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RESEARCH PAPER



From the Wuhan-Hu-1 strain to the XD and XE variants: is targeting the SARS-CoV-2 spike protein still a pharmaceutically relevant option against COVID-19?

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ABSTRACT

Since the outbreak of the COVID-19 pandemic in December 2019, the SARS-CoV-2 genome has undergone several mutations. The emergence of such variants has resulted in multiple pandemic waves, contributing to sustaining to date the number of infections, hospitalisations, and deaths despite the swift development of vaccines, since most of these mutations are concentrated on the Spike protein, a viral surface glycoprotein that is the main target for most vaccines. A milestone in the fight against the COVID-19 pandemic has been represented by the development of Paxlovid, the first orally available drug against COVID-19, which acts on the Main Protease (Mpro). In this article, we analyse the structural features of both the Spike protein and the Mpro of the recently reported SARS-CoV-2 variant XE, as well the closely related XD and XF ones, discussing their impact on the efficacy of existing treatments against COVID-19 and on the development of future ones.

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KEYWORDS

SARS-CoV-2; COVID-19; XE variant; XD variant; Spike protein; Mpro; Paxlovid

1. Introduction

More than two years have now passed since the beginning of the COVID-19 pandemic, back in December 2019^{1,2}. Caused by a beta coronavirus known as SARS-CoV-2 and characterised by para-influenzal symptoms such as fever, cough, and dyspnoea, this world-wide-spread disease has resulted in the death of more than 6 million people around the world, becoming one of the deadliest illnesses in human history^{3,4}.

The SARS-CoV-2 virus was first identified in the Chinese city of Wuhan, where the pandemic was firstly spotted⁵. The genomic sequence of this virus (named Wuhan-Hu-1 from now on in the article) is 80% identical to the one of the SARS-CoV virus^{6,7}, which was responsible for the Severe Acute Respiratory Syndrome (SARS) that struck the South East of Asia in 2002/2003, causing the death of about 800 patients over 9000 cases (10% death rate)^{8,9}. The exact origin of the SARS-CoV-2 virus is still to this date unknown, however several pieces of evidence point out bat coronaviruses as closely related ancestors and the pangolin as the intermediate host before the human spillover^{10–13}.

Soon after the original virus started spreading all over the world, several viral variants began to emerge^{14,15}, especially in the poorest countries where public health measures such as social distancing and wearing surgical masks in public places were difficult to implement^{16–18}. Most of the genome mutations that characterised these variants were concentrated in the S gene¹⁹, which encodes for the Spike protein, a surface glycoprotein that mediates the virus entry within the human cell through an interaction with the human ACE2 receptor²⁰. Some of these mutations gathered the attention of the scientific community due to the selective advantage that they provided to the correspondent viral variants,

regarding both the virus' ability to infect human cells and to escape the immune system response²¹, gaining for these reasons the status of "variant of concern" (VOC).


The first SARS-CoV-2 variant to be labelled as VOC was the so-called Alpha variant (B.1.1.7). First identified in November 2020 in the Kent region of the United Kingdom and for this reason also known as the "English variant", B.1.1.7 was estimated to be 29% more transmissible than the original virus^{22,23}. Despite being more transmissible than other circulating viral strains^{24,25}, and despite showing the first signs of reduced protection provided by vaccines, monoclonal antibodies, and convalescent sera^{26–28}, the indication from clinical studies showed that the vaccine coverage (especially in those who had already completed the vaccination cycle) was still able to contain the impact of this variant on the sanitary system²⁹.

Soon after the identification of the Alpha variant, a second VOC arose: the Delta variant (B.1.617.2), also known as the "Indian variant" due to being first detected in India in late 2020, rapidly overcame the Alpha variant becoming the dominant strain in the world, thanks to being 97% more transmissible than the original Wuhan virus²³. The replacement of the less threatening Alpha variant with the Delta marked a significant change of pace in the pandemic trend, signing an increased burden for the health system³⁰ and posing for the first time a serious threat to the protection provided by vaccines, convalescent sera, and monoclonal antibodies^{31–33} due to its increased ability to evade the immune system response³⁴.

From November 2021 onwards, the Delta variant has been flanked by another VOC firstly identified in South Africa and defined as the Omicron variant (B.1.1.529)³⁵. The rise of the Omicron variant, fuelled by a contemporary increase in

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Table 1. The genome sequences used in this work and their origin.

Organism	Isolate	Accession Code
SARS-CoV-2	"Wuhan-Hu-1"	NC_045512.2
SARS-CoV-2 "Delta"	"SARS-CoV-2/human/JPN/SARS-CoV-2"	OK091006.1
SARS-CoV-2 "Omicron"	"SARS-CoV-2/human/NLD/EMC-Omicron-1/2021"	OM287553.1
SARS-CoV-2 "XD"	"SARS-CoV-2/human/FRA/IHUCOVID-64762/2022"	OM990851.1
SARS-CoV-2 "XE"	/	OW018845.1
SARS-CoV-2 "XF"	/	OV940149.1

transmissibility³⁶ and immune evasion³⁷, resulted in an unprecedented diffusion of the SARS-CoV-2 virus all over the world, being able to overcome even the protection provided by the full primary vaccination cycle and by most neutralising antibodies used in therapy^{38–40}, thus inducing the introduction of a "booster dose" to bring the protective effect of vaccines back to adequate levels^{41,42}.

In the face of this increasingly troublesome variant landscape, characterised by a progressive reduction of the efficacy of existing therapeutic options against COVID-19, a light at the end of the tunnel is possibly represented by the development and release on the market of Paxlovid. This therapeutic combination between the active principle Nirmatrelvir (also known as PF-07321332) and the pharmacokinetic enhancer Ritonavir, represents the first orally available drug specifically designed against SARS-CoV-2 virus⁴³. Instead of targeting the Spike protein, this peptidomimetic entity is designed to inhibit the SARS-CoV-2 Main Protease (M^{Pro}) by covalently binding to Cysteine 145, one of the two components of the protease's catalytic diad⁴⁴. Clinical studies showed a remarkable therapeutic efficacy of this novel treatment, with Paxlovid being able to lower by 89% the risk of severe complications associated with COVID-19 infection in symptomatic, non-vaccinated, and non-hospitalized adult patients⁴⁵.

Recently, three novel recombinant SARS-CoV-2 variants were identified in the United Kingdom: Xd, Xe, and Xf⁴⁶. These variants are derived from the combination of the genomes of other major circulating variants, namely Delta, Omicron, and Omicron 2^{46,47}. Among these three novel viral strains, particular worry is related to the XE variant, which derives from the recombination between two VOCs, Omicron and Omicron 2, and is supposed to be 13–20% more transmissible than the Omicron 2 variant⁴⁶.

The rise of novel SARS-CoV-2 variants derived from the recombination of other threatening and heavily diffused ones poses a serious challenge in the fight against the COVID-19 pandemic since it could contribute to rendering existing therapeutic options inefficient or practically useless. To evaluate the impact that these recently reported recombinant variants could have on the efficacy of existing vaccines and treatments (Paxlovid, in particular), we performed a computational analysis to shed light on the key structural features that characterise both the Spike glycoprotein and the Main Protease of these novel viral strains. Moreover, we analysed the structural evolution of these two viral proteins throughout the pandemic, discussing the impact that mutations found on these strains had and will have on the efficacy of existing therapeutic options against COVID-19 and the development of future ones.

2. Materials and methods

The genome sequence for the SARS-CoV-2 virus and its variants, namely Delta, Omicron, XD, XE, and XF, was obtained through GenBank⁴⁸. Accession codes for each of the considered genomes are reported in Table 1. In the case of newly discovered variants

Table 2. List of all the single-point mutations affecting the SARS-CoV-2 Spike protein for all the variants considered in our study (Delta, Omicron, XE, XD, and XF).

Delta variant	Omicron variant	XD variant	XE variant	XF variant
T19R		T19R	T19R	
		A27S	A27S	
	A67V			A67V
	T95I	T95I		T95I
	G142D	G142D	G142D	
				Y145D
R158G		R158G		
	L212I	L212I		L212I
			V213G	
	G339D	G339D	G339D	G339D
	S371L	S371L	S371L	S371L
	S373P	S373P	S373P	S373P
	S375F	S375F	S375F	S375F
			T376A	
			D405N	
			R408S	
	K417N	K417N	K417N	K417N
	N440K	N440K	N440K	N440K
	G446S	G446S		G446S
L452R		S477N	S477N	S477N
T478K	S477N	T478K	T478K	T478K
	T478K	E484A	E484A	E484A
	E484A	Q493R	Q493R	Q493R
	Q493R	G496S	G496S	G496S
	G496S	Q498R	Q498R	Q498R
	Q498R	N501Y	N501Y	N501Y
	N501Y	Y505H	Y505H	Y505H
	Y505H	T547K	T547K	T547K
	T547K	D614G	D614G	D614G
D614G	D614G	H655Y	H655Y	H655Y
	H655Y	N679K	N679K	N679K
	N679K	P681H	P681H	P681H
P681R	P681H	N764K	N764K	N764K
	N764K	D796Y	D796Y	D796Y
	D796Y	N856K	N856K	N856K
	N856K	Q954H	Q954H	Q954H
	Q954H	N969K	N969K	N969K
	N969K	L981F	L981F	L981F
	L981F			

The mutations have been aligned to give a better perspective of the ones which have been conserved through the evolutionary process. The mutations involving the RBD have been highlighted in green, while the ones involving the RBM are coloured in cyan.

XD, XE, and XF, the sequence was chosen according to the one reported by the Nextclade project⁴⁸.

All the basic molecular modelling operations have been executed with the Molecular Operating Environment (MOE) suite (version 2019.01)⁴⁹.

For what concerns the Spike protein, the approach chosen depended on the variant considered. For the wild-type (WT) Spike, the three-dimensional structure was retrieved from the Protein Data Bank (PDB code: 6ZDH⁵⁰, method: cryo-EM, resolution: 3.70 Å), as well as for the Delta (PDB code: 7W9E⁵¹, method: cryo-EM, resolution: 3.10 Å), and the Omicron (PDB code: 7WPD⁵², method: cryo-EM, 3.18 Å) variants. The cited structures were all subjected to the same preparation procedure for molecular modelling.

After being downloaded, the *Structure Preparation* tool implemented in MOE was applied to rebuild the missing loops in the structures, the proper protonation state was assigned to each amino acid with the MOE "Protonate 3D" application, and finally, the added hydrogens were minimised under the AMBER10:EHT⁵³ force field implemented in MOE. Since experimentally resolved structures for the XD, XE, and XF variants are not available in public databases, the models considered for our study were created starting from the WT SARS-CoV-2 Spike coming from the PDB code 6ZDH by manually mutating the residues, exploiting the MOE "Protein builder" tool, and subjecting each protein to the preparation procedure reported above. For the realisation of the video reported in the [Supplementary Materials](#), the program VMD 1.9.2⁵⁴ (Visual Molecular Dynamics) was used.

For what concerns M^{Pro}, the protein sequences corresponding to the main protease were extracted from the whole genome sequence and aligned to the reference sequence (Wuhan-Hu-1) making use of the appropriate tool of MOE 2019.01. Subsequently, homology models for each variant were created making use of the "Homology Model" tool of MOE 2019.01, using the structure deposited in the Protein Data Bank with accession code 6Y2E (Crystal structure of the free enzyme of the SARS-CoV-2 main protease) as a template for model generation.

3. Results and discussion

3.1. Structural analysis of spike glycoprotein mutations found in SARS-CoV-2 XD, XE, and XF variants and their impact on hACE2 binding

The SARS-CoV-2 Spike protein (S) consists of a large biological entity formed by 1273 amino acids organised in different functional domains. The main role of the Spike protein is to mediate the virus entry into the host cell, with the principal and better-characterised mechanism being the pathway involving the binding to the human ACE2 receptor (hACE2)⁵⁵, a membrane-bound enzyme that is widely expressed in various districts of the human body (from the endothelial cells of the blood vessels to kidneys, liver, intestine, lungs⁵⁶, and cells of bronchial and nasal epithelia⁵⁷).

The S protein, which works in a trimeric organisation, is divided into two main subunits, S1 and S2. The second of these has very important roles in spike protein trimerization and in mediating the virion entry into the host cell once the molecular contacts have been established. It is formed by relevant subdomains such as the fusion peptide (FP, residues 943–982, crucial for viral fusion to the host cell membrane), the transmembrane domain (TM, composed of 24 amino acids and deputed both to the anchoring

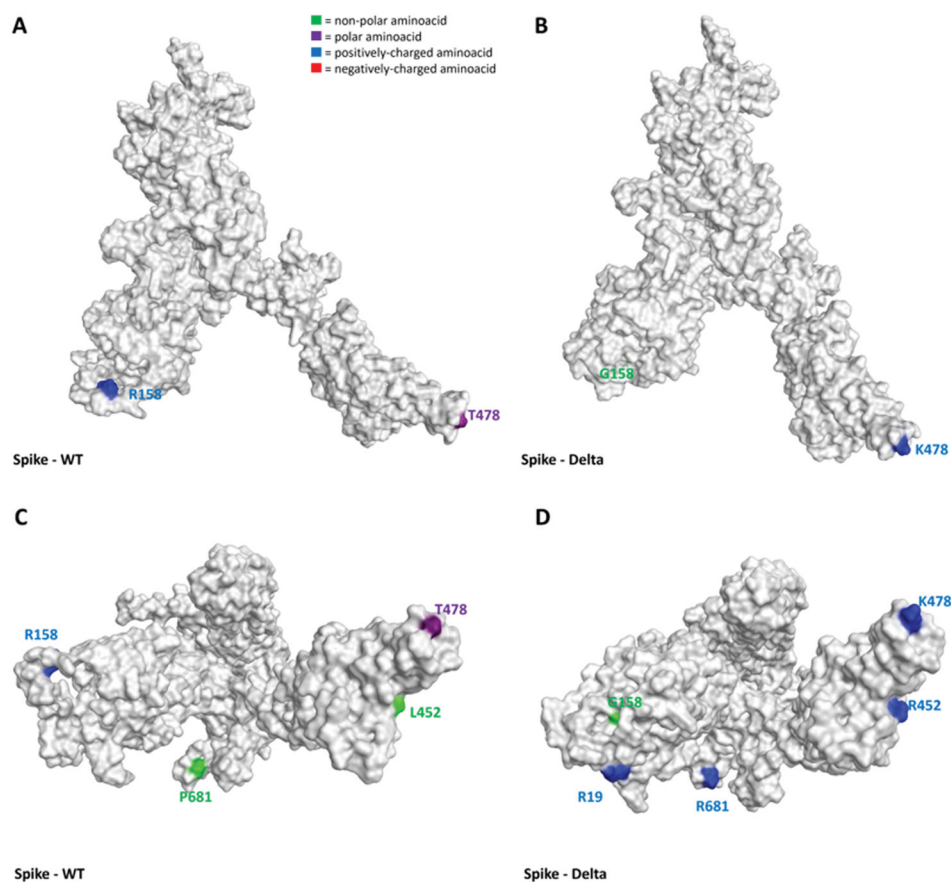


Figure 1. Representation of structural differences between the WT (taken from the Protein Data Bank⁷⁰, PDB code: 6ZDH) and the Delta variant (retrieved from the PDB, code 7W9E⁵¹) of SARS-CoV-2 Spike protein. Panels A and B offer a front view of the comparison between the structures, while panels C and D shift the point of view to the bottom of the proteins. To give a clearer view of the mutations, only one monomer was considered to create the image. The amino acids involved in mutations are labelled in the figure and are coloured based on their kind, following the legend reported in panel A. Specifically, Gly, Ala, Val, Leu, Ile, Pro, Cys, Met, Phe, and Trp are considered non-polar amino acids (green), Asp and Glu represent the negatively-charged amino acids (red), and Lys, Arg, and His form the positively-charged amino acid group (blue). Finally, Ser, Thr, Asn, Gln, and Tyr are all considered polar amino acids (purple). All images were created and rendered using the Molecular Operating Environment (MOE) suite.

of S protein to the viral membrane and the maintenance of the Spike trimeric organisation), and the cytoplasmic fusion domain (CT, mediating virus-cell fusion).

The S1 subunit, instead, contains both the N-terminal and the C-terminal domains (NTD and CTD, respectively), which are involved in the binding to host cell receptors. Specifically, the CTD contains the receptor-binding domain (RBD, aminoacids 319–541), the region deputed to the binding with hACE2. This function is more precisely carried out by a particular RBD subdomain, called receptor-binding motif (RBM), which is formed by two beta-sheets ($\beta 5$ and $\beta 6$) composed of those residues which are in close contact with hACE2 (from 438 to 506)^{58,59}.

Looking at all the S proteins of the different SARS-CoV-2 relevant variants, the RBD contains the highest “single-point mutations/sequence length” ratio in all cases. Examining the different SARS-CoV-2 variants discovered up to date, the Spike protein is surely the viral entity that has mutated the most in the evolutionary process of the virus⁶⁰. Its exposition on the viral surface and its crucial function in viral cell entry make this protein the eligible target for the host immune system⁶¹.

The SARS-CoV-2 S protein has experienced several mutations in the past two years⁴⁸, as reported in Tables 1 and 2 for the variants considered in our study. As can be noticed, variants such as Delta (but also Alpha and Beta, not specifically treated in this article) showed few mutations in the overall viral genome, and Spike

protein displayed never more than a tenth of single-point changes. The game-changing event was the advent of the Omicron variant, much different from its previous analogs, with 30 single nucleotide mutations involving the S protein only. Many of these, such as K417N, T478K, and P614G were inherited from the previous lineages (mainly Beta and Delta), but other mutations were completely new, such as G339D, G446S, or E484A.

Most of the mutations listed in Table 2 have been related to higher infectivity, mainly due to a consequent gain of affinity for hACE2 or improved shielding from the immune cells. As evidence of this, the most successful vaccination campaigns for COVID-19 always involved the forced recognition of Spike protein from the patients’ immune cells⁶².

Specifically, mutations such as S371L, K417N, and Q493R were related to a diminished binding to the anti-coronavirus monoclonal antibody Casivirimab, while mutations like N440K and G446S confer resistance towards the antibody Imdevimab⁶³. The combination of Casivirimab and Imdevimab has been used to treat COVID-19 patients but has demonstrated to be ineffective against the Omicron variant⁶⁴. Other mutations external to the RBD have been linked to different outcomes, such as increased viral replication ($\Delta 69-70$ ⁶⁵ and D614G⁶⁶) and higher viral resistance (G339D and N440K⁶⁷). In other scenarios, mutations have been reported to influence tropism of the S1/S2 cleavage, as in the cases of N679K and P681H⁶⁸.

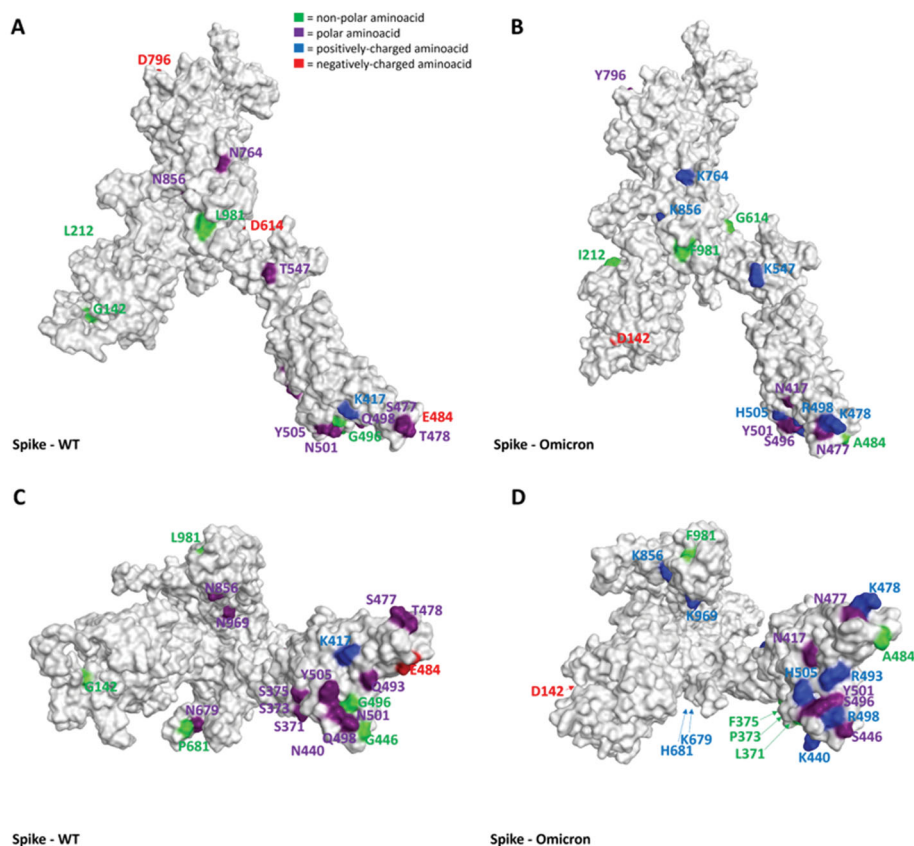


Figure 2. Representation of the differences between the WT (taken from PDB code: 6ZDH) and the Omicron variant (retrieved from the PDB, code 7WPD) of SARS-CoV-2 Spike protein. Panels A and B offer a front view of the comparison between the structures, while panels C and D shift the point of view to the bottom of the proteins. To give a clearer view of the mutations, only one monomer was considered to create the image. The amino acids involved in mutations are labelled in the figure and are coloured based on their kind, following the legend reported in panel A. Specifically, Gly, Ala, Val, Leu, Ile, Pro, Cys, Met, Phe, and Trp are considered non-polar amino acids (green), Asp and Glu represent the negatively-charged amino acids (red), and Lys, Arg, and His form the positively-charged amino acid group (blue). Finally, Ser, Thr, Asn, Gln, and Tyr are all considered polar amino acids (purple). All images were created and rendered using the Molecular Operating Environment (MOE) suite.

The majority of the mutations highlighted up to date on the SARS-CoV-2 S protein impact the binding with hACE2, as in the cases of S477N, Q498R, and N501Y⁶⁹. These last mutations, as can be seen in Table 2, have been conserved from all the variants following Omicron, assessing their importance for the viral evolutionary process.

As can be seen in Figures 1–5 and video.mp4 (Supplementary Materials), the highest “number of mutations/sequence length” ratio is owned by the RDB, as previously mentioned. Indeed, taking Omicron as an example, among the 30 mutations in the overall 1273-residues structure, 15 are located just in the 222 residues-sequence forming the RBD. The insertions and the deletions (summarized in Table 3), on the other hand, are located far outside the hACE2-binding domain in all the variants examined, allowing us to assert that these mutations should not impact all the host cell recognition process.

Interestingly, as depicted in the aforementioned figures, single-point mutations that are present in the RBD for all considered variants tend to progressively increase the positively-charged character of this protein region. Moreover, of all the changes operated by the evolutionary process of Spike protein, the very few mutations which transform a residue into a negatively-charged one (Asp or Glu) are always located away from the RBD (except for G339D, which is located in the posterior part of the RBD, away from the RBM that contacts hACE2). Indeed, in this region, the changes from polar amino acids to positively charged ones are

abundant (N440K, T478K, Q498R, Y505H), and there are also cases in which non-polar residues transform into polar ones (e.g. G446S and G496S, which are conserved in all examined post-Omicron variants except for the XE one). Another conserved structural feature across all post-Omicron variants is also the fact that E484, located in the RBM, mutates into an alanine, while a peculiar mutation exclusive to the XE variant is represented by the transformation of D405 into an asparagine. Taken together, all these pieces of evidence converge in assessing that an increase in the polar characteristics of the RBD (more specifically, the RBM), with particular relevance to an increase in the number of positively charged amino acids, could be the mechanism adopted by SARS-CoV-2 to continuously increase its infectivity through an increase in the interaction with hACE2.

To further support this evidence, in Figure 6 we reported the electrostatic surface of hACE2 complexed with WT-Spike RBD highlighting the prevalence of negative charge on the surface facing Spike RBM (coloured in green in the image). Coherently, the only mutation present in the RBD in which a positively-charged residue shift into a polar one (K417N) has been reported to reduce the affinity with hACE2²⁶. It is worth noting that other hACE2-independent entry routes for SARS-CoV-2 have been described in literature^{71,72}, but the lack of reliable structural information about the interaction with the target at the present date hampers and limits the possibility to analyse and discuss the impact that these mutations could have on them in a meaningful way. However, it

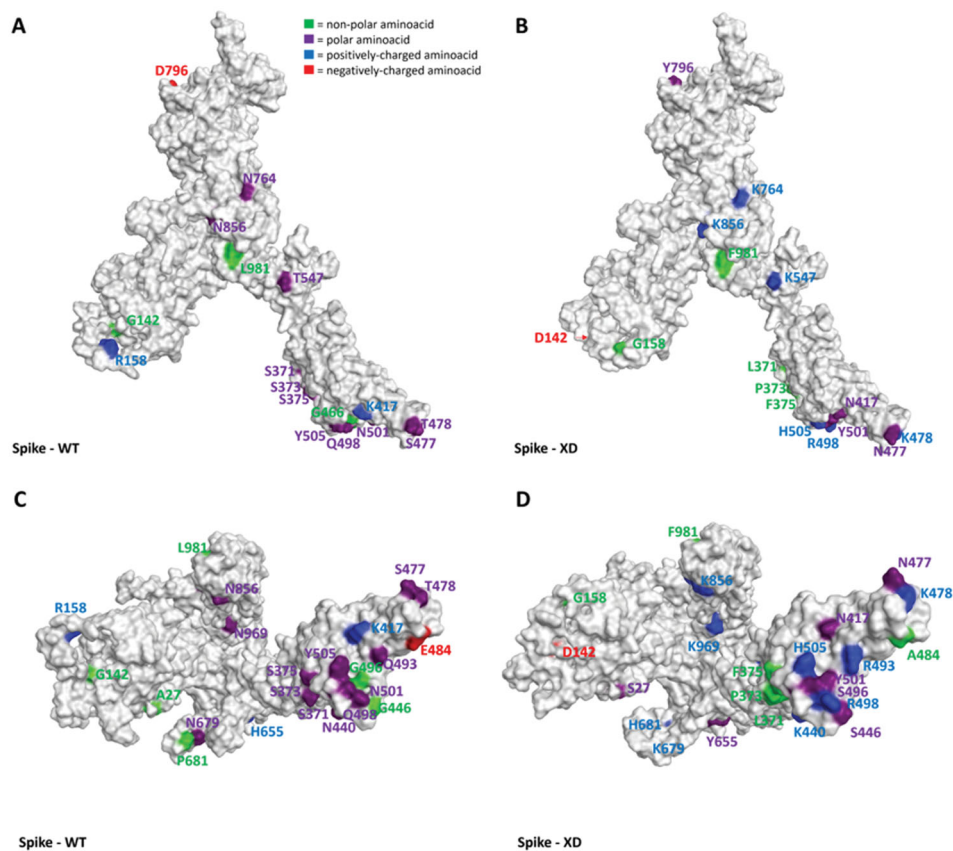


Figure 3. Representation of the differences between the WT (taken from PDB code: 6ZDH) and the XD a variant of SARS-CoV-2 Spike protein. Due to the lack of experimentally resolved structure of SARS-CoV-2 XD variant, the three-dimensional structure represented was obtained from the wild-type S protein coming from PDB code 6ZDH, and then manually mutating the residues involved in the mutations (exploiting the MOE “Protein builder” tool). Panels A and B offer a front view of the comparison between the structures, while panels C and D shift the point of view to the bottom of the proteins. To give a clearer view of the mutations, only one monomer was considered to create the image. The amino acids involved in mutations are labelled in the figure and are coloured based on their kind, following the legend reported in panel A. Specifically, Gly, Ala, Val, Leu, Ile, Pro, Cys, Met, Phe, and Trp are considered non-polar amino acids (green), Asp and Glu represent the negatively-charged amino acids (red), and Lys, Arg, and His form the positively-charged amino acid group (blue). Finally, Ser, Thr, Asn, Gln, and Tyr are all considered polar amino acids (purple). All images were created and rendered using the Molecular Operating Environment (MOE) suite.

cannot be excluded that this mutation pattern and other future Spike mutations could also impact these other entry pathways, contributing to making them more relevant for the ability of SARS-CoV-2 to infect human cells and increasing its overall infectivity.

3.2. Structural analysis of main protease mutations found in SARS-CoV-2 XD, XE, and XF variants and their impact on the recognition of known inhibitors

The main protease M^{pro}, also known as 3C-like protease or 3CL^{pro}, is a cysteine peptidase that is essential for the replication cycle of SARS-CoV-2^{73,74}. Its catalytic activity revolves around the processing of two overlapping polyproteins, namely pp1a and pp1ab, which leads to the formation of 16 mature non-structural proteins (NSPs)⁷⁵. Composed of 306 amino acids, the SARS-CoV-2 M^{pro} shares 96% sequence identity and a highly conserved three-dimensional structure with the SARS-CoV M^{pro} (0.53 Å R.M.SD between PDB entries 6Y2E and 2BX4)^{76,77}. Although a dynamic equilibrium between a monomeric and a dimeric form exists, only the dimer is responsible for the protease's enzymatic activity^{78,79}. Each protomer composing the catalytically active dimer is composed of three different domains: the chymotrypsin-like β -barrel domains I (residues 1–99) and II (residues 100–182), which comprehend the substrate-binding site and directly control the catalytic event, and the extra α -helical domain III (residues 198–306),

which is connected to the remaining domains through a 16 residues loop and is involved in the dimerisation process, thus playing an indirect role in the regulation of M^{pro} catalytic activity^{78,80}.

The catalytic site is a shallow, solvent-exposed cavity that is formed by several sub-pockets that are responsible for the recognition of various residues composing the substrate peptide sequences^{76,80}. Concerning this, particularly important is the conserved sequence Gln↓-Ser, where Gln↓ indicates the glutamine residue that precedes the cleavage site⁸¹.

Despite its peculiar structural features, which make it a difficult target for drugs, rational structure-based approaches such as Molecular Docking⁸² and Molecular Dynamics⁸³ have proven to be useful tools in the identification and characterisation of M^{pro} small molecule inhibitors, leading to the discovery of both covalent and non-covalent lead compounds^{84,85}. Further reinforcing the importance of the Main Protease as a key drug target against COVID-19, is the discovery and approval by regulatory agencies of Nirmatrelvir, the first drug specifically designed against SARS-CoV-2 to enter the market⁴³.

Due to its pivotal role in the virus replication cycle, the Main Protease is, on the contrary to Spike, particularly conserved in its primary sequence and its three-dimensional structural features among different viral strains. Taking a closer look at the main proteases from previously mentioned SARS-CoV-2 variants, only one out of 306 amino acids is mutated compared to the reference sequence, precisely residue 132, which is a proline in the case of

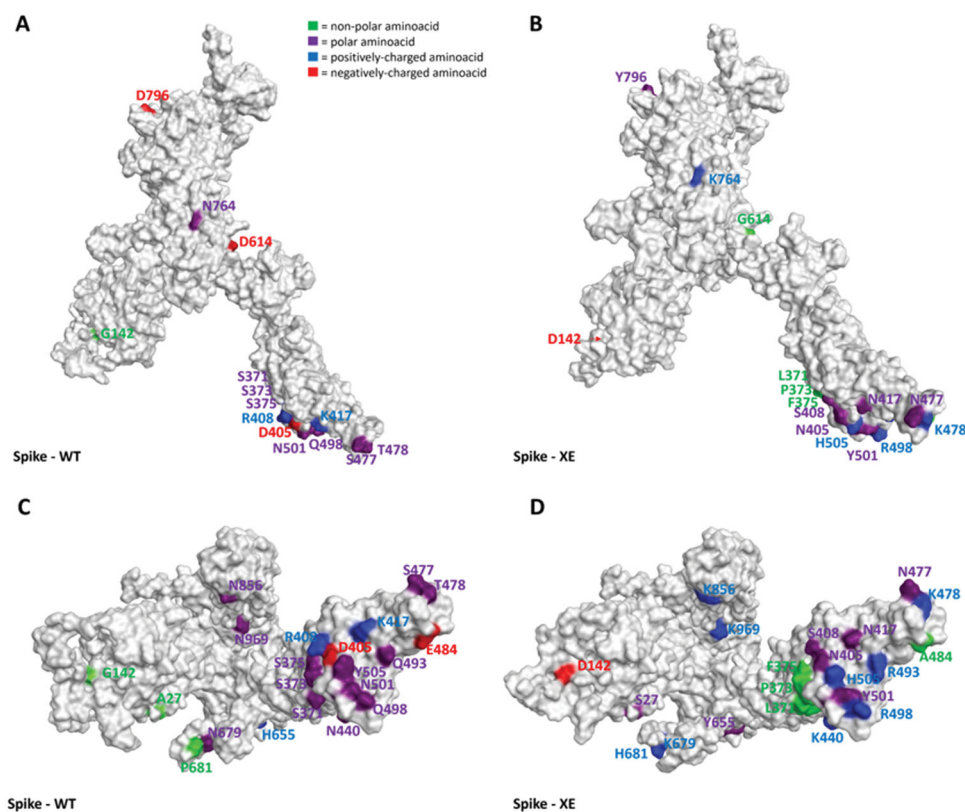


Figure 4. Representation of the differences between the WT (taken from PDB code: 6ZDH) and the XE variant of SARS-CoV-2 Spike protein. Due to the lack of experimentally resolved structure of the SARS-CoV-2 XE variant, the three-dimensional structure represented was obtained from the wild-type S protein coming from PDB code 6ZDH, and then manually mutating the residues involved in the mutations (exploiting the MOE “Protein builder” tool). Panels A and B offer a front view of the comparison between the structures, while panels C and D shift the point of view to the bottom of the proteins. To give a clearer view of the mutations, only one monomer was considered to create the image. The amino acids involved in mutations are labelled in the figure and are coloured based on their kind, following the legend reported in panel A. Specifically, Gly, Ala, Val, Leu, Ile, Pro, Cys, Met, Phe, and Trp are considered non-polar amino acids (green), Asp and Glu represent the negatively-charged amino acids (red), and Lys, Arg, and His form the positively-charged amino acid group (blue). Finally, Ser, Thr, Asn, Gln, and Tyr are all considered polar amino acids (purple). All images were created and rendered using the Molecular Operating Environment (MOE) suite.

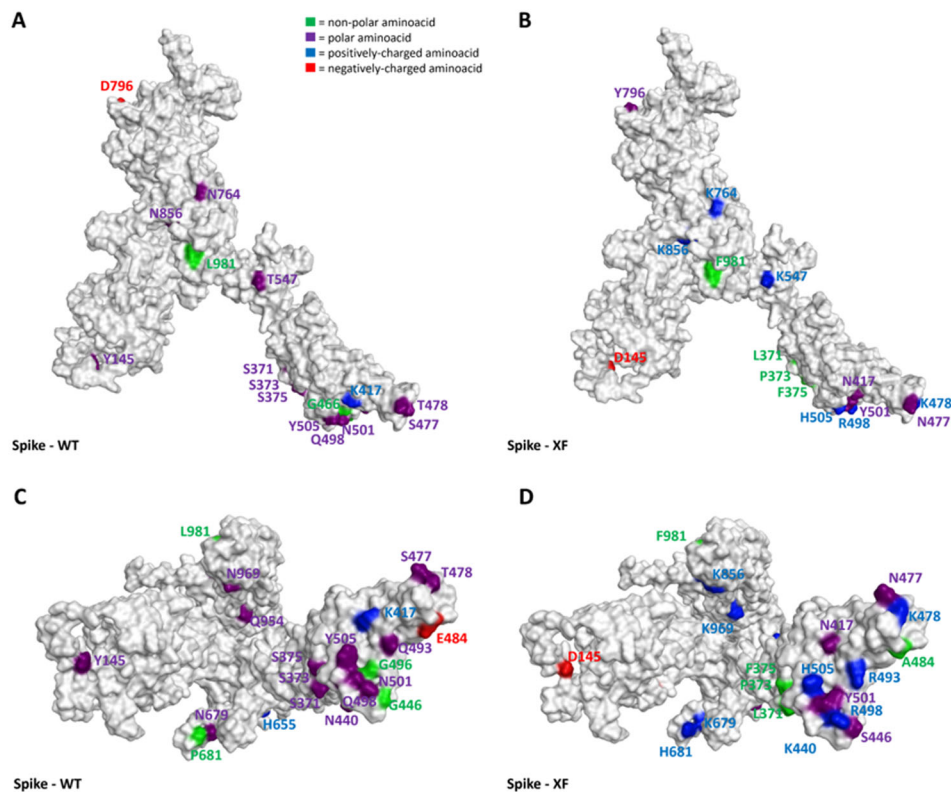


Figure 5. Representation of the differences between the WT (taken from PDB code: 6ZDH) and the XF variant of SARS-CoV-2 Spike protein. Due to the lack of experimentally resolved structure of the SARS-CoV-2 XF variant, the three-dimensional structure represented was obtained from the wild-type S protein coming from PDB code 6ZDH, and then manually mutating the residues involved in the mutations (exploiting the MOE “Protein builder” tool). Panels A and B offer a front view of the comparison between the structures, while panels C and D shift the point of view to the bottom of the proteins. To give a clearer view of the mutations, only one monomer was considered to create the image. The amino acids involved in mutations are labelled in the figure and are coloured based on their kind, following the legend reported in panel A. Specifically, Gly, Ala, Val, Leu, Ile, Pro, Cys, Met, Phe, and Trp are considered non-polar amino acids (green), Asp and Glu represent the negatively-charged aminoacids (red), and Lys, Arg, and His form the positively-charged aminoacid group (blue). Finally, Ser, Thr, Asn, Gln, and Tyr are all considered polar amino acids (purple). All images were created and rendered using the Molecular Operating Environment (MOE) suite.

Table 3. List of all the insertions and deletions operated by SARS-CoV-2 Spike protein for all the variants considered in our study (Delta, Omicron, XD, XE, and XF).

Delta variant	Omicron variant	XD variant	XE variant	XF variant
			$\Delta 24-26$	
	$\Delta 69-70$			$\Delta 69-70$ $\Delta 142-144$
	$\Delta 143-145$			
$\Delta 156-157$		$\Delta 156-157$		$\Delta 211$
	$\Delta 211$	$\Delta 211$		$\Delta 211$
	ins214EPE	ins214EPE		ins214EPE

The mutations have been aligned to give a better perspective of the ones which have been conserved through the evolutionary process.

Wuhan-Hu-1, Delta, and XD viral strains, while it is mutated to histidine in the case of Omicron, XE, and XF variants.

As can be seen in Figure 7, this mutated residue is located outside the substrate-binding site, specifically in a turn that precedes the sequence leading to the oxyanion loop (residues 138–145), which is a vital part of the catalytic machinery that is responsible for the processing of substrate peptides⁷⁶. Although the position of such mutation could suggest a possible destabilisation of the catalytic site related to an alteration of the enzymatic activity of the protease, a visual inspection of the surroundings of residue 132 suggests that this mutation should not affect in any way the stability of the three-dimensional structure of the protease, thereby not harming its ability to correctly process the substrate.

As can be seen in Figure 8, indeed, the proline residue is not involved in any intermolecular interaction relevant to the structural stability of the protease, suggesting that its only role could be limited to a joint between more relevant residues such as R131, which mediates several interactions through its sidechain guanidium group (specifically, a salt bridge with both D289 and D197, and a hydrogen bond with the backbone of T198) and its backbone (a hydrogen bond between its backbone amide proton and the amide carbonyl oxygen of T135 and another one between its carbonyl oxygen and the amide proton of F134), and N133, which is itself involved in a network of intermolecular interactions with both its backbone (hydrogen bond between its amide proton and the carboxyl oxygen of D197) and its sidechain (the amide proton donates to the carbonyl oxygen of G195 while the carbonyl oxygen receives from the hydroxyl group of T135). These structural insights are confirmed also by a functional screening performed by Flynn et al., which showed that mutations at this position, especially the P132H found in these viral variants, are generally well-tolerated, while mutations of both R131 and N133 drastically reduce or even abolish the catalytic activity of the protease⁸⁶.

Concerning the relevance of this mutation for the efficacy of existing treatments and the development of future ones, a recent study from Greasley et al. reported the crystal structure of Nirmatrelvir in complex with the main protease from three different viral variants that presented a mutation on M^{Pro}⁸⁷. The analysed mutations included the P132H, which characterises both the Omicron SARS-CoV-2 variant and the recently found XE and XF.

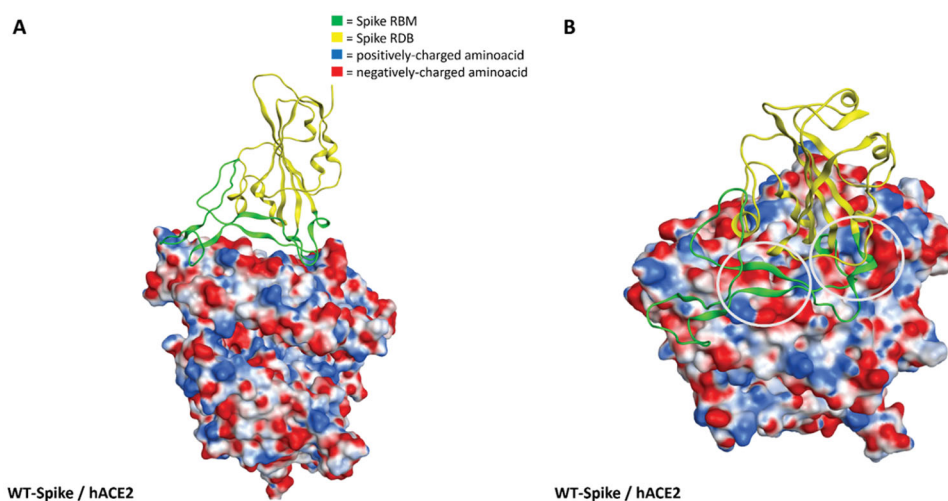


Figure 6. Representation of the interaction between WT-Spike receptor-binding domain (RBD) and hACE2 (coming from PDB code: 6M0J⁵⁸). The Spike RBD is coloured in yellow, while the receptor-binding motif (RBM) is coloured in green. The hACE2 surface is coloured according to the electrostatic properties of underlying residues (blue, positively-charged regions, red, negatively-charged regions, white, neutral regions). Panel A offers a lateral view of the complex, while panel B focuses the attention on a top-lateral perspective. As can be seen from panel B, the hACE2 regions in contact with Spike-RBM are prevalently negatively-charged (red color): concerning this, for visualisation purposes, the most extended negative regions at the Spike-hACE2 interface have also been highlighted with grey circles.

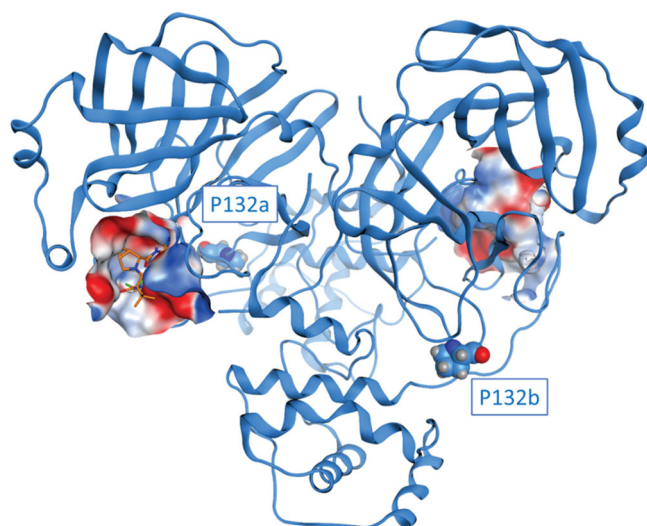


Figure 7. The structure of SARS-CoV-2 M^{Pro} (PDB ID: 6Y2E) in its free form. The protein is depicted in blue ribbons, while the mutated residue P132 in comparison with considered SARS-CoV-2 variants (Delta, Omicron, XD, XE, and XF) is highlighted and depicted as a CPK model. For visual reference, Nirmatrelvir (also known as PF-07321332, commercial name Paxlovid) from structure 7RFS is also shown in the picture, alongside the binding site surface coloured according to electrostatic properties.

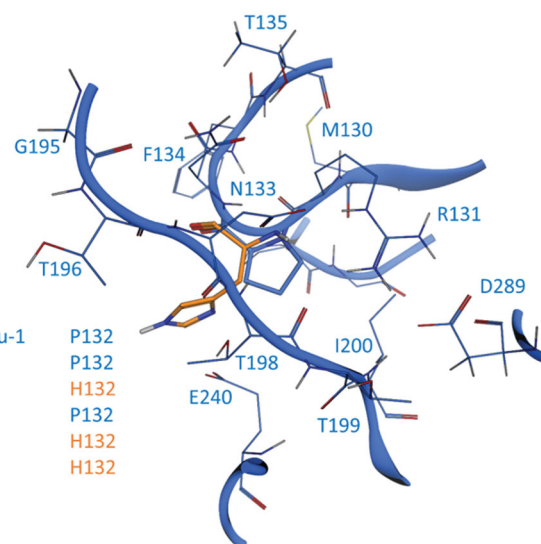


Figure 8. Comparison between SARS-CoV-2 3CL protease (M^{Pro}) from crystal structure 6Y2E (blue) and homology models of M^{Pro} from five different SARS-CoV-2 variants, reported in Table 1: the focus is on residue 132 (either proline or a histidine) of SARS-CoV-2 M^{Pro} and homology models of Delta, Omicron, XD, XE and XF variants M^{Pro}.

Greasley and collaborators established that the P132H mutation does not affect the affinity of Nirmatrelvir for the main protease catalytic site, thereby indicating the same data would be extendable also to XE and XF variants considering that they share the same P132H mutation as the Omicron variant.

As can be seen in Figure 9, although our homology model of the XE/XF variant is based on the structure 6Y2E, which represents the SARS-CoV-2 main protease in its free/unliganded form, there is an almost perfect structural superposition between our homology model and the experimentally resolved structure of the complex between the M^{Pro} from the Omicron variant and Nirmatrelvir (PDB ID: 7TLL), as is also quantitatively assessed by the 0.67 Å R.M.SD between the two structures after optimal superimposition of the backbone. The congruence between our structural prediction and the experimental data

supports the idea that the overall fold of the main protease is conserved across several variants and that the structural effect that residue mutations could have on the effectiveness of the main protease inhibitor could be accurately predicted through the combination of computational techniques such as homology modelling, molecular docking, and molecular dynamics. Moreover, based on available structural information, the high degree of structural similarity between the main proteases is not only shared by variants of the SARS-CoV-2 virus but also by other coronaviruses such as bat coronavirus¹³, the Porcine transmissible Gastroenteritis virus (TGEV)⁷⁸, Human coronavirus strain 229E (HCoV)⁸⁰, Infectious bronchitis virus (IBV)⁸⁸ and MERS-CoV⁸⁹, thereby validating the pursue of novel M^{Pro} inhibitors that could act as pan-coronaviral drugs and help to prevent future coronavirus associated pandemics.

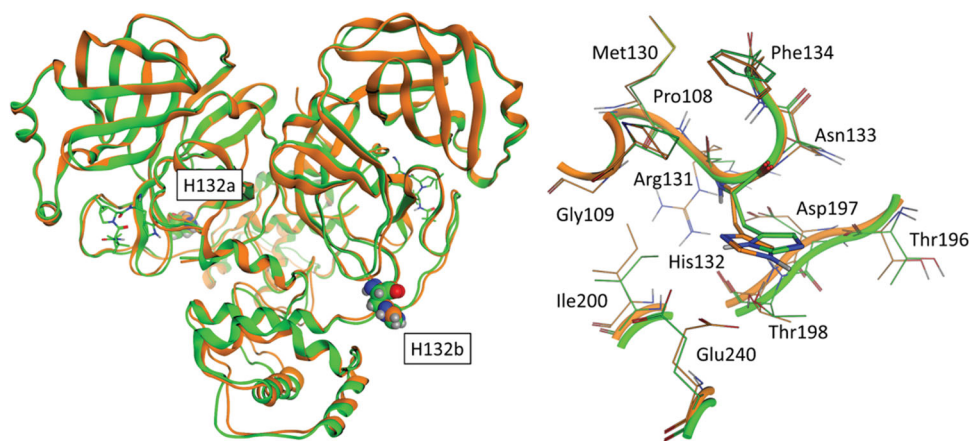


Figure 9. This panel reports the structural superposition between the crystal structure of SARS-CoV-2 Omicron variant M^{Pro} in complex with Nirmatrelvir (PDB ID: 7TLL, green) and the homology model of the XE/XF variant based on the unliganded state of the original SARS-CoV-2 virus (orange). In panel A, the whole protease structure is shown in the ribbon: for visual reference, both Nirmatrelvir and H132 are reported in liquorice and CPK models respectively. In panel B, a focused view of residue 132 and its surrounding residues is reported.

4. Conclusions

The recent emergence of novel recombinant SARS-CoV-2 variants, namely XD, XE, and XF, poses a serious threat to the efficacy of existing therapeutic options against COVID-19. In the face of a continuous evolution of the SARS-CoV-2 genome under an evolutionary pressure opposed by the development of vaccines and by the natural immunity induced by infections, the more recent viral variants have increased both their infectiveness and their ability to escape the immune response. The structural analysis reported in this article depicts a scenario where the Spike protein, which is responsible for the ability of the virus to infect human cells by interaction with the hACE2 receptor, is the viral entity that is accumulating the highest number of mutations in an attempt to increase its affinity towards the hACE2 and decrease the one towards antibodies, while the main protease M^{Pro}, a key enzyme for the virus replication cycle, is still practically identical to the wild-type virus. The different behaviour of these two proteins in response to SARS-CoV-2 genome evolution could be vital not only for the development of efficient therapies against COVID-19 but, considering the striking structural similarities between the main protease from different viruses, also in the development of pan-coronaviral drugs that could prevent the development of future coronavirus-associated pandemics.

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