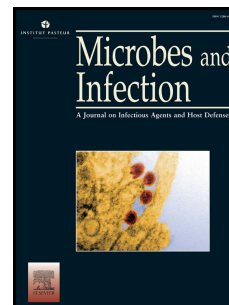


# Journal Pre-proof

Molecular basis of COVID-19 relationships in different species: a one health perspective

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1 **Molecular basis of COVID-19 relationships in different species: a one health**  
2 **perspective.**

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19 **Abstract:**

20 Outside the Hubei province, China, the mild form of infection and the progressive recover of the  
21 COVID-19 patients suggest the intervention of “unconventional” biological mechanisms worthy of  
22 attention. Based on the high-homology between the Spike protein epitopes of taxonomically-related  
23 coronaviruses, we hypothesized that past contact with infected dogs shield humans against the  
24 circulating SARS-CoV-2. Elseways, the recurrent virus exposure over a short time-lapse might  
25 result in the Antibody Dependent Enhancement, triggering the violent immune reaction responsible

26 for the severe clinical outcomes observed in the Hubei province. Nevertheless, further experimental  
27 studies are desired for a confidential evaluation of the postulated hypotheses.

28

29 **Keywords:** SARS-CoV-2; COVID-19; coronavirus; canine respiratory coronavirus; immunization;  
30 one health.

31

32 Since late December 2019, the world is facing a novel outbreak of an infectious disease, known as  
33 COVID-19, caused by an unidentified viral agent. Examination of patients cases reports clinical  
34 picture comparable to this observed in the SARS outbreak of 2003. This evidence is also supported  
35 by the early identification of the etiologic agent as a novel Severe Acute Respiratory Syndrome  
36 Coronavirus (SARS-CoV-2) and the molecular investigations describing a high percentage of  
37 sequence homology between SARS-CoV-2 and the previous SARS-causing virus (SARS-CoV)  
38 [1,2]. Coronaviruses are viruses with single-strand positive-sense RNA genome of approximately  
39 30 kilobases [3]. They have been commonly found in a wide array of host spectrum including avian,  
40 camels, bats, masked palm civets, mice, dogs, and cats [2]. Since its early identification, in the very  
41 begin of 2020, several nucleotides-based surveys and sequence homology-based alignments have  
42 been carried out with the focus to elucidate the viral origin and delineate cell- and host-tropism of  
43 the viral agent on the attempt to guide better management and control of the virus diffusion [4].  
44 Sequence similarity alignments have reported SARS-CoV-2 being highly similar to several wildlife  
45 animals such as bats, pangolins and masked palm civets [3]. Controversial results have been  
46 reported concerning the snake coronavirus [2,5]. Nevertheless, the sole investigation of the genetic  
47 sequences might result in a biased and/or incomplete depiction of the virus source, biology and its  
48 pathogenetic mechanism [4]. In the Hubei province, where the outbreak originated, severe cases or  
49 deaths attributed to SARS-CoV-2 infections are predominantly arising from patients suffering from  
50 one or more previous pathological conditions such as diabetes, cardiovascular and cerebrovascular

51 disease leading to sequelae, sometimes fatal, such as cellular immune deficiency, coagulation  
52 activation, myocardia injury, hepatic and kidney injury, and secondary bacterial infection [4,6].  
53 Although featured by a higher fatality rate when compared with the virus of the seasonal influenza  
54 (3.4%, WHO Situation Report-37, 25/02/2020), outside the primordial outbreak site, much less  
55 warning clinical signs have been registered in the SARS CoV-2 infected patients manifesting a mild  
56 form of the disease that is not evolving into severe stages and can, in some cases, recover without  
57 medical intervention [4]. Altogether this suggests the important contribution of the biological  
58 mechanisms that cannot be explained by the sole investigation of the genetic sequences.

59 Protein repertoire of the coronaviruses consists of four main structural and approximately 16 major  
60 non-structural proteins [3,7]. Among the four structural proteins, the Spike protein (also known as  
61 S-protein) is involved in host tropism by means of recognition and attachment to the angiotensin-  
62 converting enzyme 2 (ACE2) receptor exposed in the outer layer of the host cell membrane [8]. A  
63 very recent amino acid sequence alignment of ACE2 from different animal species including  
64 humans, pets and the major domestic animals highlighted high protein sequence similarity between  
65 the tested animal species, suggesting a potential interaction of the viral particles with a wider array  
66 of host cells [8].

67 In accordance with the hypothesis of interspecies transmission of the beta coronaviruses [8], we  
68 performed sequence homology analysis of the aminoacidic sequence of the Spike protein from  
69 SARS CoV-2 against taxonomically related coronaviruses with tropism for other animal species.  
70 Although the low homology scored while comparing the whole protein sequence (Tab. 1), particular  
71 interest was dedicated to the canine respiratory coronavirus, the bovine coronavirus revealing and  
72 the human enteric coronavirus 36.93%, 38.42% and 37.68% sequence homology to SARS CoV-2  
73 spike protein, respectively.

74

75 Table1

76

77 A previous study of Hua and colleagues on the amino acid sequence of SARS CoV spike protein  
78 identified six epitopes based on the hydrophilicity, surface probability, antigenic index, and  
79 secondary structure [9] (Tab.2).

80

81 Table2

82

83 The sequence homology analysis restricted to the epitope sequences of the SARS CoV-2 revealed  
84 instead high percentage homology towards taxonomically related coronaviruses. Of these, we retain  
85 it is worth of note the high similarity occurring among 4 epitope sequences of canine respiratory  
86 coronavirus (Canine respiratory CoV BJ 232) and the circulating SARS CoV-2 (Table 3).

87

88 Table3

89

90 Highlighting the similarity at the epitope level opens new avenues in understanding the biological  
91 mechanisms undertaken during viral infection. Before the current outbreak, several studies witness  
92 the common identification of coronavirus infection in dogs [10,11]. Previous studies performed by  
93 Szczepanski and colleagues report the possibility to grow both bovine and canine coronavirus on  
94 human cells lines such as the Human Rectal Tumor cells (HRT-18G) and Human Airway  
95 Epithelium (HAE) although featured by a strongly reduced replication efficiency [12,13]. In this  
96 view, a previous “contact” with the canine virus might provide at least a partial/basal immunization  
97 that shields humans against the circulating SARS CoV-2. This would partially explain the mild  
98 symptoms registered among patients outside the Hubei province and without previous co-  
99 morbidities. Accordingly, high similarity percentage have also been observed between SARS CoV-  
100 2 spike proteins epitopes and bovine coronavirus, which genome and proteins appear among those  
101 included in the patented vaccinal formulation released against coronavirus.

102 With regard to the most severe cases of infection registered both outside and within the Hubei  
103 province, it is most likely that these individuals have been primed by one or more coronavirus

104 exposure within a narrow time window, leading to the effects of Antibody-Dependent Enhancement  
105 (ADE) [4]. Such a hypothesis is in line with the previous observations dating back to the previous  
106 SARS outcome in 2003 when human coronaviruses, known to cause mild infections, were  
107 hypothesized of priming the high mortality scored in China. Here, anti-Spike protein antibodies  
108 were indicated as potentially involved in the enhancement mechanism [14–18]. In this view, it is  
109 plausible that Hubei province patients are featured by a more severe clinical picture, since more  
110 likely to be recurrently exposed to the coronavirus.

111 To summarize, animals have (had) a critical role in this outbreak onset and evolution.  
112 Acknowledged their pivotal role as virus reservoir, they might act in the first instance as  
113 "beneficial" source of immune-stimulating virus particles; thus, providing a shield against the  
114 circulating SARS CoV-2. However, the recurrent exposure within a narrow time-lapse might results  
115 in the detrimental triggering of violent immune responses and the evolution towards a more severe,  
116 or even fatal, clinical picture.

117 Above hypotheses are based on the assumption that sufficient viral titer of the SARS CoV-2 is  
118 transmitted from patient to patient [19]. The previous investigation on animal betacoronavirus such  
119 as the canine coronavirus have demonstrated a high transmission efficiency through direct contact  
120 between infected animals, acknowledged the high viral titers found in the aerosols of respiratory  
121 secretions [20]. Novel investigations to assess the transmission rate of the canine respiratory  
122 coronavirus to humans are needed to better suit the above hypotheses. Also, care should be taken  
123 while evaluating the data on the sole basis of sequence homology and further studies employing  
124 purified forms of the spike proteins and/or its epitopes are desired.

125

126

127 **Conflict of interest:** The authors declare no conflict of interest

128

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130

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Table 2. Epitope details of the Spike glycoprotein of diverse coronaviruses.

Protein GI	Epitope AAs	Epitope sequence	Organism	NCBI TaxID
<b>QHR63290</b>	424-437	KLPDDFTGCVIAWN	SARS coronavirus 2	2697049
	447-458	NYNYLYRLFRK		
	560-571	LPFQQFGRDIAD		
	754-764	LQYGSFCTQLN		
	789-799	YKTPPIKDFGG		
	1139-1152	DPLQPELDSFKEEL		
<b>QAY30020</b>	424-437	SGYTVAATFASLFP	Canine respiratory coronavirus	215681
	447-458	FYLNQYRINGI		
	560-571	QLSDSTLVKFSA		
	754-764	TYEYVVKWPWY		
	789-799	CGTSCFKKCGG		
	1139-1152	DPLQPELDSFKEEL		
<b>AHA50776</b>	424-437	ATSCQLYYNLPAAN	Bovine coronavirus	11128
	447-458	TWNRRFGFTEQS		
	560-571	EHCSGLAIKSDH		
	754-764	SGYCVDYSTKR		
	789-799	DSLEPVGGLYE		
	1139-1152	NGNHIISLVQNAPY		
<b>ACJ35486</b>	424-437	ATSCQLYYNLPAAN	Human enteric coronavirus	166124
	447-458	TWNRRFGFTEQS		
	560-571	EHCSGLAIKSDH		
	754-764	SGYCVDYSTKR		
	789-799	DSLEPVGGLYE		
	1139-1152	NGNHIISLVQNAPY		

Table3. Epitope sequence alignment. The table summarizes results from the alignment of SARS CoV-2 spike protein epitopes against the spike protein epitopes of other coronaviruses with known tropism for other animals than humans.

<b>Epitope</b>	<b>Organism</b>	<b>% identity</b>
424-437	Bovine Coronavirus	80,00
	Canine respiratory coronavirus	80,00
	Human enteric coronavirus	80,00
447-458	Bovine Coronavirus	75,00
	Canine respiratory coronavirus	-
	Human enteric coronavirus	75,00
754-764	Bovine Coronavirus	83,33
	Canine respiratory coronavirus	83,33
	Human enteric coronavirus	83,33
789-799	Bovine Coronavirus	57,14
	Canine respiratory coronavirus	57,14
	Human enteric coronavirus	57,14
1139-1152	Bovine Coronavirus	70,00
	Canine respiratory coronavirus	100,00
	Human enteric coronavirus	70,00

Table 1. Amino acid sequence alignment of SARS CoV-2 Spike protein (GI QHR63290) against betacoronavirus database. The table display alignment relative to taxonomically-related coronavirus having a tropism for other hosts than humans.

<b>Protein Accession</b>	<b>E-value</b>	<b>Organism</b>	<b>NCBITax ID</b>	<b>% identity</b>
AGO98871	1e-154	Bovine Coronavirus	11128	38.42%
QAY30020	3e-152	Canine respiratory coronavirus	215681	36.93%
ACJ35486	3e-150	Human enteric coronavirus	166124	37.68%
ACT10865	2e-105	Feline coronavirus	12663	32.26%
ABQ57216.1	4e-100	Bat coronavirus	693998	31.23%
AID16631	1e-145	Mouse coronavirus	1508222	36.56%
AID16649	8e-149	Rat coronavirus	1508223	37.60%