1	Short	Note
1	Short	Note

- 2 The Abnormal Nature of the Fecal Swab Sample used for
- **3 NGS Analysis of RaTG13 Genome Sequence Imposes a**
- 4 Question on the Correctness of the RaTG13 Sequence
- 5

6 Monali C. Rahalkar¹* and Rahul A. Bahulikar²

- ⁷ ¹C2, Bioenergy group, MACS Agharkar Research Institute, G.G. Agarkar Road,
- 8 Pune 411004, Maharashtra, India
- 9 ²BAIF Development Research Foundation, Central Research Station,
- 10 Urulikanchan, Pune 412202
- 11 *Corresponding author: <u>monalirahalkar@aripune.org</u>
- 12

13

- 14
- 15

 \odot \odot

1 Abstract:

RaTG13 is the next relative of SARS-CoV-2 derived from bat feces. The Illumina based 2 NGS sequence of RaTG13 MN996532.1 was deposited on 27th Jan 2020 and the raw data, a 3 little later on 13th Feb 2020 https://www.ncbi.nlm.nih.gov/sra/SRX7724752[accn]. The fecal 4 swab sample shows abnormally high reads from eukaryotes which includes not only bats but 5 6 other animals, as per the NCBI site. Also, comparison of the fecal swab to other bat fecal 7 swabs deposited by the same group on the same date indicates that the fecal swab from which 8 RaTG13 sequence was derived looked abnormal. The proportion of bacteria in this RNA Seq project was only 0.7% in contrast to 70-90% abundance in other fecal swabs from bats. Also, 9 the amplicon sequencing done on the same sample showed large number of gaps and 10 inconsistencies. This poses a question on the authenticity of the RaTG13 sequence also. 11 Keywords: RaTG13; SARS-COV-2; Illumina sequencing; amplicon sequencing; NGS; fecal 12

13 swab

14

15

1 Covid-19 has been a devastating pandemic affecting more than nineteen million people in more than 200 countries and killing three quarter million people till now. SARS-CoV2, the 2 virus responsible for the disease is most similar to RaTG13 (a bat derived virus) on the 3 4 genomic level. RaTG13 has been known as the sister virus of SARS-CoV-2 as its shows 96.2% overall genomic similarity to CoV-2 genome (Zhou et al., 2020). RaTG13 has been 5 6 widely used for various comparative experiments with that of SARS-CoV-2. This includes the capacity of its spike to bind to human ACE-2, its infective capacity, etc. RaTG13 genome 7 is also used for calculations of the common ancestor and also for further calculations before 8 9 how long RaTG13 and SARS-CoV-2 got separated, etc.

RaTG13 is described as the virus (not a real virus, but available as a sequence) from the RNA of a bat fecal swab collected in July 2013, from Tongguan mines in Yunnan. The old name of RaTG13 virus is CoV4991 (Ge et al., 2016). However, the sample appears to be over or not available to the scientific community as per a recent news investigation (2020). One main condition for using RaTG13 for all future experiments is that the sequence of this virus should be accurate and based on a good raw data.

RaTG13 never seemed to have existed before SARS-COV-2 was described, as the genome
sequence was not available on NCBI before (Zhou et al., 2020) .The Illumina based NGS
sequence of RaTG13 MN996532.1 was deposited on 27th Jan 2020 and the raw data, a little
later on 13th Feb 2020 <u>https://www.ncbi.nlm.nih.gov/sra/SRX7724752[accn]</u>.

The earlier name of RaTG13 is CoV/4991. A 370 base RdRp fragment (KP378696.1) of CoV/4991 and showed highest similarity to SARS-CoV-2 RdRp fragment with only 3-5 bases different (NCBI blast analysis). Also, 4991 or RaTG13 has a great significance as it was recovered from the same site where a COVID-19 like disease occurred (2020, Rahalkar

1	and Bahulikar,	2020). Co	V 4991 i	s also the	first and	only	SARS-like CoV	associated	with
---	----------------	-----------	----------	------------	-----------	------	---------------	------------	------

2 human pneumonia cases, before SARS-COV-2 (Rahalkar and Bahulikar, 2020).

3 Problems seen in the RAW DATA of RaTG13: Illumina sequence SRX7724752

4 Here are the basic discrepancies encountered after the analysis of the Illumina raw data

5 <u>https://www.ncbi.nlm.nih.gov/sra/SRX7724752[accn]</u>:

1. The genome of RaTG13 is derived from a fecal or anal swab (MN996532.1). However in
the Illumina sequencing description, SRX7724752, the sample is described to be of a BAL
fluid (broncho alveolar lavage).

9 2. The total raw data is 3.3 Gb. After the Krona analysis it is seen that ~30% reads are
10 unidentified (no matches) and only ~ 70% reads are identified. Out of the 70%, a vast
11 majority i.e. 68% was contributed by eukaryotes (fig. 1). This is highly unusual as it is a fecal
12 swab and the analysis of other bat fecal or anal swabs cannot show such high proportion of
13 eukaryotic RNA.

3. Within the 68% eukaryote sequences, the bat sequences are about 36-40% (Fig 1a.), and rest of the 30% sequences are contributed by squirrels, flying foxes, foxes, and other types of animals (Fig.1 b). First of all, why would such high proportion of eukaryotic sequences appear in the RNA when it's a fecal swab? From where do these animal sequences come when it is supposed to be a *Rhinophus affinis* swab? Also, even though the *Rhinophus affinis* sequence may not be present in the database, why are they similar to so many bat sequences? Some of these bats are found only in Mexico or USA (Zhang, 2020).

4. The RNA Seq data shows extremely less abundance of bacteria, only 0.65%. This is far too
less in comparison to other fecal or anal swab of bats, which show a very high proportion of
bacterial sequences ~76-90% (Fig.2 and.3). SRA data of six other fecal swabs submitted by

the same group were used for comparison (data not shown). Bacteria are the highest
 constituents of a fecal sample.

5. The coronavirus sequence (RaTG13) contributes to only ~0.003% of the total sequence reads. These raw reads were used to build an almost complete assembly, though the overall coverage is very less ~8X. Though there were less overlaps in some regions there are only 2-3 gaps. The Wuhan Institute of Virology has recently described methods like probe-capture for getting the whole genome of viruses from samples like bat feces (Li et al 2019). In this case, without the use of any other methods, and after using so old fecal swab or fecal swab RNA with no bacteria in it, how did they recover such good quality viral reads?

6. The assembly method and the actual assembly accession for RaTG13 is not described or
linked to MN669532 and also no assembly method is specified in the raw data SRX7724752
and the Illumina run. Therefore, no assembly data is available for RaTG13 genome.

7. After blasting the RaTG13 genome against the SRA, ~1700 reads can be retrieved which
covers only 252 Kb of the total 3.3 Gb. The genome size of RaTG13 is known to be ~30 kb.
Therefore this is ~8x coverage, which is quite less and insufficient to arrive to a definitive
assembly. Then how was the sequence MN669532 used so confidently by various researchers
without any doubt?

8. We also compared the fecal/anal swab from the same species, i.e. *Rhinolophus affinis*(Fig.2) and fecal swab from another bat (Fig. 3) and it clearly shows that the other two swabs
showed normal findings, with 70-90% bacterial reads and very few reads associated with the
host. Also these swabs do not show sequences coming from other animals.

9. Similar findings have been documented in a latest preprint by Zhang, D. (Zhang,
 2020) <u>https://zenodo.org/record/3969272#.Xypwfn5S-Un.</u>

1 Problems in the Amplicon sequencing data:

We found that some amplicon sequencing data for RaTG13 (SRX8357956) was submitted in
May 2020.

4 1. No indications of amplicon sequencing given by Zhou et al 2020 about the amplicon
5 sequencing of RaTG13. There are in total 33 spots with forward and reverse sequences.

2. This sequencing shows that the dates are 2017 and 2018. However, the submission has
been done in 2020. This sequencing has never been mentioned in any publications. Also, it
does not cover the entire genome and major gaps are seen in various regions.

9 3. There are two contrasting sequences for a single patch (spots 23 and spot 24), e.g. shows
94-96% similarity to that of MN669532.1. However, another spot the same sequence showed
99% similarity to the described RaTG13 consensus MN669532.1.

4. In general, the amplicons show 97-99% similarity with the MN669532.1. However, it doesnot cover the entire genome and major gaps are seen in various regions.

5. Also the RdRp derived from the amplicon sequencing is incomplete and does not match
with RdRp of 4991 KP876546.1. Around 170 bases from 370 base sequences are missing and
it shows 2 base mismatches.

17 **Conclusions:**

a. Our main objection is that the fecal swab from which RaTG13 sequence
is derived does not appear like a normal fecal sample due to the above
listed things.

b. RaTG13 sequence has been used extensively for all genomic comparisons
as it is believed to be the next relative of SARS-CoV-2.

c. However, the nature of the fecal swab appears very suspicious, with 70%
 of eukaryotic sequences also from sources which should not have been
 detected in bat feces like mexican bats, squirrels, flying foxes, red foxes,
 etc.).

d. And most importantly, there is negligible abundance of bacteria.
Bacteria constitute a major part of any feces, irrespective if it is an animal
or bird or any eukaryote.

8 e. The reads from which the viral sequence of RaTG13 was derived 9 appears not to be affected. An almost complete assembly is assumed to be 10 had been built from this raw data (Illumina reads). How did so good data 11 come from an otherwise abnormal looking, old and degraded fecal swab 12 sample preserved for 7-8 years?

f. The amplicon data is incomplete and submitted much later and
undescribed anywhere.

g. The question is why are these anomalies? And if these are there, should
the scientific community really rely on the RaTG13 genome sequence
MN996532.1? Should this data be used for further important experiments?

18

19

1 Figures:

2 Fig.1 RNA-Seq of Rhinolophus affinis:Fecal swabTaxonomy Analysis (RaTG13)

Full 🗸					Send to
<u>SRX7724752</u> : RN 1 ILLUMINA (Illum				s, 1.7Gb downlo	ads
-	nen constructed	d using the TruSe	q Stranded m	RNA Library Pr	Aamp Viral RNA Mini Kit following the manufacturers instructions. eparation Kit (Illumina, USA). Paired-end (150 bp) sequencing of th
Submitted by: W	uhan Institute o	of Virology, Chines	se Academy	of Sciences	
Study: Bat corona PRJNA60616 show Abstrac	5 • <u>SRP249482</u>	Genome sequenci · <u>All experiments</u>	•		
	201 • SRS61468 identified coron	537 • <u>All experime</u> avirus	ents • <u>All run</u> s	2	
Library: Name: RaTG Instrument: II Strategy: RNJ Source: MET Selection: RA Layout: PAIR	lumina HiSeq 3 A-Seq AGENOMIC NDOM	000			
Runs: 1 run, 11.6	M spots, 3.3G k	oases, <u>1.7Gb</u>			
Run	# of Spots	# of Bases	Size	Published	
		3.3G	4 701	2020-02-13	

4 Fig1a. RNA-Seq of *Rhinolophus affinis*:Fecal swab (RaTG13)

5

RNA-Seq of Rhinolophus affinis:Fecal swab (SRR11085797)

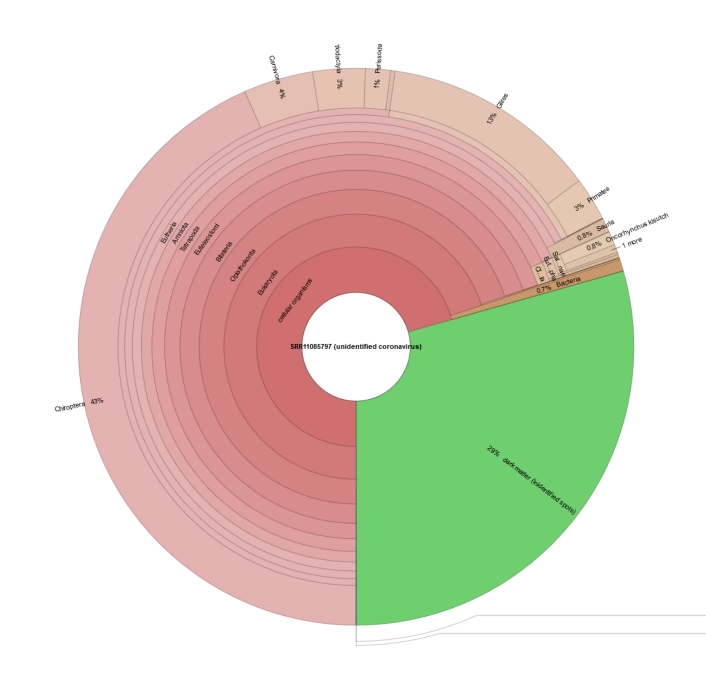
axonomy An	alysis				
Unidentified rea	ds: 29.38%				
Identified reads	: 70.62%				
ellular organ	isms: 70.61%				
Eukaryota:					
	onta: 49.7% a: 49.23%				
	a: 49.23% eria: 48.9%				
	eleostomi: 41.62%				
	mniota: 14.99%				
Ê Î	Eutheria: 11.52%				
	Boreceutheria: 10.81% Laurasiatheria: 6.61%				
	Chiroptera: 4.27%				
	Euarchontoglires: 1.91%				
	< 0.01% (7 Kbp)				
	ntae: 0.09%)1% (10 Kbp)				
Bacteria: 0.	65%				
 Bacteria: 0. Viruses: 0.019 					
Viruses: 0.019	6				
	6				
Viruses: 0.019	6				
Viruses: 0.019	6	Rank	%%	Кbp	Coverage
Viruses: 0.019	6	Rank species		Kbp 1,048,945	Coverage
Viruses: 0.019 View in Krona Strong signals SuperKingdom	Organism				Coverage
Viruses: 0.019 View in Krona Strong signals SuperKingdom Eukaryota	6 Organism Hipposideros armiger	species	31.8 4.6	1,048,945	Coverage
Viruses: 0.019 View in Krona Strong signals SuperKingdom Eukaryota Eukaryota	6 Organism Hipposideros armiger Rousettus aegyptiacus	species species	31.8 4.6	1,048,945 151,010	Coverage
Viruses: 0.019 View in Krona Strong signals SuperKingdom Eukaryota Eukaryota Eukaryota	6 Organism Hipposideros armiger Rousettus aegyptiacus Marmota marmota marmota	species species subspecies	31.8 4.6 4.6	1,048,945 151,010 150,069	Coverage
Viruses: 0.019 View in Krona Strong signals SuperKingdom Eukaryota Eukaryota Eukaryota Eukaryota	6 Organism Hipposideros armiger Rousettus aegyptiacus Marmota marmota marmota Vulpes vulpes	species species subspecies species	31.8 4.6 4.6 4.0	1,048,945 151,010 150,069 131,805	Coverage
Viruses: 0.019 View in Krona Strong signals SuperKingdom Eukaryota Eukaryota Eukaryota Eukaryota Eukaryota	6 Organism Hipposideros armiger Rousettus aegyptiacus Marmota marmota marmota Vulpes vulpes Marmota flaviventris	species species subspecies species species	31.8 4.6 4.6 4.0 3.6	1,048,945 151,010 150,069 131,805 118,361	Coverage
Viruses: 0.019 View in Krona Strong signals SuperKingdom Eukaryota Eukaryota Eukaryota Eukaryota Eukaryota Eukaryota	6 Organism Hipposideros armiger Rousettus aegyptiacus Marmota marmota marmota Vulpes vulpes Marmota flaviventris Pteropus	species species subspecies species species genus	31.8 4.6 4.6 3.6 3.0	1,048,945 151,010 150,069 131,805 118,361 100,495	Coverage

1 2

3

Fig. 1b. Distribution of the reads in the raw data. The individual distribution is given and in thesecond part, the reads which contribute to a higher extent are given.

6



2

- 3 Fig.1 c. Krona chart of RaTG13 raw data, 29% unidentified reads, 43% Chiroptera, 13% Gileres, 3%
- 4 Primates, 0.7% bacteria and 0.024% RaTG13 reads
- 5

Fig 2. RNA-Seq of Rhinolophus affinis: Fecal swab Taxonomy Analysis 1

2 https://www.ncbi.nlm.nih.gov/sra/SRX7724693[accn]

Full 🗸 SRX7724693: RNA-Seq of Rhinolophus affinis: Anal swab 1 ILLUMINA (Illumina HiSeq 3000) run: 11.9M spots, 3.5G bases, 1.6Gb downloads Design: Total RNA was extracted from bronchoalveolar lavage fluid using the QIAamp Viral RNA Mini Kit following the manufacturers instructions. An RNA library was then constructed using the TruSeq Stranded mRNA Library Preparation Kit (Illumina, USA). Paired-end (150 bp) sequencing of the RNA library was performed on the HiSeq 3000 platform (Illumina). Submitted by: Wuhan Institute of Virology, Chinese Academy of Sciences Study: Discovery of Bat Coronaviruses through Surveillance and Probe Capture-Based Next-Generation Sequencing. PRJNA606159 • SRP249478 • All experiments • All runs show Abstract Sample: SAMN14086235 • SRS6146479 • All experiments • All runs Organism: unclassified Rhinacovirus Library: Name: 160660 Instrument: Illumina HiSeq 3000 Strategy: RNA-Seq Source: METAGENOMIC Selection: RANDOM Layout: PAIRED Runs: 1 run, 11.9M spots, 3.5G bases, 1.6Gb Run # of Spots # of Bases Size Published SRR11085736 11,924,182 3.5G 1.6Gb 2020-02-13

ID: 10102706

3

Fig. 2a. RNA-Seq of Rhinolophus affinis: Anal swab (SRR11085736) 4

5

Send to: -

Taxonomy Analysis

Unidentified reads: 0.86%

Identified reads: 99.14% cellular organisms: 99.11% Bacteria: 91.07% Eukaryota: 4.36% Viruses: 0.03%

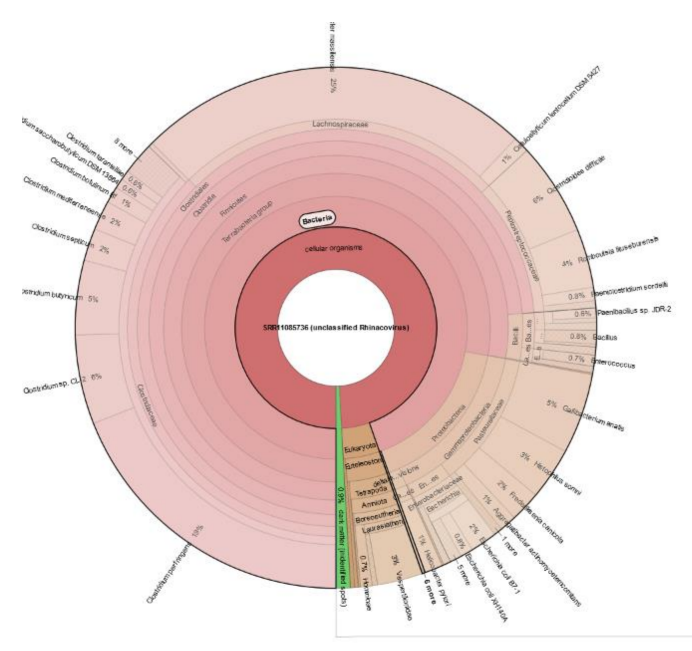
🕼 <u>View in Krona</u>

BacteriaClostridiumgenus37.31,288,845BacteriaNiameybacter massiliensisspecies24.6849,347BacteriaPasteurellaceaefamily11.7404,812BacteriaClostridioides difficilespecies5.8199,35347.6EukaryotaBoreoeutheria-4.2145,9691BacteriaRomboutsia lituseburensisspecies3.7126,405BacteriaEscherichia colispecies3.2110,84321.5BacteriaHelicobactergenus1.138,5811	Strong signals					
BacteriaNiameybacter massiliensisspecies24.6849,347BacteriaPasteurellaceaefamily11.7404,812BacteriaClostridioides difficilespecies5.8199,35347.6EukaryotaBoreoeutheria-4.2145,9691BacteriaRomboutsia lituseburensisspecies3.7126,405BacteriaEscherichia colispecies3.2110,84321.5BacteriaPaenibacillusgenus1.447,8481BacteriaHelicobactergenus1.138,5811	SuperKingdom	Organism	Rank	%%	Kbp	Coverage
BacteriaPasteurellaceaefamily11.7404,812BacteriaClostridioides difficilespecies5.8199,35347.6EukaryotaBoreoeutheria-4.2145,969-BacteriaRomboutsia lituseburensisspecies3.7126,405BacteriaEscherichia colispecies3.2110,84321.5BacteriaPaenibacillusgenus1.447,848BacteriaHelicobactergenus1.138,581	Bacteria	Clostridium	genus	37.3	1,288,845	
BacteriaClostridioides difficilespecies5.8199,35347.6BacteriaBoreoeutheria4.2145,96947.6BacteriaRomboutsia lituseburensisspecies3.7126,405BacteriaEscherichia colispecies3.2110,84321.5BacteriaPaenibacillusgenus1.447,848BacteriaHelicobactergenus1.138,581	Bacteria	Niameybacter massiliensis	species	24.6	849,347	
EukaryotaBoreoeutheria4.2145,969BacteriaRomboutsia lituseburensisspecies3.7126,405BacteriaEscherichia colispecies3.2110,84321.5BacteriaPaenibacillusgenus1.447,84847,848BacteriaHelicobactergenus1.138,5811	Bacteria	Pasteurellaceae	family	11.7	404,812	
BacteriaRomboutsia lituseburensisspecies3.7126,405BacteriaEscherichia colispecies3.2110,84321.5BacteriaPaenibacillusgenus1.447,848BacteriaHelicobactergenus1.138,581	Bacteria	Clostridioides difficile	species	5.8	199,353	47.6
BacteriaEscherichia colispecies3.2110,84321.5BacteriaPaenibacillusgenus1.447,848BacteriaHelicobactergenus1.138,581	Eukaryota	Boreoeutheria		4.2	145,969	
BacteriaPaenibacillusgenus1.447,848BacteriaHelicobactergenus1.138,581	Bacteria	Romboutsia lituseburensis	species	3.7	126,405	
Bacteria Helicobacter genus 1.1 38,581	Bacteria	Escherichia coli	species	3.2	110,843	21.5
	Bacteria	Paenibacillus	genus	1.4	47,848	
Bacteria Paeniclostridium sordellii species 0.8 28,640 8.2	Bacteria	Helicobacter	genus	1.1	38,581	
	Bacteria	Paeniclostridium sordellii	species	0.8	28,640	8.2
Bacteria Enterococcus faecalis species 0.4 14,079 4.7	Bacteria	Enterococcus faecalis	species	0.4	14,079	4.7
Bacteria Staphylococcus aureus species 0.3 11,072 3.9	Bacteria	Staphylococcus aureus	species	0.3	11,072	3.9
Bacteria Enterococcus faecium species 0.3 10,030 3.4	Bacteria	Enterococcus faecium	species	0.3	10,030	3.4

1

2 Fig. 2b. Distribution of the reads in the raw data. The individual distribution is given and in the

3 second part, the reads which contribute to a higher extent are given.



2 Fig. 2c. Krona chart of the anal swab of Rhinolophus affinis: Fecal swab Taxonomy

- 3
- 4

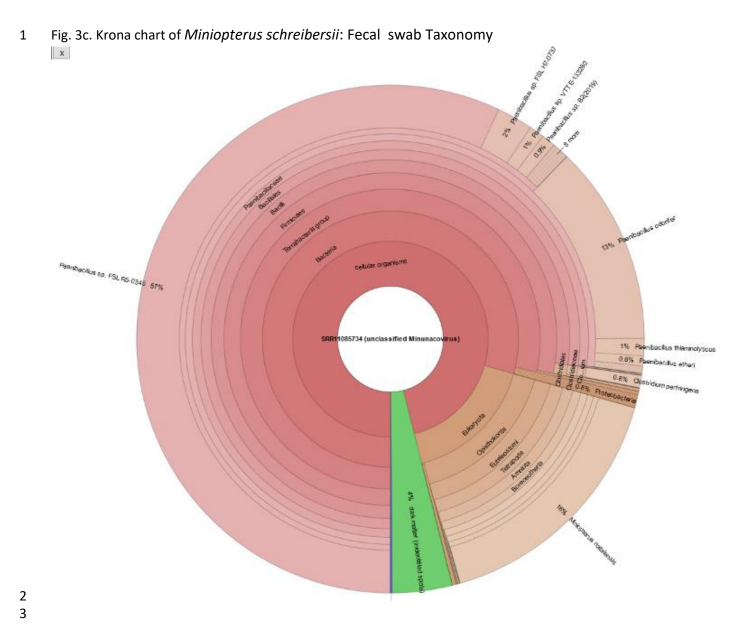
1 Fig 3 RNA-Seq of *Miniopterus schreibersii*: Fecal swab Taxonomy Analysis

RNA-Seq of Miniopterus schreibersii: Anal swab (SRR11085734)

Faxonomy An	alysis				
⊡Am ⊢N ■Fungi:	ds: 3.81% : 96.19% isms: 96.08% 5.15% 16.03% onta: 10% a: 9.99% aria: 9.99% niota: 7.67% 1iniopterus natalensis: 5.98% < 0.01% (1 Kbp) ntae: 0.16%				
Strong signals					
SuperKingdom	Organism	Rank	%%	Кbр	Coverage
Bacteria	Paenibacillus	genus	77.2	2,124,474	
Eukaryota	Miniopterus natalensis	species	16.3	449,835	
Bacteria	unclassified Paenibacillus		5.9	162,892	
Bacteria	Paenibacillus sp. FSL R5-0345	species	1.6	44,539	
Bacteria	Paenibacillus odorifer	species	1.2	33,308	4.8
Bacteria	Clostridium perfringens	species	0.8	22,508	6.3
Bacteria	unclassified Massilia		0.5	14,457	
			0.2	5,115	
Bacteria	Mycoplasma	genus	0.2	0,110	

2

3 Fig. 3a. RNA-Seq of fecal swab *Miniopterus schreibersii*



1 Fig. 4

Full 🗸

Send to: -

```
<u>SRX8357956</u>: amplicon_sequences of RaTG13
1 CAPILLARY (AB 310 Genetic Analyzer) run: 33 spots, 30,576 bases, 1.1Mb downloads
```

Design: Primer-based amplicon sequences

Submitted by: Wuhan Institute of Virology, Chinese Academy of Sciences

Study: Bat coronavirus RaTG13 Genome sequencing <u>PRJNA606165</u> • <u>SRP249482</u> • <u>All experiments</u> • <u>All runs</u> <u>show Abstract</u> Sample:

SAMN14082201 • SRS6146537 • All experiments • All runs Organism: unidentified coronavirus

Library:

Name: RaTG13_amplicon_sequences Instrument: AB 310 Genetic Analyzer Strategy: AMPLICON Source: METAGENOMIC Selection: PCR Layout: SINGLE

Runs: 1 run, 33 spots, 30,576 bases, 1.1Mb

Run	# of Spots	# of Bases	Size	Published
SRR11806578	33	30,576	1.1Mb	2020-05-19

```
2 ID: 10870921
```

```
3
```

4 **References:**

- 5 2020. <u>https://www.thetimes.co.uk/article/seven</u> year covid trail revealed I5vxt7jqp. *The Sunday* 6 *Times.*
- Ge, X. Y., Wang, N., Zhang, W., Hu, B., Li, B., Zhang, Y. Z., Zhou, J. H., Luo, C. M., Yang, X. L., Wu, L. J.,
 Wang, B., Zhang, Y., Li, Z. X. & Shi, Z. L. 2016. Coexistence of multiple coronaviruses in
 several bat colonies in an abandoned mineshaft. *Virol. Sin.*, 31, 31-40.
- Rahalkar, Monali C. & Bahulikar, Rahul A. 2020. Understanding the origin of 'BatCoVRaTG13', a virus
 closest to SARS-CoV-2.
- 12 Zhang, Daoyu 2020. Anomalies in BatCoV/RaTG13 sequencing and provenance.
 13 <u>https://zenodo.org/record/3969272#.Xy0m5jVS_IX</u>.
- Zhou, P., Yang, X. L., Wang, X. G., Hu, B., Zhang, L., Zhang, W., Si, H. R., Zhu, Y., Li, B., Huang, C. L.,
 Chen, H. D., Chen, J., Luo, Y., Guo, H., Jiang, R. D., Liu, M. Q., Chen, Y., Shen, X. R., Wang, X.,
 Zheng, X. S., Zhao, K., Chen, Q. J., Deng, F., Liu, L. L., Yan, B., Zhan, F. X., Wang, Y. Y., Xiao, G.
 F. & Shi, Z. L. 2020. A pneumonia outbreak associated with a new coronavirus of probable
 bat origin. *Nature*, 579, 270-273.