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# Probable Pangolin Origin of SARS-CoV-2 Associated with the COVID-19 Outbreak



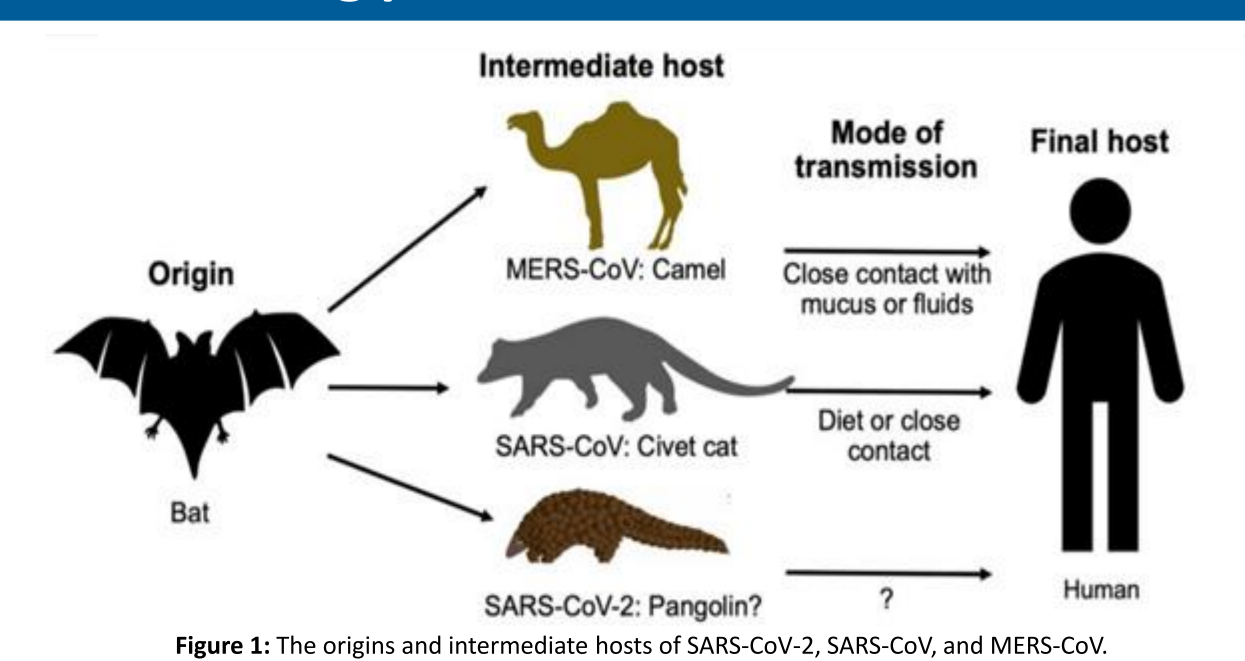
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### **Abstract**

The outbreak of the 2019 novel coronavirus (SARS-CoV-2) in the city of Wuhan in China and its extensive impact on human health requires further research on affected animal species with CoV-like viruses. Phylogenetic analysis is essential to examine key information about CoV-like viruses that could potentially help combat the spread of COVID-19. To discover evolutionary relatives, it is necessary to determine the presence and similarity of spike protein genes as well as specific motifs like the furin recognition sequence. However, apart from BatCoV RaTG13, there are more known species affected by potentially highly similar CoV-Like viruses, with their affinity to the SARS-CoV-2 virus being still unexplored. Here they show that Pangolin-CoV is the second-closest known relative to SARS-CoV-2, right behind BatCoV RatG13, with an identity of 91.02 % on the whole genome level. Additionally, they analyzed the identity of the spike protein, which is around 97.5 %, indicating functional similarities between the Pangolin-CoV and the SARS-CoV-2 virus. Moreover, they found that the highly conserved Nucleocapsid (N) protein is also present in the Pangolin-CoV. The N-Protein is often used as a marker in diagnostic assays, while the spike protein is essential for the cell-entry process of the virus, giving information about the pathogenic potential. With these findings, the pangolin species can be classified as a natural reservoir of SARS-CoV-2 like CoVs. Further experiments should be performed to deepen the knowledge about the interconnection of potentially related viruses to be able to block interspecies transmission.

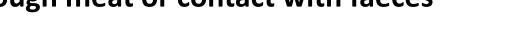
# Finding potential intermediate hosts



- Pangolins are the most poached and trafficked mammal in the world
- → all eight species are considered endangered

Low immunity and poor health condition

- Known infection with SARS-CoV-Like viruses
- Potential intermediate host for SARS-CoV-2
- infection through meat or contact with faeces





Why pangolins?

# Figure 2: Adult pangolin.

# **Workflow and Results**

### Data collection & preprocessing

- Raw RNA reads from collected pangolin lung samples
- Quality control/contamination removal [Trimmomatic]
- Identify virus reads with reference based mapping [Bowtie2]

# De novo assembly & gene prediction

- De novo assembly & taxonomical annotation of the assembled contigs
- Draft genome from annotated contigs [reference guided scaffolding]
- Align draft genome with SARS-CoV-2 for potential ORF's [MUSCLE]
- Compare ORF-identity [PangoCoV/BatCoV and PangoCoV/Sars-CoV]

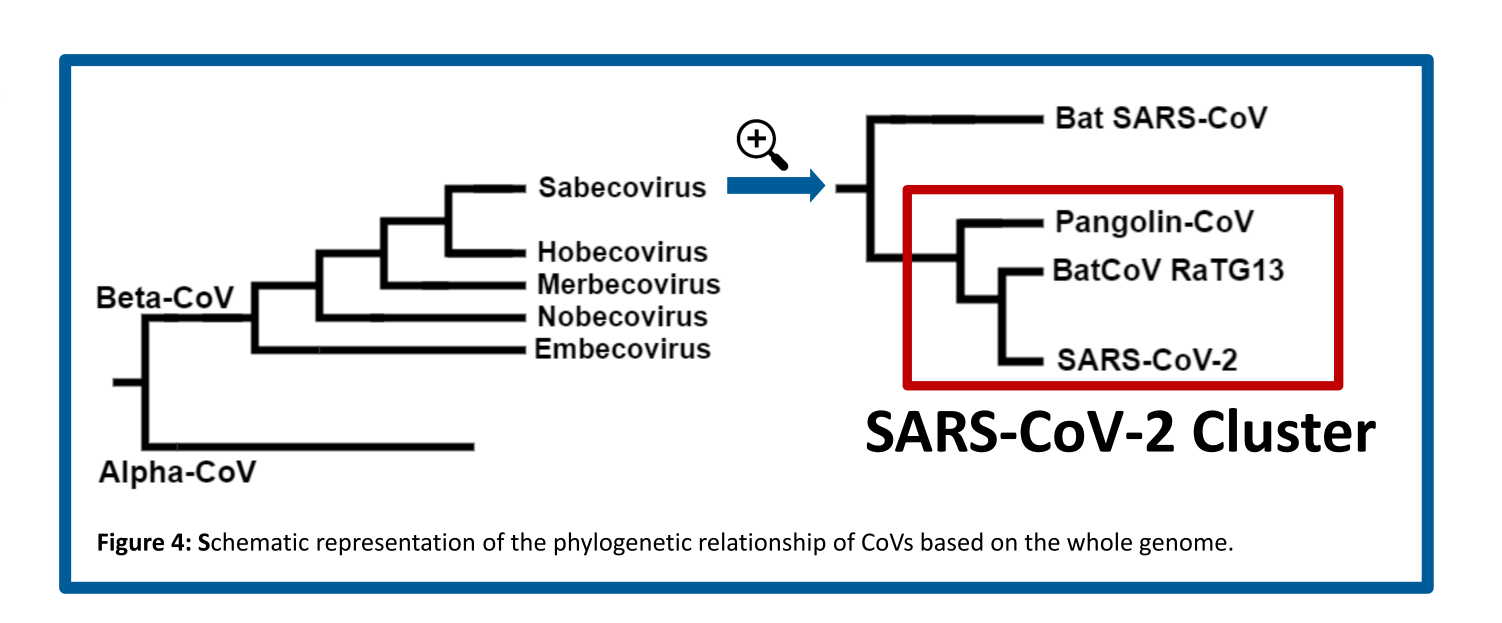
### Phylogenetic analysis

- Generate phylogenetic trees [MegaX]
- Based on whole genome, orf1a, orf1b, M-gene, S-gene, RdRp-gene
- SARS-CoV-2 cluster [BatCoV RatG13, SARS-CoV-2, Pangolin-CoV]

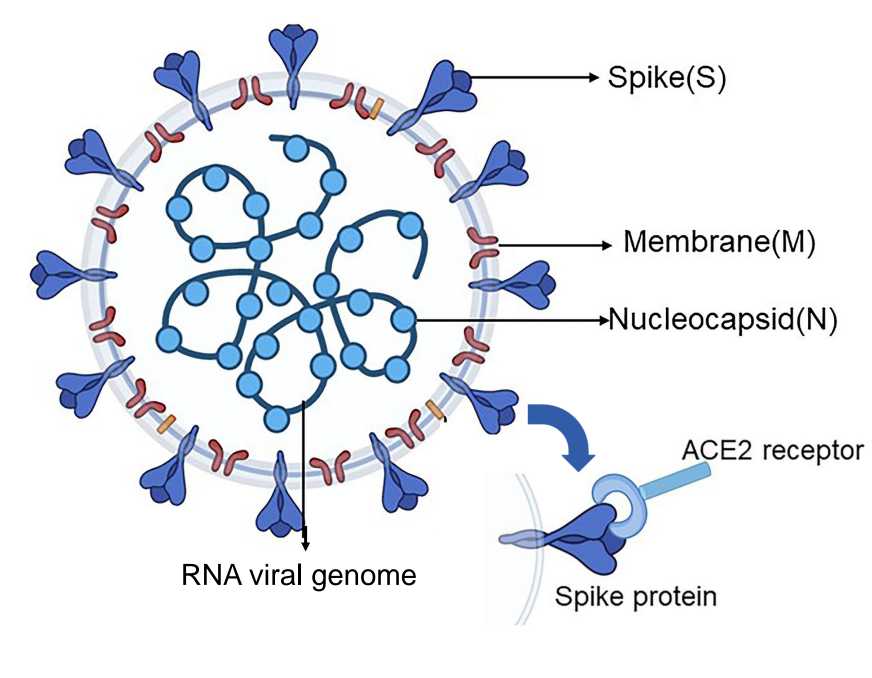
### Main results summarized

- Whole genome identity to SARS-CoV-2: **91.02** %
  - second closest known relative to SARS-CoV-2
- Potential common ancestor of SARS-CoV-2 and BatCoV RatG13
- Highly similar spike protein and nucleocapsid protein
  - potential functional similarities

### Non-structural Structural SARS-CoV-2 **|97.5 %|** 73.4 % 72.8 % |94.6 %| BatCoV RaTG13 Figure 3: Comparison of common genome organization similarity among SARS-CoV-2, Pangolin-CoV, and BatCov RaTG13. **Comparison of the genomic structure** Some Pangolin-CoV showed higher Whole genome identity: similarity to SARS-CoV-2 than to Pango-CoV/SARS-CoV: 91.02 % RaTG13 • Pango-CoV/BatCoV : **90.55** % → orf1b, orf7a, orf10, spike protein



# Structure of the coronavirus



### SARS-CoV-2 Spike protein S2 RBD S1

### Figure 5: Schematic representation of the structure of SARS-CoV-2 coronavirus and the organization of its spike protein.

### Spike protein:

Enables cell entry into receptor-expressing cells [RBD binding to ACE2 receptors in human cells] → Insights into the pathogenic potential

- S-protein gets cleaved into S1/S2-Subunits
- → Cleavage motif necessary

### **Nucleocapsid protein:**

**Priming-phase:** 

- Highly immunogenic phosphoprotein
- → Very conserved & used for diagnostic assays

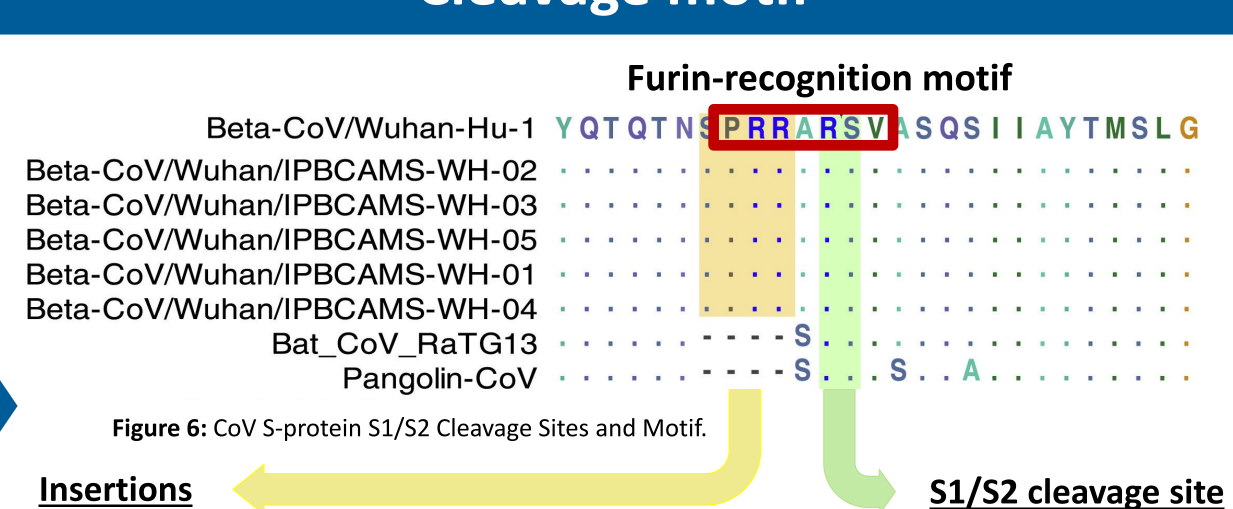
### **Envelope protein:**

support protein helping newly generated virons fold correctly

### Membrane protein:

support protein for the assembly of newly generated virons

# Cleavage motif



- Insertions in SARS-CoV-2 cause an furin-recognition cleavage site
  - No insertions in RaTG13 and Pangolin-CoV
- **PRRARSV**-sequence works as cleavage-recognition-site for host-cell proteases [furin]

# Conclusion

- First report on a potential closely related kin pangolin-CoV of SARS-CoV-2
- Further research necessary to be able to determine if pangolins are a potential candidate related to the COVID-19 outbreak
- The given findings suggest to classify the pangolin species as a natural reservoir for SARS-CoV-like CoV's