCCAACCACTATGCATGCACTGTCCATAAC

>gnl|SRA|SRR11085797.17.2 17 (Biological)

GGTTAGGGTTGGGAACAGGATAGGGTTAGGGTTAGGGTTAGGGTTAGGGTTAGGAAACAGGATAGGG  
GTAGGGTTAGGGCTAGGGTGAGAAACAGGGTAGGGGTAGGGTGAGGATAAGGGATAGGGT  
TGGGGTTGGGAACAGAGAAGGGGAAGGGCA

>gnl|SRA|SRR11085797.18.1 18 (Biological)

CTAACCTGTTCCCAAACTTAAATCCAATCCTAACCCATCCTGTTCCCAACCCATAACCC  
AAACCTATACCTATCCTGCCCCACACCCGACCCATACACCACCCATAAACGCAACCCTA  
ACCCCATCCTGTTATCGAAGCATACCCCCAC

>gnl|SRA|SRR11085797.18.2 18 (Biological)

GTTAGGGTTAGGGTTAGGGTTGGTAACAGGATAGGGTTAGGGTTAGGGTTAGTGTGGGA  
ACAGGATAGGGGGAGGGAGAGGGTTAGGGTTGGGAAGAGGATAGGGATAGGGGTAGGGAG  
AGGGTTGGGGATAGGGAAGAGAGAGGAG

>gnl|SRA|SRR11085797.19.1 19 (Biological)

CCTAACCCATAACCTAACCCCTCCTGTTTCCAACCAATAACCCATAACCCATCCTAACCC  
CTAACCCACTCCTGTTCTTAACACTAACCTTAACCTCTGAGCTCATCCCCAACCCATAACCA  
TAACCCACCAGTTCGGATACCATCACCCCC

>gnl|SRA|SRR11085797.19.2 19 (Biological)

AGGGTTAGGGTTGGAAACAGGATAGGGTTAGGGTTAGGGTTAGGGTTAGGGTTAGGGTTG  
GGGACAGGATAGGGTTAGGGTTAGGGTTAGGGTTGGGGTTAGGGTTAGGTGTGGGGTTGG  
GATTGGGTATGGGTAGTGGTCAGGGATAGTG

```

>gnl|SRA|SRR11085797.20.1 20 (Biological)
CCTGTTTCCAACCCCTCACCCTGACACTGACCCTAACACTAACCCTAACCCGATC
CTGTTTCTGACCCTAACGACAAGCCTGGCACTAAACTGATCGCGTTTCCAATCGTTACCG
CTTCCCTAACACCCGTCTGTGAAGATACTCCG

>gnl|SRA|SRR11085797.20.2 20 (Biological)
CTTTAGGTTTGTGTTAGGGATAGGGTTAGGGTTAGGGTTAGGGTTAGGGTTAGGGTTAG
GGTTGGGAACAGGATAGGGTTAGGGTTAGGGATAGGGTTGGGGTCTGGATAGGGTTGGGG
GTAGGGTTAGAGTGAGTGTGGGCAGCAGCG

```

Figure 1: The reads that contained Telomere-like repeat sequences within the first 20 reads of SRX7724752.

Despite the theoretical presence of traces of Telomere-like repeats in total RNA of most cells, such repeats comprise only a tiny fraction of the total cellular RNA within real biological samples, and normally does not show up in the first 100 reads. RaTG13 contained an anomalous amount of such repeats, which comprises 63% of the dataset and exist in nearly any set of 10 reads within this dataset. In comparison, the next highest content of such repeats within any other sample of similar context on NCBI, contained merely 4% of these repeats, which does not show up in the first 20 reads of the dataset. Telomere-like repeats are not detected in the first 100 reads of any other datasets examined.

In comparison, the related SRX7724693 lacked such reads within the first 100 reads of the dataset.

```

>gnl|SRA|SRR11085736.100.1 100 (Biological)
CTACTGTGTCTATCCCATTTCCAAAACGCCTTATTGGCGGTACAGGAATATCAACCTGTTGT
CCATCACCTACGCCTTTCGGCCTCGGCCTAGGTCTCTGACTAACCCAGGGCAGAAGAACCCT
TCCCCCTGGAAACCTTGGGTGTGACGGCCCCGTG

>gnl|SRA|SRR11085736.100.2 100 (Biological)
ATCCACGGGCCGTAAACCCAAAGGTTTCCAGGGGAAGGTTTCGTCCGCCCTGGGTAGTCA
GGACCTAAGCCGACGCCGAAAGGCGTAGGTGATGGACAACAGGTTGATATTCCTGTAACC
GCAATAAGCGTTTGAGAGATGGGATGACAGT

```

Figure 2: the first 100 reads in SRX7724693 did not show any Telomere-like repeats.

In addition, SRX7724752 contained 6% all-N sequences that were exactly 35nt long, which is not found at levels any close in other datasets that had the same design section.

### Reads (separated)

```

>gnl|SRA|SRR11085797.11.1 11 (Biological)
NNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNN
>gnl|SRA|SRR11085797.11.2 11 (Biological)
NNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNN

```

Figure 3: an example of All-N read in SRX7724752.

## Anomalous enrichment of non-attributable and low-match data within SRX7724752

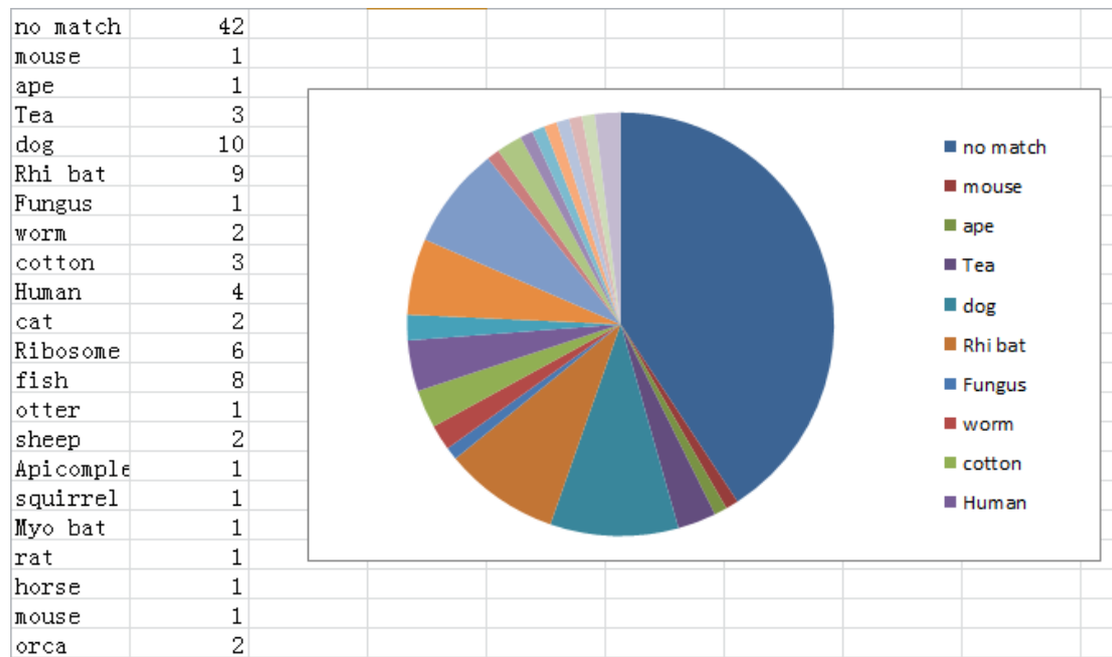


Figure 4: BLAST result of 100 random reads obtained from RaTG13 using BLASTn.

In addition to the anomalous enrichment of repeats, The vast majority of the non-repeat sequences in SRX7724752 does not show any clear matches when examined using BLASTn. With matching results ranging from nearly all domains of life—all of which were partial and low-quality matches, including that of bats.

Only 2 out of 7 Non-repeat and non-PolyN sequences from the first 20 reads from SRX7724752 had any matches, and the match was only partial matches to certain hypothetical proteins

Description

Molecule type

Query Length

Other reports [Distance tree of results](#) [MSA viewer](#) [?](#)

Percent Identity  to

E value  to

Query Coverage  to

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**Descriptions** | [Graphic Summary](#) | [Alignments](#) | [Taxonomy](#)

**Sequences producing significant alignments** Download  Manage Columns  Show  [?](#)

select all 6 sequences selected [GenBank](#) [Graphics](#) [Distance tree of results](#)

Description	Max Score	Total Score	Query Cover	E value	Per. Ident	Accession
<input checked="" type="checkbox"/> <a href="#">Eimeria mitis hypothetical protein conserved partial mRNA</a>	102	102	44%	4e-18	94.03%	<a href="#">XM_013494305.1</a>

Results for: 1:icl|Query\_61914.gn|SRA|SRR11085797.7.1.7 (Biological)(150bp) ▾

Program: BLASTN [Citation](#) ▾

Database: nt [See details](#) ▾

Query ID: icl|Query\_61914

Description: gn|SRA|SRR11085797.7.1.7 (Biological)

Molecule type: dna

Query Length: 150

Other reports: [Distance tree of results](#) [?](#)

Organism: *only top 20 will appear*  exclude

Type common name, binomial, taxid or group name

[+ Add organism](#)

Percent Identity:  to  E value:  to  Query Coverage:  to

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select all 2 sequences selected

[GenBank](#) | [Graphics](#) | [Distance tree of results](#)

Description	Max Score	Total Score	Query Cover	E value	Per. Ident	Accession
<input checked="" type="checkbox"/> <a href="#">Eimeria mitis hypothetical protein conserved partial mRNA</a>	73.1	73.1	40%	3e-09	88.33%	<a href="#">XM_013494305.1</a>
<input checked="" type="checkbox"/> <a href="#">Cyprinus carpio genome assembly common carp genome scaffold 000012284</a>	73.1	73.1	40%	3e-09	88.33%	<a href="#">LN595243.1</a>

Figure 5: the BLASTn result of the 2 non-repeat and non-PolyN sequences in the first 20 reads of SRX7724752. The rest can not be matched to any known organisms.

**Depletion of bacterial-like reads in SRX7724272 which is inconsistent with fecal samples prepared using the methods as indicated by the “Design” section of the SRX7724752 metadata.**

Fecal matter [3], is primarily bacteria by composition. All other fecal swabs prepared using the methods indicated by the metadata correctly showed the presence of bacteria as the majority of the reads. In contrast, SRX7724272 contained only 0.65% bacteria-like reads, all of which were 16S rRNA.





**Description**

gn|SRA|SRR11085797.11550005.1 11550005 (B ...

**Molecule type**

dna

**Query Length**

151

**Other reports**

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**Descriptions** | Graphic Summary | Alignments | Taxonomy

**Sequences producing significant alignments** Download Manage Columns Show 100 ?

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	Description	Max Score	Total Score	Query Cover	E value	Per. Ident	Accession
<input checked="" type="checkbox"/>	<a href="#">PREDICTED: Phyllostomus discolor 28S ribosomal RNA (LOC114512504)_rRNA</a>	279	279	100%	2e-71	100.00%	<a href="#">XR_003685809.1</a>
<input checked="" type="checkbox"/>	<a href="#">PREDICTED: Phyllostomus discolor basic proline-rich protein-like (LOC114512442)_mRNA</a>	279	279	100%	2e-71	100.00%	<a href="#">XM_028531404.1</a>
<input checked="" type="checkbox"/>	<a href="#">Homo sapiens lncAB370.3 lncRNA gene complete sequence</a>	274	274	100%	9e-70	99.34%	<a href="#">MK280359.1</a>
<input checked="" type="checkbox"/>	<a href="#">Homo sapiens lncAB366.1 lncRNA gene complete sequence</a>	274	274	100%	9e-70	99.34%	<a href="#">MK280356.1</a>
<input checked="" type="checkbox"/>	<a href="#">Felis catus Senzu DNA_chromosome: E1_American Shorthair breed</a>	274	1372	100%	9e-70	99.34%	<a href="#">AP023165.1</a>
<input checked="" type="checkbox"/>	<a href="#">PREDICTED: Marmota flaviventris 28S ribosomal RNA (LOC117794687)_rRNA</a>	274	274	100%	9e-70	99.34%	<a href="#">XR_004618536.1</a>
<input checked="" type="checkbox"/>	<a href="#">PREDICTED: Avicanthus niloticus 28S ribosomal RNA (LOC117704856)_rRNA</a>	274	274	100%	9e-70	99.34%	<a href="#">XR_004606369.1</a>
<input checked="" type="checkbox"/>	<a href="#">PREDICTED: Avicanthus niloticus 28S ribosomal RNA (LOC117704855)_rRNA</a>	274	274	100%	9e-70	99.34%	<a href="#">XR_004606368.1</a>

Description gn|SRA|SRR11085797.11550005.1 11550005 (Biological)

Molecule type dna

Query Length 151

Other reports [Distance tree of results](#) [MSA viewer](#) [?](#)

**Percent Identity**  to  **E value**  to  **Query Coverage**  to

**Descriptions** | Graphic Summary | Alignments

**Sequences producing significant alignments** Download Manage Columns Show 100 ?

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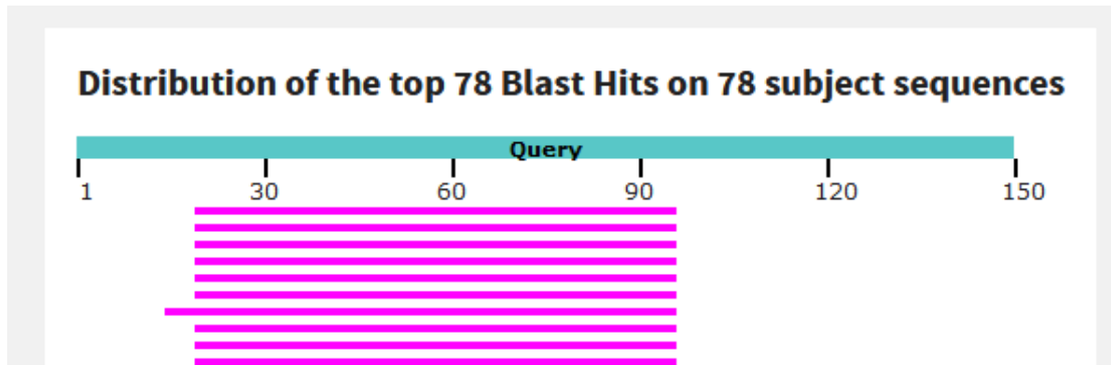
	Description	Max Score	Total Score	Query Cover	E value	Per. Ident	Accession
<input checked="" type="checkbox"/>	<a href="#">SRX7724752</a>	279	279	100%	2e-73	100.00%	<a href="#">SRA:SRR11085797.11601488.1</a>
<input checked="" type="checkbox"/>	<a href="#">SRX7724752</a>	279	279	100%	2e-73	100.00%	<a href="#">SRA:SRR11085797.11639592.1</a>
<input checked="" type="checkbox"/>	<a href="#">SRX7724752</a>	279	279	100%	2e-73	100.00%	<a href="#">SRA:SRR11085797.11598985.1</a>
<input checked="" type="checkbox"/>	<a href="#">SRX7724752</a>	279	279	100%	2e-73	100.00%	<a href="#">SRA:SRR11085797.11598757.1</a>
<input checked="" type="checkbox"/>	<a href="#">SRX7724752</a>	279	279	100%	2e-73	100.00%	<a href="#">SRA:SRR11085797.11588533.1</a>
<input checked="" type="checkbox"/>	<a href="#">SRX7724752</a>	279	279	100%	2e-73	100.00%	<a href="#">SRA:SRR11085797.11585765.1</a>
<input checked="" type="checkbox"/>	<a href="#">SRX7724752</a>	279	279	100%	2e-73	100.00%	<a href="#">SRA:SRR11085797.11585419.1</a>
<input checked="" type="checkbox"/>	<a href="#">SRX7724752</a>	279	279	100%	2e-73	100.00%	<a href="#">SRA:SRR11085797.11583582.1</a>
<input checked="" type="checkbox"/>	<a href="#">SRX7724752</a>	279	279	100%	2e-73	100.00%	<a href="#">SRA:SRR11085797.11582051.1</a>
<input checked="" type="checkbox"/>	<a href="#">SRX7724752</a>	279	279	100%	2e-73	100.00%	<a href="#">SRA:SRR11085797.11580259.1</a>
<input checked="" type="checkbox"/>	<a href="#">SRX7724752</a>	279	279	100%	2e-73	100.00%	<a href="#">SRA:SRR11085797.11579956.1</a>
<input checked="" type="checkbox"/>	<a href="#">SRX7724752</a>	279	279	100%	2e-73	100.00%	<a href="#">SRA:SRR11085797.11579609.1</a>
<input checked="" type="checkbox"/>	<a href="#">SRX7724752</a>	279	279	100%	2e-73	100.00%	<a href="#">SRA:SRR11085797.11573984.1</a>
<input checked="" type="checkbox"/>	<a href="#">SRX7724752</a>	279	279	100%	2e-73	100.00%	<a href="#">SRA:SRR11085797.11570618.1</a>
<input checked="" type="checkbox"/>	<a href="#">SRX7724752</a>	279	279	100%	2e-73	100.00%	<a href="#">SRA:SRR11085797.11568464.1</a>
<input checked="" type="checkbox"/>	<a href="#">SRX7724752</a>	279	279	100%	2e-73	100.00%	<a href="#">SRA:SRR11085797.11567362.1</a>
<input checked="" type="checkbox"/>	<a href="#">SRX7724752</a>	279	279	100%	2e-73	100.00%	<a href="#">SRA:SRR11085797.11566985.1</a>
<input checked="" type="checkbox"/>	<a href="#">SRX7724752</a>	279	279	100%	2e-73	100.00%	<a href="#">SRA:SRR11085797.11558788.1</a>



## Pale spear-nosed bat

Figure 9: *Phyllostomus Discolor*, or Pale spear-nosed bat, a species of bat native to the Americas, is found in high abundance within SRX7724752.

```
>gnl|SRA|SRR11085797.11550023.2 11550023 (Biological)
TTTGTTTTGTTTTTATAATTTATTTTTAAAATTTATTGGGGTGACAATTGTTAGTAAAA
TTACATAGATTTTCAGGTGTACAATTCCTGTATTACATGTGGACGGTCCAGCCGCCACGAGT
TCAACGTTTTACATGAAAGGGGGTGTGGGA
```



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### *Rhinolophus ferrumequinum* clone VMRC7-331J24, complete sequence

Sequence ID: [AC150242.3](#) Length: 120894 Number of Matches: 1

Range 1: 32707 to 32786 [GenBank](#) [Graphics](#)

[Next Match](#) [Previous Match](#)

Score	Expect	Identities	Gaps	Strand
122 bits(66)	3e-24	76/80(95%)	3/80(3%)	Plus/Plus
Query 20	tttattttttaa---at	tttattGGGGTGACA	ATTGTTAGTAAAA	TACATAGATTCAGG 76
Sbjct 32707	TTTTTTTTTAAATTT	ATTTATTGGGGTGACA	ATTGTTAGTAAAA	TACATAGATTCAGG 32766
Query 77	TGTACAATTCCTGT	TATTACAT 96		
Sbjct 32767	TGTACAATTCCTGT	TATTACAT 32786		

Figure 10: A sequence which was matched to a bat mRNA clone in the first 96 nucleotides, but then matching nothing on the later nucleotides. This match end with a T.



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### Bat coronavirus RaTG13, complete genome

Sequence ID: [MN996532.1](#) Length: 29855 Number of Matches: 2

Range 1: 26736 to 26844 [GenBank](#) [Graphics](#) [Next Match](#) [Previous Match](#)

Score	Expect	Identities	Gaps	Strand
161 bits(87)	7e-36	102/109(94%)	2/109(1%)	Plus/Plus
Query 45	GCAC	TGACTTG-CTT-TAGCACTGATGTGGCTGAGCTACTTCATTGCTTCTTTCAGGCTA	102	
Sbjct 26736	GCAATGGCTTGTCTTGTAGGCTTGATGTGGCTGAGCTACTTCATTGCTTCTTTCAGGCTA	26795		
Query 103	TTTGCACGTACGCGTTCCATGTGGTCATTCAATCCAGAACTAACATTT	151		
Sbjct 26796	TTTGCACGTACGCGTTCCATGTGGTCATTCAATCCAGAACTAACATTT	26844		

Range 2: 27478 to 27542 [GenBank](#) [Graphics](#) [Next Match](#) [Previous Match](#) [First Match](#)

Score	Expect	Identities	Gaps	Strand
121 bits(65)	1e-23	65/65(100%)	0/65(0%)	Plus/Plus
Query 1	ATGAAGGCAATTCACCATTCCATCCTCTAGCTGATAATAAATTTGCACTGACTTGCTTTA	60		
Sbjct 27478	ATGAAGGCAATTCACCATTCCATCCTCTAGCTGATAATAAATTTGCACTGACTTGCTTTA	27537		
Query 61	GCACT 65			
Sbjct 27538	GCACT 27542			

Figure 12: an anomalous fusion of two non-canonical regions of the RaTG13 genome. The fusion again happens on a T.

### Bat coronavirus RaTG13, complete genome

Sequence ID: [MN996532.1](#) Length: 29855 Number of Matches: 2

Range 1: 28217 to 28346 [GenBank](#) [Graphics](#) [Next Match](#) [Previous Match](#)

Score	Expect	Identities	Gaps	Strand
233 bits(126)	2e-57	129/130(99%)	1/130(0%)	Plus/Plus
Query 23	TTC-TCTAAACGAACAACTAAAATGTCTGATAATGGACCCCAAAACCAACGAAATGCAC	81		
Sbjct 28217	TTCATCTAAACGAACAACTAAAATGTCTGATAATGGACCCCAAAACCAACGAAATGCAC	28276		
Query 82	CCCGCATTACGTTTGGTGGACCCTCAGATTCAACTGGCAGTAACCAGAATGGAGAACGCA	141		
Sbjct 28277	CCCGCATTACGTTTGGTGGACCCTCAGATTCAACTGGCAGTAACCAGAATGGAGAACGCA	28336		
Query 142	GTGGAGCACG 151			
Sbjct 28337	GTGGAGCACG 28346			

Range 2: 25 to 60 [GenBank](#) [Graphics](#) [Next Match](#) [Previous Match](#) [First Match](#)

Score	Expect	Identities	Gaps	Strand
67.6 bits(36)	2e-07	36/36(100%)	0/36(0%)	Plus/Plus
Query 1	CTCTCGATCTCTTGTAGATCTGTCTCTAAACGAAC	36		
Sbjct 25	CTCTCGATCTCTTGTAGATCTGTCTCTAAACGAAC	60		

Figure 13: the only canonical sgRNA-like read\* in SRX7724752. Furthermore, SRX7724752 contained significant amount of reads that had higher query coverage on the DNA sequence than on the corresponding mRNA. This most likely indicate a clonal, rather than cDNA, library, was responsible for most of the bat-like reads observed in SRX7724752.

```

>gnl|SRA|SRR11085797.76.1 76 (Biological)
CATCAAACTGAGGTTTCAGCAAGGCAAAGATAGCCAGCAACAAAACAAAAAGGCATCCTA
CTGAATGGAAGCAGATAATTGCCAATAGTACATCAGTAAGGAGTTAATATTAAGAATTAG
TTTTTAAAAAGCTCTATATGATGTCAGAAAT

>gnl|SRA|SRR11085797.76.2 76 (Biological)
GTTTTTCACTTGCATTTCTCTAATAAATTAGTGATGTTGAGCATCTTTTCATATGTCATTTG
GCCATCTGTATGTCGTCTTTGGAGAAATGCTATTCAGATTTCTGCCCAATTTTAAATTG
GCTTGTTTTGTTTTTTGTTTTTGAAATTGAGTT

```

Descriptions    Graphic Summary    Alignments    Taxonomy

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select all 11 sequences selected    GenBank    Graphics    Distance tree of results

	Description	Max Score	Total Score	Query Cover	E value	Per. Ident	Accession
<input checked="" type="checkbox"/>	<a href="#">Rhinolophus ferrumequinum clone VMRC7-71A7 . complete sequence</a>	267	267	99%	1e-67	98.68%	<a href="#">AC150307.3</a>
<input checked="" type="checkbox"/>	<a href="#">Rhinolophus ferrumequinum clone VMRC7-251C10 . complete sequence</a>	185	185	94%	4e-43	90.14%	<a href="#">AC149630.3</a>
<input checked="" type="checkbox"/>	<a href="#">Myotis lucifugus clone CH235-427D16 . complete sequence</a>	137	137	82%	1e-28	87.20%	<a href="#">AC174832.3</a>
<input checked="" type="checkbox"/>	<a href="#">Pteropus alecto clone BAC.P100M20 .BAC.P103A18 . complete sequence</a>	135	135	86%	4e-28	85.38%	<a href="#">KP862827.1</a>
<input checked="" type="checkbox"/>	<a href="#">Pteropus alecto clone BAC.P201M3 .BAC.P216K21 . complete sequence</a>	135	135	86%	4e-28	85.38%	<a href="#">KP862826.1</a>
<input checked="" type="checkbox"/>	<a href="#">Pteropus alecto clone BAC.P56N20 . complete sequence</a>	135	135	86%	4e-28	85.38%	<a href="#">KP862825.1</a>
<input checked="" type="checkbox"/>	<a href="#">Rhinolophus euryale isolate REM0134 microsatellite RM1198 . sequence</a>	128	128	47%	7e-26	98.61%	<a href="#">KC910215.1</a>
<input checked="" type="checkbox"/>	<a href="#">Pteropus alecto clone BAC.P212O7-1 .BAC.P229M21 . complete sequence</a>	126	126	84%	3e-25	84.38%	<a href="#">KP862828.1</a>
<input checked="" type="checkbox"/>	<a href="#">PREDICTED: Miniopterus natalensis zinc finger protein 713 (ZNF713) . transcript variant X3 . mRNA</a>	106	106	90%	3e-19	81.02%	<a href="#">XM_016196283.1</a>
<input checked="" type="checkbox"/>	<a href="#">PREDICTED: Miniopterus natalensis zinc finger protein 713 (ZNF713) . transcript variant X2 . mRNA</a>	106	106	90%	3e-19	81.02%	<a href="#">XM_016196281.1</a>
<input checked="" type="checkbox"/>	<a href="#">PREDICTED: Miniopterus natalensis zinc finger protein 713 (ZNF713) . transcript variant X1 . mRNA</a>	106	106	90%	3e-19	81.02%	<a href="#">XM_016196280.1</a>

Figure 14: a read from SRX7724752 which have higher coverage on the clone than on the corresponding mRNA. E.g. the read contained nucleotide sequences that were not supposed to be transcribed in actual cells/bats.

## Inability of SRX8357956 to prove the promises claimed in [5]

Date	RaTG13		SARS-CoV-2		Sequence	Blast Archive	Name
	Identity	Query Cover	Identity	Query Cover			
14-Oct-18	95.53%	56%	93.15%	56%	24	archive.is/H107n	gnl SRA SRR11806578.24 RaTG13-9-5-5_9-5-f1_2018-10-14_B02
14-Oct-18	97.31%	76%	93.18%	75%	23	archive.is/8phs4	gnl SRA SRR11806578.23 RaTG13-9-5-4_9-5-r1_2018-10-14_C02
14-Oct-18	97.55%	53%	93.00%	52%	25	archive.is/ycQ89	gnl SRA SRR11806578.25 RaTG13-9-5-5_9-5-r1_2018-10-14_D02
14-Oct-18	99.43%	97%	96.39%	97%	22	archive.is/abSp6	gnl SRA SRR11806578.22 RaTG13-9-5-4_9-5-f1_2018-10-14_A02
11-Oct-18	97.37%	98%	92.23%	98%	20	archive.is/B20Et	gnl SRA SRR11806578.20 RaTG13-9-5-1_21230-F_2018-10-11_A12
11-Oct-18	98.54%	99%	88.26%	99%	21	archive.is/L2pTq	gnl SRA SRR11806578.21 RaTG13-9-5-1_23258-R_2018-10-11_B12
08-Oct-18	98.64%	98%	96.19%	99%	5	archive.is/W7Fxp	gnl SRA SRR11806578.5 RaTG13-11-2_18297-F_TSS20181008-027-0303_G10
08-Oct-18	99.19%	98%	91.83%	98%	7	archive.is/h8810	gnl SRA SRR11806578.7 RaTG13-12-2_24144-R_TSS20181008-027-0303_C11
08-Oct-18	99.89%	99%	87.86%	99%	6	archive.is/VJY2	gnl SRA SRR11806578.6 RaTG13-12-2_22717-F_TSS20181008-027-0303_H10
30-Sep-18	99.50%	99%	92.71%	99%	9	archive.is/ONBUX	gnl SRA SRR11806578.9 RaTG13-2-3_RaTG13-2-R1_2018-09-30_B11
30-Sep-18	99.79%	99%	92.89%	99%	8	archive.is/udSil	gnl SRA SRR11806578.8 RaTG13-2-3_RaTG13-2-F_2018-09-30_A02
29-Sep-18	99.00%	99%	94.20%	99%	10	archive.is/jdzvN	gnl SRA SRR11806578.10 RaTG13-2-3_RaTG13-2-R2_2018-09-29_D05
29-Sep-18	99.09%	98%	94.97%	98%	3	archive.is/lI9eI	gnl SRA SRR11806578.3 RaTG13-10-3_RaTG13-10-F_2018-09-29_G04
29-Sep-18	99.72%	98%	98.06%	98%	11	archive.is/7Kioa	gnl SRA SRR11806578.11 RaTG13-20-1_RaTG13-F_2018-09-29_H04
29-Sep-18	99.72%	98%	95.19%	98%	4	archive.is/tKTXg	gnl SRA SRR11806578.4 RaTG13-10-3_RaTG13-10-R_2018-09-29_E05
27-Sep-18	95.03%	98%	90.88%	98%	14	archive.is/NNfnm	gnl SRA SRR11806578.14 RaTG13-4-2_RaTG13-4-R_2018-09-27_G06
27-Sep-18	95.82%	98%	93.46%	93%	13	archive.is/udSil	gnl SRA SRR11806578.13 RaTG13-4-2_RaTG13-4-F_2018-09-27_G05
27-Sep-18	98.08%	98%	94.50%	98%	1	archive.is/kcHAi	gnl SRA SRR11806578.1 RaTG13-1-2_RaTG13-1-F_2018-09-27_E05
27-Sep-18	98.81%	99%	96.90%	99%	17	archive.is/nhvd2	gnl SRA SRR11806578.17 RaTG13-6-2_RaTG13-6-R_2018-09-27_H06
27-Sep-18	98.91%	99%	94.54%	99%	2	archive.is/veLPW	gnl SRA SRR11806578.2 RaTG13-1-2_RaTG13-1-R_2018-09-27_F06
27-Sep-18	99.09%	99%	96.92%	99%	16	archive.is/dtqMp	gnl SRA SRR11806578.16 RaTG13-6-2_RaTG13-6-F_2018-09-27_A06
27-Sep-18	99.28%	98%	96.81%	98%	12	archive.is/ZHjMY	gnl SRA SRR11806578.12 RaTG13-3-2_RaTG13-3-F_2018-09-27_F05
27-Sep-18	99.46%	98%	96.74%	98%	15	archive.is/Epig7	gnl SRA SRR11806578.15 RaTG13-5-2_RaTG13-5-F_2018-09-27_H05
27-Sep-18	99.50%	98%	98.49%	98%	18	archive.is/NdyHK	gnl SRA SRR11806578.18 RaTG13-7-2_RaTG13-7-F_2018-09-27_B06
27-Sep-18	99.53%	99%	95.67%	97%	19	archive.is/2qg0a	gnl SRA SRR11806578.19 RaTG13-8-2_RaTG13-8-F_2018-09-27_C06
20-Jun-17	99.10%	99%	96.61%	99%	28	archive.is/ve7nN	gnl SRA SRR11806578.28 RaTG13-R-1-1_7896-1-F1_2017-06-20_E03
20-Jun-17	99.61%	99%	97.43%	99%	32	archive.is/ehzBr	gnl SRA SRR11806578.32 RaTG13-R-4-1_7896-4-F_2017-06-20_F03
20-Jun-17	99.87%	98%	97.42%	98%	33	archive.is/do9Rt	gnl SRA SRR11806578.33 RaTG13-R-4-1_7896-4-R_2017-06-20_H03
20-Jun-17	99.90%	98%	97.44%	98%	29	archive.is/HjQD8	gnl SRA SRR11806578.29 RaTG13-R-1-1_7896-1-R1_2017-06-20_G03
17-Jun-17	98.56%	99%	95.85%	99%	26	archive.is/fqWWF	gnl SRA SRR11806578.26 RaTG13-ORF8-1-1_ORF8-F_2017-06-17_A05
17-Jun-17	98.99%	98%	96.52%	98%	27	archive.is/N01Ah	gnl SRA SRR11806578.27 RaTG13-ORF8-1-1_ORF8-R1_2017-06-17_A06
03-Jun-17	99.07%	97%	97.49%	97%	30	archive.is/WwyWy	gnl SRA SRR11806578.30 RaTG13-R-2-1_7896-2-F1_2017-06-03_A07
03-Jun-17	99.46%	99%	98.01%	99%	31	archive.is/tCLHu	gnl SRA SRR11806578.31 RaTG13-R-2-1_7896-2-R1_2017-06-03_A08

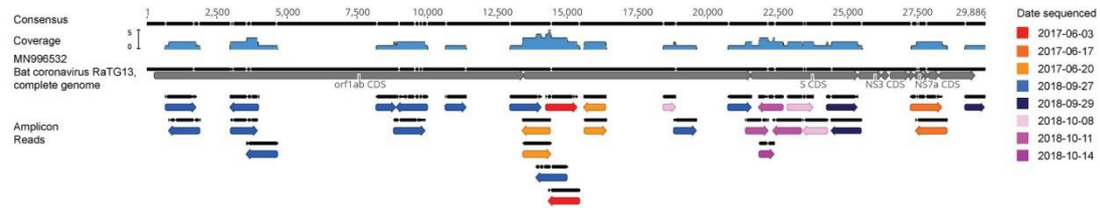


Figure 15: A complete analysis [4] of all Amplicon sequences in SRX8357956. Including the location of these amplicons and the similarity of such amplicon to the RaTG13 and SARS-CoV-2 genome.

Chuan Xiao et.al claimed that RaTG13 contained all the 3 S1 variable loops that were previously considered unique in SARS-CoV-2. [5] However, such claims can not be verified using the amplicons listed in SRX8357956.

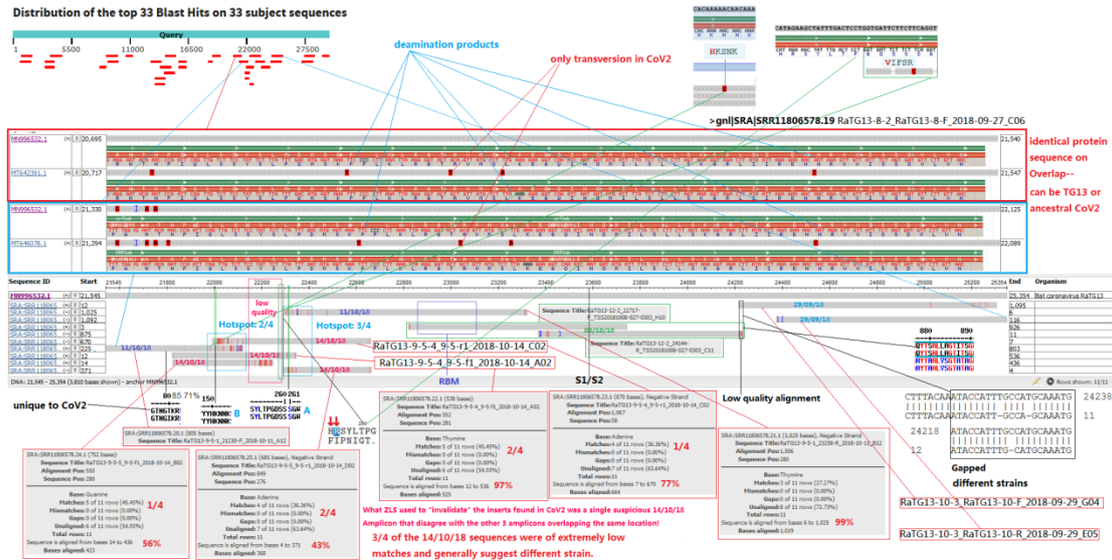


Figure 16: a thorough analysis of the amplicons located on the S locus of RaTG13 in SRX8357956. Notice that the last 4 amplicons sequenced in 14/10/2018 was of very low quality matches, and matched other organisms—including Mouse(mus musculus).

select all 99 sequences selected

Description	Max Score	Total Score	Query Cover	E value	Per. Ident	Accession
<input checked="" type="checkbox"/> Gadus morhua genome assembly_chromosome: 16	60.8	105	27%	4e-05	78.05%	LR633958.1
<input checked="" type="checkbox"/> Mus musculus BAC clone RP24-498P8 from chromosome 9_complete sequence	59.9	59.9	47%	1e-04	72.92%	AC168217.2
<input checked="" type="checkbox"/> Mus musculus chromosome 9_clone RP24-484G16_complete sequence	59.9	59.9	47%	1e-04	72.92%	AC137678.11

select all 100 sequences selected

Description	Max Score	Total Score	Query Cover	E value	Per. Ident	Accession
<input checked="" type="checkbox"/> Homo sapiens BAC clone RP11-792A8 from 7_complete sequence	54.5	54.5	44%	0.006	71.15%	AC027644.9
<input checked="" type="checkbox"/> Coregonus sp. 'balchen' genome assembly_chromosome: 7	52.7	52.7	29%	0.022	73.68%	LR778259.1
<input checked="" type="checkbox"/> Coregonus sp. 'balchen' genome assembly_chromosome: 15	50.9	50.9	16%	0.078	83.33%	LR778267.1
<input checked="" type="checkbox"/> Salmo trutta genome assembly_chromosome: 21	50.9	50.9	37%	0.078	71.90%	LR584437.1
<input checked="" type="checkbox"/> Xanthophylomyces dendrorhous genome assembly_Xden1_scaffold Scaffold_79	50.9	50.9	9%	0.078	96.88%	LN483167.1
<input checked="" type="checkbox"/> Coregonus sp. 'balchen' genome assembly_chromosome: 20	50.0	50.0	72%	0.078	67.49%	LR778272.1
<input checked="" type="checkbox"/> Aquila chrysaetos chrysaetos genome assembly_chromosome: 14	50.0	141	42%	0.078	69.50%	LR606194.1
<input checked="" type="checkbox"/> Bos mutus isolate yakQH1 chromosome 16	50.0	50.0	24%	0.078	74.68%	CP027084.1
<input checked="" type="checkbox"/> Mus musculus BAC clone RP23-128D11 from 7_complete sequence	50.0	50.0	36%	0.078	71.90%	AC122222.6
<input checked="" type="checkbox"/> Mus musculus BAC clone RP23-66E21 from 7_complete sequence	50.0	50.0	36%	0.078	71.90%	AC131741.4

select all 24 sequences selected

Description	Max Score	Total Score	Query Cover	E value	Per. Ident	Accession
<input checked="" type="checkbox"/> Mus musculus targeted KO-first_conditional ready_lacZ-tagged mutant allele Fabp4.tm1a(KOMP)Wts	50.9	50.9	23%	0.045	85.11%	JN963014.1
<input checked="" type="checkbox"/> Mus musculus targeted non-conditional_lacZ-tagged mutant allele Fabp4.tm1e(KOMP)Wtsj_transger	50.9	50.9	23%	0.045	85.11%	JN947213.1
<input checked="" type="checkbox"/> Mus musculus chromosome 3_clone RP23-436F15_complete sequence	50.9	50.9	23%	0.045	85.11%	AC123726.11
<input checked="" type="checkbox"/> Mus musculus chromosome 3_clone RP24-137C19_complete sequence	50.9	50.9	23%	0.045	85.11%	AC113990.10

Figure 16: BLAST result of the non-RaTG13 matched parts of Amplicons 25, 24 and 23 in SRX8357956. Using the remaining amplicons, the 3 variable loops, GTNGIKR, HKSNNK and VIFSQ was obtained.



The second overlap, one with BtCoV/4991, contained only 1 C-T transition, which have a probable origin in the primers used to generate the amplicons in the first place.

## DISCUSSIONS

### Origins of the anomalies in SRX7724752

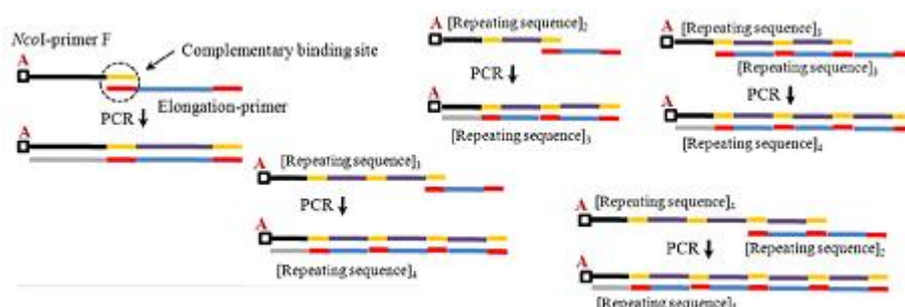


Figure 18A: Origin of repeating sequences in SRX7724752.

The only satiable explanation for the anomalous enrichment of the Telomere-like repeats in SRX7724752, involves the self-amplification of such sequences in a PCR reaction with little to no template.

Normally, with significant amount of template, the random primers normally used in RT-PCR amplifies most sequences evenly and outcompetes the repeat sequences in the reaction, and the result was an accurate reflection of such repeats within cellular samples—extremely poor. However, in samples that have little to no template, such that the random primers/random hexamers used in the reaction were not able to prime the amplification of most sequences—e.g. the amount of normal templates within the reaction falls below the timescale needed for the amplification of the repeating sequences, Repeating sequences, of which telomere-like repeats forms the vast majority of it in the environment and in most samples, can self-amplify in a primer-independent fashion, eventually reaching very high dominance, through repeated denaturing, sliding, reannealing and extension.

As this is a linear process, the self-amplification process is very slow, and is normally outcompeted by the normal amplicons as long as any usable amount of templates were present. Therefore, the presence of anomalously enriched telomere-like repeats within SRX7724752 indicate that the original sample couldn't have contained enough templates for the generation of the complete genome, through any means possible.

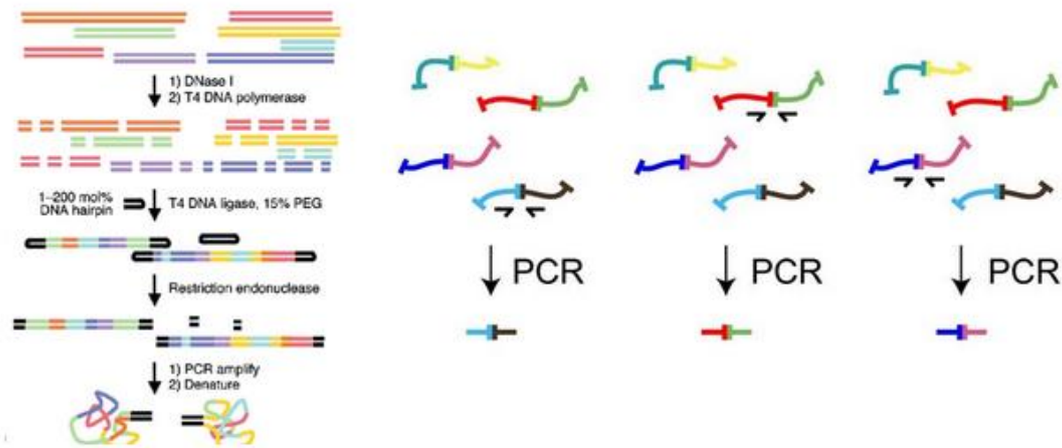


Figure 18B: Origin of the random matched sequences and partial sequences in SRX7724752  
 The majority of the sequences that were not repeats, when BLASTed, does not match any known organisms. There were also many sequences that matches—only partially, to many diverse organisms. What was striking, is, however, is that these matches often ends with an “A” or a “T”. The most possible explanation of this anomaly is that the Library preparation process of ILLUMINA RNA-seq, which include strand synthesis and A-T ligation to adaptors, were fed dsDNA rather than ssRNA, as input. Such dsDNA input may be PCR products, or it may be a pre-made ILLUMINA sequencing library—Certain mRNA-like reads were inadvertently inverted, indicating double-stranded cDNA was likely used instead of single stranded mRNA.

<b>RID</b>	<a href="#">J85DMESK016</a> Search expires on 08-01 20:59 pm <a href="#">Download All</a> ▼
<b>Results for</b>	2:lcl Query_5937 gnl SRA SRR11085797.66.2 66 (Biological)(150bp) ▼
<b>Program</b>	BLASTN <a href="#">?</a> <a href="#">Citation</a> ▼
<b>Database</b>	nt <a href="#">See details</a> ▼
<b>Query ID</b>	lcl Query_5937
<b>Description</b>	gnl SRA SRR11085797.66.2 66 (Biological)
<b>Molecule type</b>	dna
<b>Query Length</b>	150
<b>Other reports</b>	<a href="#">Distance tree of results</a> <a href="#">?</a>

Download ▾ GenBank Graphics

**PREDICTED: Hipposideros armiger putative P2Y purinoceptor 10 (LOC109385656), mRNA**

Sequence ID: [XM\\_019648164.1](#) Length: 2682 Number of Matches: 1

Range 1: 2162 to 2293 [GenBank](#) [Graphics](#) [Next Match](#) [Previous Match](#)

Score	Expect	Identities	Gaps	Strand
154 bits(83)	1e-33	116/132(88%)	1/132(0%)	Plus/Minus
Query 12	TTT-TCATTATTAAGTATTATGTACTGTACATAAATGTATGTACTATACTTTTATACAAC	70		
Sbjct 2293	TTTATCATATATCAAGTGTATGTACTGTACAGTATTGTATGTGTATATACTTTTATATGAC	2234		
Query 71	TGGCAGCACAGCAGGTTTGGTTTATACCAGCATCACCACAAAAATGTGAGTAATGCATTAC	130		
Sbjct 2233	TGACAGCATAGTAGGCTTGGTTTACACCAGCATCACCACAAAAATGTGAGTAATGCATTAC	2174		
Query 131	ACTACAATGTTA	142		
Sbjct 2173	ACTATGATGTTA	2162		

Figure 19: An inverted mRNA-like read.

\*: Analysis of the sole sgRNA-like read reveal the usage of a leader/F primer and the mispriming of Amplicon DNA

```

          ttagatttcattctaaacgaacaaactaaaatgtctgataatggacccccaaacgaatgcaccccgattacgtttgggtgacccct
          CTCTCGATCTCTGTAGATCTGTTCTCTAAACGAAC
ACA AACCAACGAACTCTCGATCTCTGTAGATCTGTTCTCTAAACGAAC
          TAACTCCTTTTIGCCCTAGTTTACCGATCTCATCGCCCTGCCAGGGTCCATGGACTGTGTGATCTGCTCTCTGCTCCTC

```

Figure 20a: the match analysis between different genomic fragments of RaTG13, of the sole sgRNS-like read\* in SRX7724752. \*:Figure 13

Despite being sgRNA-like in the first glance, analysis of the exact overlapping region of this particular read reveal that this region is identical to BtCoV/ZC45 and BtCoV/ZXC21—indicating it's identity as likely being a consensus primer.

select all 100 sequences selected [GenBank](#) [Graphics](#) [Distance tree of results](#)

Description	Max Score	Total Score	Query Cover	E value	Per. Ident	Accession
<input checked="" type="checkbox"/> <a href="#">Pangolin coronavirus isolate PCoV_GX-P3B_genomic_sequence</a>	67.6	67.6	100%	8e-09	100.00%	<a href="#">MT072865.1</a>
<input checked="" type="checkbox"/> <a href="#">Pangolin coronavirus isolate PCoV_GX-P2V_complete_genome</a>	67.6	67.6	100%	8e-09	100.00%	<a href="#">MT072864.1</a>
<input checked="" type="checkbox"/> <a href="#">Pangolin coronavirus isolate PCoV_GX-P5E_complete_genome</a>	67.6	67.6	100%	8e-09	100.00%	<a href="#">MT040336.1</a>
<input checked="" type="checkbox"/> <a href="#">Pangolin coronavirus isolate PCoV_GX-P5L_complete_genome</a>	67.6	67.6	100%	8e-09	100.00%	<a href="#">MT040335.1</a>
<input checked="" type="checkbox"/> <a href="#">Pangolin coronavirus isolate PCoV_GX-P1E_complete_genome</a>	67.6	67.6	100%	8e-09	100.00%	<a href="#">MT040334.1</a>
<input checked="" type="checkbox"/> <a href="#">Pangolin coronavirus isolate PCoV_GX-P4L_complete_genome</a>	67.6	67.6	100%	8e-09	100.00%	<a href="#">MT040333.1</a>
<input checked="" type="checkbox"/> <a href="#">Bat coronavirus RaTG13_complete_genome</a>	67.6	67.6	100%	8e-09	100.00%	<a href="#">MN996532.1</a>
<input checked="" type="checkbox"/> <a href="#">Bat SARS-like coronavirus isolate bat-SL-CoVZXC21_complete_genome</a>	67.6	67.6	100%	8e-09	100.00%	<a href="#">MG772934.1</a>
<input checked="" type="checkbox"/> <a href="#">Bat SARS-like coronavirus isolate bat-SL-CoVZC45_complete_genome</a>	67.6	67.6	100%	8e-09	100.00%	<a href="#">MG772933.1</a>
<input checked="" type="checkbox"/> <a href="#">Bat coronavirus (BtCoV/279/2005)_complete_genome</a>	65.8	65.8	97%	3e-08	100.00%	<a href="#">DQ648857.1</a>
<input checked="" type="checkbox"/> <a href="#">Mutant SARS coronavirus Urbani clone SARS-Urbani-MA_SHC014-spike_complete_genome</a>	63.9	63.9	94%	1e-07	100.00%	<a href="#">MT308984.1</a>
<input checked="" type="checkbox"/> <a href="#">Coronavirus BtRs-BetaCoV/YN2018D_complete_genome</a>	63.9	63.9	94%	1e-07	100.00%	<a href="#">MK211378.1</a>

Figure 20b: BLAST result of CTCTCGATCTCTGTAGATCTGTTCTCTAAACGAAC.

This particular sequence have extended overlap to the beginning of the N gene, which was coincidentally at the end of the last 2017/06/17 amplicon. This indicate it was most likely the product of mispriming, rather than a true sgRNA-like read.

**SRX7724752 is a mixed library consists of a matrix of dried American bat guano, a bat WGS/RNA-seq ILLUMINA library, a synthetic 16S library and megaprimer PCR products from the SRX8357956 Amplicons and a degraded sample of SARS-CoV-2 cDNA**

Phyllostomus Discolor, a species of bat native to Mexico and southern United states, leaves numerous Full-length 100% matched reads that don't match anything else. Coincidentally, Mexico is one of the major supplier of bat guano used for fertilizer and other commercial purposes[6]. The confirmed presence of this particular bat species, suggest the use of a commercial dried bat guano matrix as the bulk of the sample being sequenced. As in PRJNA494391[7] which synthetic metagenome samples were constructed using cDNA amplicons and a specific material matrix to simulate realistic metagenomic reads of a desired virus in a sample.

Traces of the original template used in the megaprimer PCR process can be seen as traces of low-matched virus-like reads within this dataset, which are found across the entire RaTG13 genome.



Figure 21: Read coverage of SRX7724752 on the RaTG13 genome. The red pixels represent significant mismatches on the reads in the dataset.

The Bacterial-like reads in SRX7724752 is also likely a synthetic 16S library—as the only other dataset with Telomere-like repeats(4%), still contained significant amount of bacterial mRNA.

```
>gnl|SRA|SRR11085733.2232944.1 2232944 (Biological)
GCCTTCGTTTGATATAGTTTTAATGCAAATCCCCTAACATCTCTTTCAGCATCTGCTGC
ACCTCTTTCACCAGCAACTGTAGAAAATCTTAAAAGGGCTTTTGTTTTTTTACCAACTTT
GTTAAAAATATCTGCTTTAGAATATTTTGT

>gnl|SRA|SRR11085733.2232944.2 2232944 (Biological)
AGAGGTCCACTCTTTTACAAGATACTTGGCTTTTAGAAAACTTGCACATTTTCGATAGG
GAAAGGATACCAGAAAGAGTTGTGCACGCTAAAGGAAGTGCTGCATACGGCGAATTAACA
ATTACTAATGATATTACAAAATATTCATAA
```



<b>Job Title</b>	<b>AE017125:Helicobacter hepaticus ATCC 51449,...</b>
<b>RID</b>	<a href="#">J88SF43U01R</a> Search expires on 08-01 21:56 pm <a href="#">Download All</a> ▼
<b>Program</b>	<a href="#">?</a> <a href="#">Citation</a> ▼
<b>Database</b>	SRA <a href="#">See details</a> ▼
<b>Query ID</b>	<a href="#">AE017125.1</a>
<b>Description</b>	Helicobacter hepaticus ATCC 51449, complete genome
<b>Molecule type</b>	nucleic acid
<b>Query Length</b>	934935
<b>Other reports</b>	<a href="#">?</a>



No significant similarity found. For reasons why, [click here](#)

Figure 22b: the same species of bacteria in SRX7724752. No significant matches were found. This dataset is likely subjected to probe-capture sequencing similar to these other datasets—the use of a positive-sense CoV probe resulted in the selective presentation of the negative ssDNA strand of the ligation products to show up. This is supported by the observation that while most of the virus-like reads were on the negative strand, the Repeats does not show a bias in strand polarity, and the mRNA-like reads have a much higher chance of being on the wrong polarity for RNA-seq. This is likely due to the ligation process being used.

## CONCLUSION

The raw data of BtCoV/RaTG13 Contained multiple anomalies that signifies that the original sample could not have contained enough RNA template for the extraction of a complete viral genome as in MN996532.1

Furthermore, many of these anomalies points toward the fraudulent use of a mixed DNA library, rather than genuine mRNA, for the sequencing of SRX7724752, evident by the presence of widespread A-T ligation of unrelated dsDNA fragments that can only happen if the same library preparation process have been ran on dsDNA instead of ssRNA. which would constitute Academic fraud.

Therefore, the sequencing of BtCoV/RaTG13 can not be considered to be valid or honest as is, and any publications, including [2], and other publications that cites or use RaTG13 as critical pieces of evidence or proof, must be immediately invalidated and retracted.

## REFERENCES

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