Key genetic elements, single and in clusters, underlying geographically dependent SARS-CoV-2 genetic adaptation and their impact on binding affinity for drugs and immune control

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Objectives: To define key genetic elements, single or in clusters, underlying SARS-CoV-2 (severe acute respiratory syndrome coronavirus-2) evolutionary diversification across continents, and their impact on drug-binding affinity and viral antigenicity.

Methods: A total of 12150 SARS-CoV-2 sequences (publicly available) from 69 countries were analysed. Mutational clusters were assessed by hierarchical clustering. Structure-based virtual screening (SBVS) was used to select the best inhibitors of 3-chymotrypsin-like protease (3CL-Pr) and RNA-dependent RNA polymerase (RdRp) among the FDA-approved drugs and to evaluate the impact of mutations on binding affinity of these drugs. The impact of mutations on epitope recognition was predicted following Grifoni *et al.* (*Cell Host Microbe* 2020; **27**: 671–80.)

Results: Thirty-five key mutations were identified (prevalence: $\geq 0.5\%$), residing in different viral proteins. Sixteen out of 35 formed tight clusters involving multiple SARS-CoV-2 proteins, highlighting intergenic co-evolution. Some clusters (including D614G_{Spike} + P323L_{RdRp} + R203K_N + G204R_N) occurred in all continents, while others showed a geographically restricted circulation (T1198K_{PL-Pr} + P13L_N + A97V_{RdRp} in Asia, L84S_{ORF-8} + S197L_N in Europe, Y541C_{Hel} + H504C_{Hel} + L84S_{ORF-8} in America and Oceania). SBVS identified 20 best RdRp inhibitors and 21 best 3CL-Pr inhibitors belonging to different drug classes. Notably, mutations in RdRp or 3CL-Pr modulate, positively or negatively, the binding affinity of these drugs. Among them, P323L_{RdRp} (prevalence: 61.9%) reduced the binding affinity of specific compounds including remdesivir while it increased the binding affinity of the purine analogues penciclovir and tenofovir, suggesting potential hypersusceptibility. Finally, specific mutations (including Y541C_{Hel} + H504C_{Hel}) strongly hampered recognition of Class I/II epitopes, while D614G_{Spike} profoundly altered the structural stability of a recently identified B cell epitope target of neutralizing antibodies (amino acids 592–620).

Conclusions: Key genetic elements reflect geographically dependent SARS-CoV-2 genetic adaptation, and may play a potential role in modulating drug susceptibility and hampering viral antigenicity. Thus, a close monitoring of SARS-CoV-2 mutational patterns is crucial to ensure the effectiveness of treatments and vaccines worldwide.

Introduction

The new coronavirus, termed SARS-CoV-2 (severe acute respiratory syndrome coronavirus-2), emerged in China at the end of

2019.^{1,2} Afterwards, SARS-CoV-2 was declared a pandemic and has been responsible for over 16 million cases with >650 000 deaths (https://www.gisaid.org/, updated 29 July 2020), causing a global health emergency of inconceivable magnitude.^{2,3}

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SARS-CoV-2 is an enveloped positive-sense RNA virus characterized by a genome encoding four structural proteins, 16 nonstructural proteins (NSPs) and other regulatory proteins. The four structural proteins are: the envelope (E), spike (S), membrane (M) and nucleocapsid (N) protein. The 16 NSPs include the 3chymotrypsin-like protease (3CL-Pr), the papain-like protease (PL-Pr), the replication complex comprising the RNA-dependent RNA polymerase (RdRp), the helicase (Hel), the 3',5'-exonuclease (NSP-14) and other NSPs involved in the different steps of viral replication.⁴ So far, 3CL-Pr and RdRp have been explored as the main drug targets for therapeutic approaches against SARS-CoV-2 infection.⁵

Preliminary studies suggest that SARS-CoV-2 is evolving during its spread worldwide and its genome is accumulating some new variations with respect to the SARS-CoV-2 strains that originated in China.^{6,7} Nevertheless, an in-depth definition of mutational profiles underlying SARS-CoV-2 genetic diversification across geographical areas and their functional characterization has not been extensively addressed. Furthermore, given the urgency of the SARS-CoV-2 outbreak, there has been considerable interest in repurposing existing drugs approved for treating other infections or for other medical indications.⁸ Nevertheless, no information is available on the role of SARS-CoV-2 mutations in affecting, positively or negatively, the binding affinity of these drug candidates. Understanding this issue can provide important information for the development of effective antiviral agents and universal vaccines, as well as for the design of accurate diagnostic assays, thus representing a crucial aspect to consider in ongoing public health measures to contain infection worldwide.

In this light, by analysing one of the largest sets of SARS-CoV-2 sequences, this study aimed to define key genetic elements, single or in clusters, underlying the evolutionary diversification of SARS-CoV-2 across continents, and their impact on protein structural stability by molecular dynamics simulations, on binding affinity of drug candidates by docking analysis and on epitope recognition by *in silico* prediction models.

Methods

SARS-CoV-2 sequences

A total of 12 150 high-quality and nearly complete SARS-CoV-2 genomic sequences were retrieved from https://www.gisaid.org/ (see Supplementary Information available as Supplementary data at JAC Online). Sequences were obtained from samples collected between 24 December 2019 and 20 April 2020, and cover 69 countries with the following geographic distribution: Europe (N = 6680), America (N = 3274), Oceania (N = 1321), Asia (N = 777) and Africa (N = 98).

The quality filters for sequences inclusion are reported in the Supplementary Material. Sequences were aligned using the NC_045512.2 SARS-Cov-2-Wuhan-Hu-1 isolate as the reference sequence by Bioedit.

Amino acid variability of the different SARS-CoV-2 proteins

The amino acid variability in viral proteins (S, M, N, E, 3CL-Pr, PL-Pr, RdRp, Hel, NSP-14, NSP-7, NSP-8 and ORF-8) was evaluated by estimating the mean evolutionary divergence (ED) compared with the reference NC_045512.2 using the Poisson correction included in the Mega X software.

The Shannon entropy was calculated to measure the extent of amino acid variability at each position of SARS-CoV-2 protein sequences using the

formula [Sn = $-\sum_i (p_i \ln p_i)/\ln N$], where p_i was the frequency of each amino acid and N was the total number of sequences analysed.

For each protein, we assessed the number of amino acid positions with a variability ${\geq}0.5\%.$

Mutational analysis

SARS-CoV-2 mutations were defined according to the sequence of each specific protein using the NC_045512.2 SARS-Cov-2-Wuhan-Hu-1 isolate as the reference sequence. The prevalence of mutations was calculated in the overall population and according to the continent of sequence isolation. Statistically significant differences in the prevalence of mutations between Asia (continent of origin for the epidemic) and the other continents were assessed by Fisher's exact test and corrected for multiple testing by the Benjamini–Hochberg method (false discovery rate = 0.05).

Covariation analysis

Statistically significant pairs of mutations were investigated by calculating the binomial correlation coefficient (phi) for the simultaneous presence of mutations at two positions in the same isolate, while clusters of mutations were identified by average linkage hierarchical agglomerative clustering described elsewhere⁹ and in the Supplementary Material.

Phylogenetic analysis

Phylogenetic trees were performed by MEGA X^{10} using a maximum likelihood tree based on the Jukes–Cantor model,¹¹ and the bootstrap method of 1000 replicates. Phylogenetic trees were rooted and viewed using FigTree v1.4 (http://tree.bio.ed.ac.uk/software/figtree/).

Epitope localization of the identified SARS-CoV-2 mutations and predicted impact on epitope recognition

Mutations were localized in SARS-CoV-2 major histocompatibility complex (MHC) Class I/II T cell epitopes and B cell epitopes defined by Grifoni *et al.* (2020).¹² The impact of each mutation in altering the binding affinity between the epitopes and the MHC molecules was estimated *in silico* through the Immune Epitope Database and Analysis Resource (IEDB), by following the approach recently used in Grifoni *et al.* (2020)¹² and described in the Supplementary Material.

Structural analysis

Molecular dynamics simulations (described in the Supplementary Material) were performed to assess the impact of mutations on the stability of RdRp, Hel, 3CL-Pr and S proteins, with the available crystallographic models. The molecular recognition studies of 3CL-Pr and RdRp were carried out by structure-based virtual screening techniques using the DrugBank database as library (details in the Supplementary Material).

Results

Differential amino acid variability in SARS-CoV-2 proteins

The amino acid ED from the SARS-CoV-2 reference sequence (NC_045512.2) varied according to the proteins analysed and never exceeded a mean value of 0.0028 amino acid substitutions per site, indicating limited genetic divergence. The highest ED was observed in the N protein and in the ORF-8-encoded regulatory protein (mean ED \pm SD: 0.0021 \pm 0.0011 and 0.0028 \pm 0.0020 amino acid substitutions per site) (Table 1). Notably, in the N protein, the highest ED occurred in the Ser/Arg-rich motif, crucial

Protein	Amino acid length	Number of sequences	ED (±SD)ª	Number of variable amino acid positions ^b	Entropy (min-max value) ^c
Structural proteins					
Spike (S)	1-1273	9111	0.0006 (±0.0005)	3	0.04-0.67
Membrane (M)	1-222	11 760	0.0005 (±0.0003)	2	0.07-0.13
Envelope (E)	1-75	12 003	0.0002 (±0.0001)	0	0
Nucleocapsid (N)	1-419	11 380	0.0021 (±0.0011)	7	0.04-0.46
Viral enzymes					
RNA polymerase	1-932	11 185	0.0007 (±0.0006)	4	0.03-0.67
NSP-7	1-83	12087	0.0005 (±0.0004)	1	0.08
NSP-8	1-198	12024	0.0002 (±0.0000)	0	0
3′,5′-Exonuclease (NSP-14)	1-527	9526	0.0002 (±0.0001)	2	0.05-0.07
Helicase	1-601	11662	0.0007 (±0.0003)	4	0.03-0.28
3CL-protease	1-306	11918	0.0003 (±0.0001)	3	0.03-0.06
PL-protease	1-1945	9903	0.0002 (±0.0000)	6	0.04-0.1
Regulatory protein					
ORF-8	1–121	12023	0.0028 (±0.0020)	3	0.07-0.39

Table 1. Amino acid evolutionary divergence (ED) and Shannon entropy of SARS-CoV-2 main proteins

NS, non-structural; 3CL, 3-chymotrypsin-like; PL, papain-like.

^aThe amino acid ED of each SARS-CoV-2 protein compared with the reference NC_045512.2 sequence was calculated by Mega X software.

^bA position was defined as variable if amino acid substitutions were detected with a frequency \geq 0.5%.

^cShannon entropy was used to measure the amino acid variability at each position. The minimum and maximum entropy value is reported.

to mediate the interaction of the nucleocapsid with viral and cellular proteins. $^{\rm 13-15}$

Despite limited ED, Shannon entropy analysis revealed the presence of key amino acid mutations (with a frequency $\geq 0.5\%$) at 35 hot-spot positions (Table 1 and Figure 1). Mutations detected with the highest frequency were D614G_{Spike} (61.2%) and P323L_{RdRp} (61.8%), followed by R203K_N and G204R_N (17.2% and 17.2%, respectively), L84S_{ORF-8} (13%), P504L_{Hel} and Y541C_{Hel} (8.1% and 8.2%, respectively) (Figure 1). The remaining mutations occurred with a prevalence <5% (Figure 1). Among them, S193I_N, S194L_N, S197L_N and S202N_N (along with R203K_N and G204R_N) resided in the Ser/Arg-rich motif, corroborating the selective pressure acting on this domain (Figure 1). Furthermore, all the mutations observed in 3CL-Pr and PL-Pr showed a prevalence <5% (Figure 1).

Genetic diversification of SARS-CoV-2 proteins across continents

A dramatic increase (up to 60%) in the prevalence of D614G_{Spike} and P323L_{RdRp} was observed in all continents compared with Asia (adjusted $P \le 0.01$ for all comparisons), indicating that these mutations are emerging as the major circulating viral variants despite a limited initial circulation in Asia (Figure 2).

Furthermore, specific mutations circulate with higher prevalence in Europe than in other geographical regions. Among them, T175M_M, S193I_N and K90R_{3CL-Pr} showed a prevalence of 4.8%, 3.5% and 1.8%, respectively, in Europe while being absent or nearly absent in other continents (prevalence <0.9% for T175M_M, <0.1% for S193I_N and 0.3% for K90R_{3CL-Pr}, adjusted $P \le 0.05$ for all comparisons). A similar scenario was observed for R203K_N and G204R_N, showing a remarkable increase in their circulation in Europe and Oceania (adjusted P < 0.05) compared with Asia and

America (Figure 2). Similarly, $G15S_{3CL-Pr}$ showed a higher prevalence in Europe and Africa compared with the other geographic areas (adjusted P < 0.05) (Figure 2).

Interestingly, the genetic diversification in America mainly involved proteins acting as cofactors of viral polymerase. In particular, P504L_{Hel} and Y541C_{Hel} circulate predominantly in America (prevalence: 26.5% and 27%, respectively) while being rarely detected in Asia and Europe (prevalence <0.3%) and never in Africa (adjusted P< 0.05 for each comparison) (Figure 2). Similarly, S25L_{NSP-7} and A320V_{NSP-14} circulate in America with a prevalence of 5.1% and 4.3%, while their prevalence never exceeded 1.1% and 0.3%, respectively, in other geographic areas.

A peculiar situation was observed in Oceania characterized by an increased circulation of a large variety of specific mutations throughout the different viral proteins, rarely found in Asia. Among them, $G282V_{PL-Pr}$ was the only detected exclusively in Oceania (prevalence: 5.8%) and never in the other continents (Figure 2).

Covariation profiles among SARS-CoV-2 mutations across continents

Statistically significant pairs of mutations

Covariation analysis identified several statistically significant pairs involving mutations localized in different viral proteins, highlighting a process of intergenic co-evolution. In particular, among the above-mentioned 35 SARS-CoV-2 mutations, 16 were tightly associated with each other.

Specific pairs of mutations were detected in all continents, although with a diverse frequency of circulation (Table 2). This is the case of D614G_{Spik} + P323L_{RdRp} (phi from 0.7 to 0.9 in the different continents) predominantly circulating in Europe (71.9%) and Africa (83.3%), followed by America (54.6%) and Oceania (46.6%),



Figure 1. Mutation prevalence across SARS-CoV-2 proteins. The histogram reports the prevalence of mutations detected with a prevalence $\geq 0.5\%$ in the Spike protein [number (*N*) of sequences analysed = 9111)], Membrane protein (*N* = 11760), Nucleocapsid protein (*N* = 11380) (a), RNA-dependent RNA polymerase (*N* = 11185), NSP-7 (*N* = 12087), NSP-14 (*N* = 9526), Helicase (*N* = 11662) (b), 3CL-protease (*N* = 11918), PL-protease (*N* = 9903) and ORF-8-encoded protein (*N* = 12023) (c). No mutation with a prevalence $\geq 0.5\%$ was detected in the Envelope protein.



Figure 2. Distribution of mutations across continents. The histograms show the mutations with a prevalence $\geq 1\%$ in Spike, Membrane and Nucleocapsid proteins (a), in RNA-dependent RNA polymerase, NSP-7, NSP-14 and Helicase (b) and in 3CL-protease, PL-protease and ORF-8-encoded protein (c) across continents. Statistically significant differences were calculated by χ^2 test comparing each continent with Asia. The Benjamini-Hochberg method was used for correction of multiple comparisons. *P < 0.05, **P < 0.01. G282V_{PL-Pr}, although with an overall prevalence of <1%, was reported due to its peculiar geographic distribution.

Table 2. Significant pairs of mutations across continents

Mutation	Mutation prevalence, N (%)	Mutation	Mutation prevalence, N (%)	Covariation frequency ^a , N (%)	Phi ^b	Adjusted <i>P</i> value
Asia						
P3231 a la	108 (18 7)	D614Ge	89 (15 4)	79 (13.6)	07	<1 0e-20
1 JZ JERdRp	100 (10.7)	P203K	37 (6 4)	33 (5 7)	0.7	4 50-18
		G204R.	37 (6.4)	33 (5.7)	0.5	4.50-18
D614Ga	89 (15 4)	R203K.	37 (6.4)	37 (6.4)	0.5	3 50-18
DOITOSpike	05 (15.4)	G204P.	37 (6.4)	37 (6,4)	0.6	3 50-18
P203K.	37 (6 /)	G204RN	37 (6.4)	37 (0.4)	0.0	<1.0e-20
	27 (0.4)	0204KN	22 (2.0)	22 (2.0)	1.0	<1.0e-20
TIJLN	27 (4.0)	T1108K	23 (3.3)	23 (3.0)	0.5	<1.0e-20
A97V _{RdRp}	23 (3.9)	T1198K _{PL-Pr}	23 (3.9)	23 (3.9)	1	<1.0e-20 <1.0e-20
Europe						
D614Gsailea	2373 (74.3)	P3231 pdps	2318 (72.6)	2298 (71.9)	0.9	<1.0e-20
D 0 1 1 0 Spike	2070 (7 110)	R203K	771 (24.1)	771 (24.1)	0.3	<1.0e-20
		G204RN	771 (24.1)	771 (24.1)	0.3	<1.0e-20
		T175M	201 (6 2)	200 (6 3)	0.1	6 4e-13
P3231 p p-	2318 (72.6)		771 (24.1)	748 (23.9)	0.1	<1 0e-20
ГЭДЭСкакр	2510 (72.0)	G204R.	771 (24.1)	748 (23.9)	0.3	<1.0e-20
		T175M.	201 (6 2)	197 (6 3)	0.5	3 5e-11
R203K	771 (24 1)	G204R.	771 (24.1)	771 (24.1)	1.0	<1 0e-20
N205NN	//1(24.1)	T175M.	201 (6 2)	198 (6 1)	0.4	<1.00 20
G204R.	771 (24 1)	T175M _M	201 (6.2)	198 (6.1)	0.4	<1.0e-20
L84S _{ORE-8}	107 (3.3)	S197L _N	62 (1.9)	62 (1.9)	0.4	<1.0e-20
America						
D614Ge 1	1554 (60.9)	P3231 a. 15	1433 (56.1)	1393 (54.6)	0.8	<1 00-20
D614G _{Spike}	1554 (00.5)		106 (4 1)	106 (4 2)	0.0	<1.0C 20 / 0o-18
		G204P.	105 (4.1)	105 (/, 1)	0.2	6 80-18
D3231	1/33 (56 1)	P203K.	105 (4.1)	101 (/, 0)	0.2	1 10-14
FJZJLRdRp	1455 (50.1)	6203RN,	100 (4.1)	101 (4.0)	0.2	1.1e-14
1845	781 (30.6)	0204K _N	682 (26 7)	681 (36 7)	0.2	1.00-20
LO4JORF-8	701 (30.0)	VE41C	60/2 (20.7)	604 (20.7)	0.9	<1.0e-20
		1541C _{Hel} ,	094 (27.2)	094 (27.2) 20 (1 E)	0.9	<1.0e-20
		PIJJL _{PL-Pr} ,	42 (1.0)	59 (1.5) (0 (1.6)	1.7	J.2e-13
VE/1C	(0, (27, 2))	VOZLORF-8	44 (1.0)	40 (1.0)	1.7	1.5e-12
1541C _{Hel}	094 (Z7.Z) 124 (E 2)	PSU4L _{Hel}	002 (20.7)	001 (20.7)	0.9	<1.0e-20
LZDS _{NSP-7}	154 (5.2)	ASZUV _{NSP-14}	105 (4.0) 105 (7.1)	105 (4.0) 105 (7.1)	0.9	<1.0e-20
V621 or «	44 (1.6)	P1531 pt pr	42 (1.6)	39 (1.5)	0.9	<1.0e-20 <1.0e-20
Oceania	(1.0)	. 100 CFL-FI	.2 (1.0)	55 (1.5)	0.5	100 20
D61/(G	331 (/(0 //)	D3231	31/ (//6.9)	312 (46.6)	0.9	<1.00-20
DO14O _{Spike}	314 (46.0)	P202K	314(40.9)	S12 (40.0) 80 (12.3)	0.9	<1.0e-20
FJZJL _{RdRp}	514 (40.5)	C204D	93 (13.9) 02 (12 7)	09 (13.3) 90 (12.3)	0.4	-1 0o 20
19/5	1/0 (22.1)	6204K _N	92 (15.7)	09 (15.5)	0.4	<1.0e-20
L043 _{ORF-8}	140 (22.1)	VE(1C	09 (10.5) E7 (9 E)	40 (0.0) EC (0.2)	0.4	2.00-13
	1/0 (22.1)	1541C _{Hel}	57 (0.5) FF (0.5)	20 (0.3) FF (0.2)	0.6	<1.0e-20
LO43ORF-8	148 (22.1)	POU4L _{Hel} ,	55 (8.2)	>> (ð.2)	0.6	<1.0e-20
		SIY/L _N ,	49 (7.3)	49 (7.3)	0.5	<1.Ue-20
		PIDJL _{PL-Pr} ,	29 (4.3)	29 (4.3)	0.4	2.9e-16
		V62L _{ORF-8} ,	29 (4.3)	29 (4.3)	0.4	2.9e-16
D2021/	00 (40 0)	FZ33L _{NSP-14}	29 (4.3)	29 (4.3)	0.4	2.9e-16
κ2υ3κ _N	93 (13.9)	GZU4K _N	92 (13.7)	89 (13.3)	0.4	<1.0e-20

Continued

Table 2. Continued

Mutation	Mutation prevalence, N (%)	Mutation	Mutation prevalence, N (%)	Covariation frequency ^a , N (%)	Phi ^b	Adjusted <i>P</i> value
P13L _N	69 (10.3)	S197L _N ,	49 (7.3)	46 (6.8)	0.8	<1.0e-20
		A97V _{RdRp} ,	23 (3.4)	23 (3.4)	0.6	1.1e-20
		T1198K _{PL-Pr}	22 (3.2)	22 (3.2)	0.5	1.1e-19
Y541C _{Hel}	57 (8.5)	P504L _{Hel}	55 (8.2)	55 (8.2)	0.9	<1.0e-20
P153L _{PL-Pr}	29 (4.3)	F233L _{NSP-14} ,	29 (4.3)	29 (4.3)	1.0	<1.0e-20
		V62L _{ORF-8}	29 (4.3)	29 (4.3)	1.0	<1.0e-20
T1198K _{PL-Pr}	22 (3.2)	A97V _{RdRp}	23 (3.4)	22 (3.2)	1.0	<1.0e-20
Africa						
D614G _{Spike}	30 (8.3)	P323L _{RdRp}	30 (83.3)	30 (83.3)	1.0	1.9e-3
R203K _N	4 (11.1)	G204R _N	4 (11.1)	4 (11.1)	1.0	6.0e-3

^aCovariation analysis was performed using 7024 sequences stratified across continents (Asia, N = 577; Europe, N = 3192; America, N = 2550; Oceania, N = 669; Africa, N = 36). Covariation frequency was calculated as the prevalence of the pairs of mutations in the different continents.

^bStatistically significant pairs of mutations were assessed by Fisher's exact test using the Benjamini–Hochberg method for multiple comparison correction [false discovery rate (FDR) = 0.001]. For Africa, FDR = 0.01 was considered due to restricted sample size.

and limitedly in Asia (13.6%) (Table 2). Similarly, $R203K_N + G204R_N$ (phi = 1.0 except for Oceania) circulate in all continents, with the highest frequency in Europe (24.1%), followed by Oceania (13.3%), Africa (11.1%), Asia (6.4%) and America (4.1%) (Table 2).

Other specific pairs of mutations showed a preferential circulation in specific geographic areas. In particular, P504L_{Hel} + Y541C_{Hel} circulate in America and Oceania (phi = 0.9 for both) with a prevalence of 26.7% and 8.6%, respectively (Table 2). Similarly, P13L_N + A97V_{RdRp} and P13L_N + T1198K_{PL-Pr} were detected in Asia and Oceania with a prevalence ranging from 3.2% to 3.9% and from 3.2% to 7.3%, respectively (phi = 0.9 and 0.6 for P13L_N + A97V_{RdRp} and phi = 0.8 and 0.5 for P13L_N + T1198K_{PL-Pr}) (Table 2).

Again, a peculiar scenario was observed in Oceania, characterized by the circulation of multiple pairs of mutations occurring solely in this continent: $L84S_{ORF-8} + P13L_N$ (phi=0.4), P153L_{PL-Pr} + F233L_{NSP-14} (phi=1.0) and P13L_N + S197L_N (phi=0.8).

The 3CL-Pr was the only viral protein whose mutations (G15S, K90R and A266V) were not involved in statistically significant pairs.

Clusters of correlated mutations

By hierarchical clustering analysis, the pair D614G_{Spike} + P323L_{RdRp} was linked to R203K_N + G204R_N, forming a tight cluster that was detected in all continents, with the highest prevalence in Oceania (13.2%), followed by Europe (6.1%), Asia (5.7%) and America (3.9%) (bootstrap = 1.0 for all continents) (Figure 3). In Europe, this cluster D614G_{Spike} + P323L_{RdRp} + R203K_N + G204R_N was accompanied by the continent-specific mutation T175M_M (bootstrap = 1.0) (Figure 3).

Furthermore, the cluster made up of P504L_{Hel} + Y541C_{Hel} + L84S_{ORF-8} was detected in America and in Oceania (bootstrap = 1.0 and 0.97, respectively) with a prevalence of 26.7% and 8.2%, respectively.

Finally, two continent-specific clusters were identified: T1198K_{PL-Pr} + P13L_N + A97V_{RdRp} in Asia (bootstrap = 1.0) and P153L_{3CL-Pr} + V62L_{ORF-8} + F233L_{NSP-14} in Oceania (bootstrap = 1.0) (Figure 3). No continent-specific clusters of mutations were found in Africa, presumably due to the limited number of sequences available.

Phylogenetic analysis confirmed the geographically dependent clustering of mutations (Figure S1).

Functional characterization of the identified mutations on SARS-CoV-2 proteins

Predicted impact of the mutations on SARS-CoV-2 recognition by T cell- and B cell-mediated immune response

Among the 35 above-mentioned mutations, 16 resided in Class I/II-restricted T cell epitopes (Table S1). Remarkably, by *in silico* prediction, 12/16 mutations reduced the binding affinity for specific human leukocyte antigens (HLAs) compared with the WT epitope (Table 3). Importantly, a drastic drop in the binding affinity was observed for P1263L_{Spike} (score for HLA-B*07:02 of the WT versus the mutated epitope: 0.649 versus 0.001), P504L_{Hel} (score for HLA-B*07:02 of the WT versus the mutated epitope: 0.725 versus 0.001) and Y541C_{Hel} (score for HLA-A*01:01 of the WT versus the mutated epitope: 0.976 versus 0.008) (Table 3). This suggests a process of antigenic drift favouring SARS-CoV-2 escape from T cell-mediated immune responses.

Furthermore, nine mutations were localized in B cell epitopes: two in the S protein, including $D614G_{Spike}$, and seven in the N protein (Table S1). Notably, six out of seven N mutations were mapped within the same putative B cell epitope spanning positions 177–215.

Impact of mutations on the stability of structural proteins

Structural analysis was focused on mutations with a prevalence >1% localized in 3CL-Pr, RdRp, Hel and S (the only SARS-CoV-2 proteins whose crystallographic models are available).

Most of the analysed mutations determined minimal changes in the stability of the entire proteins compared with the WT (Figure S2a–d). The only exception is represented for $K90R_{3CL-Pr}$ associated with an increased stability of the entire 3CL-Pr compared with the WT (RMSD_{WT}=2.07 Å, RMSD_{K90R}=1.57 Å) (Figure S2d). Salpini et al.



Figure 3. Dendrogram of correlated mutations. The dendrogram, obtained from average linkage hierarchical agglomerative clustering, shows clusters of mutations localized in different SARS-CoV-2 proteins. The length of branches reflects distances between mutations in the original distance matrix. Bootstrap values, indicating the significance of clusters, are reported in the boxes.

Focusing in more detail on the structural localization of mutations, $P323L_{RdRp}$ is located close to the interface with NSP-8. Compared with the WT model, $P323L_{RdR}$ increased the number of total and average hydrogen bonds (HBs) between RdRp and NSP-8, supporting a more stable interaction between these two proteins (WT_{tot} = 241 075 versus P323L_{tot} = 247 839; WT_{ave} = 241 versus P323L_{ave} = 248) (Figure S3a).

Furthermore, as mentioned above, $D614G_{Spike}$ is located in a B cell epitope encompassing residues 592–620. By the root mean square fluctuation analysis, this mutation profoundly decreased

Table 3. Impact of SARS-CoV-2 mutations on bindin	g affinity between	epitopes and Class I/II-MHCs
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		CD8+ T ly	mphocytes				
Mutations	T cell epitopes ^a	HLA alleles	Score ^b (WT versus mutated)	Rank ^b (WT versus mutated)			
P1263L _{Spike}	SE P VLKGVKL	HLA-B*07:02	0.649 versus 0.001	0.12 versus 11.00			
T175M _M	ATSR T LSYYK	HLA-A*11:01	0.787 versus 0.672	0.08 versus 0.17			
S126L _{PL-Pr}	GDCEEEEFEP S TQY	HLA-B*44:03	0.742 versus 0.647	0.09 versus 0.15			
T1198K _{PL-Pr}	T KPVETSNSF	HLA-B*35:01	0.772 versus 0.537	0.10 versus 0.22			
		HLA-B*07:02	0.376 versus 0.645	0.33 versus 0.12			
P323L _{RdRp}	TVFP P TSFGPLVRK	HLA-A*03:01	0.721 versus 0.533	0.12 versus 0.29			
P504L _{Hel}	RNP AWRKAVF	HLA-B*07:02	0.725 versus 0.001	0.08 versus 9.00			
Y541C _{Hel}	TVDSSQGSE Y	HLA-A*01:01	0.976 versus 0.008	0.02 versus 8.90			
R392C _{Hel}	VVNARL R AK	HLA-A*03:01	0.773 versus 0.596	0.08 versus 0.22			
A320V _{NSP-14}	HMVVKA a l	HLA-B*08:01	0.629 versus 0.492	0.09 versus 0.19			
		CD4+ T lyr	nphocytes ^a				
Mutations		T cell epitopes ^c	3	Rank (WT versus mutated)			
Т175М _м		KEITVATSR T LSY	/K	11.00 versus 6.80			
K90R _{3CL-Pr}		QNCVLKL K VDTAN	IPK	14.00 versus 9.80			
S25L _{NSP-7}		LQQLRVES S SKLWAQ					
A320V _{NSP-14}		KVQHMVVKA A LLA	ADK	16.00 versus 19.00			
F233L _{NSP-14}		IG F DYVYNPFMIDVQ 16.00 versus 36.00					

^aEpitopes are based on T cell epitope prediction by Grifoni *et al.* (2020).¹² Letters in bold in the epitopes indicate the mutated amino acid. ^bThe score and the rank were assessed by the Immune Epitope DataBase. A reduced score and an increased rank indicate a decreased binding affinity between the epitope and the related MHC Class I/II allele.

the stability of this B cell epitope compared with the WT, suggesting an altered epitope conformation (Figure S3b).

Impact of mutations on viral susceptibility to potential inhibitors of 3CL-Pr and RdRp

Firstly, we applied an *in silico* drug-repurposing approach to select the best inhibitors of 3CL-Pr and RdRp (the main anticoronavirus pharmacological targets) among the DrugBank compounds. Then, we evaluated if the mutations P323L_{RdRp}, K90R_{3CL-Pr} and G15S_{3CL-Pr} can modulate, positively or negatively, the binding affinity between the enzyme and the inhibitor.

Based on our structure-based virtual screening (SBVS), 20 potential inhibitors of RdRp were identified: five purine analogues (including remdesivir), four cephalosporins, two acetamide derivatives, two flavone compounds, two peptide derivatives, two triazoles, an oxoazepanyl compound, a polyphenol derivative and a pyrimidine analogue (Figure 4). The majority of the identified compounds establish several HBs with specific RdRp residues (including K545, S549, K551, R553, D623 and D761) and with some RNA nucleobases, such as the uracil at position 18, adenine at position 19 and uracil at position 20 (Figure S4 to Figure S23).

As shown in Figure 5, the best ranked compound cefoperazone in the WT complex was involved in four coordination bonds with $\rm Mg^{2+}$ cations, two π -cation interactions and two HBs, while in the presence of P323L_{RdRp} the oxoazepanyl derivative RU82209 was well stabilized into the enzyme-binding pocket by means of six coordination bonds between its phosphate groups and the Mg^{2+} cations.

Notably, several candidates were better recognized in the P323L complex, as in the case of penciclovir, tenofovir, PF-00610355, zanamivir, diosmin, isavuconazole, resveratrol and, above all, RU82209, associated with the absolute best G-score value (Figure 4).

Conversely, in the presence of $P323L_{RdRp}$, all cephalosporins and both peptide derivatives showed a decreased binding affinity compared with the WT, as well as rutin and PF-03715455. Interestingly, $P323L_{RdRp}$ decreased the binding affinity of remdesivir compared with the WT (Figure 4).

Regarding 3CL-Pr, SBVS allowed selection of 21 promising inhibitors: four cephalosporins, four peptide derivatives, four purine analogues (including remdesivir), three flavone compounds, three pyrimidine analogues, two triazoles and a benzeneacetamide (Figure 6). The majority of the identified compounds establish several HBs with crucial residues of 3CL-Pr (including N142, G143, C145, E166 and T190), as well as Van der Waals contacts and a pivotal π - π stacking interaction with H41. Both C145 and H41 were involved in the 3CL-Pr catalytic dyad (Figure S24 to Figure S44). Figure 7 reports the best ranked compounds screened against the WT and mutated models.

Notably, the NAD 3-pentanone adduct, PF-00610355, PF-03715455, reproterol and rutin were found to be better recognized in both $K90R_{3CL-Pr}$ and $G15S_{3CL-Pr}$ compared with the WT, with the NAD 3-pentanone adduct being associated with the absolute best G-score in the presence of $K90R_{3CL-Pr}$ (Figures 6 and 7). This increased theoretical binding affinity could be justified by an additional salt bridge between the ligand pyridine charged nitrogen and 3CL-Pr glutamate at position 166, missing

DrugBank ID	G score WT (-11.54)*	G score P323L (-11.35)*	2D Structure	Drug Name
			Purine analogues	
DB00157	-9.69	-9.07	grand	NADH
DB00299	-8.22	-10.12		Penciclovir
DB14761	-9.91	-8.80		Remdesivir
DB12846	-8.14	-8.01		Reproterol
DB14126	-8.25	-8.45	and the second s	Tenofovir
			Cephalosporins	
DB01329	-11.54	-10.50		Cefoperazone
DB00430	-11.07	-10.11		Cefpiramide
DB01415	-10.07	-9.48	, prints	Ceftibuten

Figure 4. Binding affinity of the different drug candidates for WT and mutated RNA-dependent RNA polymerase (RdRp). This figure reports the absolute best G-score value for each analysed target, expressed as kcal/mol. The molecular recognition studies of RdRp were carried out by structure-based virtual screening techniques using the DrugBank database as library. This figure appears in colour in the online version of JAC and in black and white in the print version of JAC.

in the WT model. After reproterol recognition, in both mutated complexes, we observed a pivotal π - π stacking interaction with H41 (Figure S35), while in the best poses of both investigational PF compounds, an increased number of HBs was found (Figures S43 and S44). Furthermore, in K90R_{3CL-Pr}, the NAD 3-pentanone adduct engaged seven HBs, a salt bridge contact and a π - π stacking interaction with H41 (Figure 7).

Notably, among the protease inhibitors already used for other viral infections, $G15S_{3CL-Pr}$ determined an increased binding affinity of indinavir (Figure 6). Conversely, both $K90R_{3CL-PR}$ and $G15S_{3CL-Pr}$ strongly reduced the theoretical binding affinity of isavuconazole and sofosbuvir (Figure 6). Furthermore, $K90R_{3CL-Pr}$ determined a remarkable decrease in the binding affinity of adafosbuvir, cefpiramide and ceftibuten, while $G15S_{3CL-Pr}$ affected that of hesperidin, lisinopril and presatovir (Figure 6).

DB09050	-10.74	-9.53	not an net the second	Ceftolozane		
		A	cetamide derivatives			
DB11871	-8.23	-9.18	- Cort	PF-00610355		
DB00558	-7.90	-8.02		Zanamivir		
Flavone derivatives						
DB08995	-8.07	-8.45	ng the second	Diosmin		
DB01698	-9.58	-8.02		Rutin		
			Peptide derivatives			
DB03632	-10.65	-10.18	A A A A A A A A A A A A A A A A A A A	Argifin		
DB00722	-9.88	-8.57		Lisinopril		
			Triazole derivatives			
DB11633	-7.18	-8.02		Isavuconazole		

Figure 4. Continued

DB12138	-10.72	-9.01		PF-03715455				
	Oxoazepanyl derivatives							
DB03591	-8.75	-11.35	o o o o o o o o o o o o o o o o o o o	RU82209				
		Р	olyphenol derivatives					
DB02709	-7.33	-8.42		Resveratrol				
Pyrimidine analogues								
DB00879	-9.07	-8.61	J.	Emtricitabine				

Figure 4. Continued



Figure 5. 3D representation of the best docking pose of (a) cefoperazone (green carbon sticks) and (b) RU82209 (cyan carbon sticks) in the WT RdRp and P323L mutated models, respectively. The enzyme and the RNA are shown as a pale yellow and raspberry cartoon, respectively. The residues involved in crucial contacts with the compounds are reported as pale yellow carbon sticks and the magnesium cations are shown as pink spheres. This figure appears in colour in the online version of *JAC* and in black and white in the print version of *JAC*. In the colour version, H bond, stacking and salt bridge interactions are indicated, respectively, as yellow, light blue and violet dashed lines.

Discussion

Based on one of the largest publicly available datasets of SARS-CoV-2 sequences so far analysed (>12 000), this study identified key mutations, single or in pairs or clusters, underlying geographically dependent viral evolutionary adaptation to human hosts.^{16,17} Some of these mutations can hamper Class I/II epitope recognition or can profoundly alter the conformation of specific B cell epitopes, suggesting their capability to alter viral antigenicity. Furthermore, to our knowledge, this is the first study highlighting the capability of mutations in RdRp and 3CL-Pr to modulate, either positively or negatively, the binding affinity of specific compounds, suggesting their potential involvement in mechanisms underlying SARS-CoV-2 hypersusceptibility or resistance to drugs.

DrugBank ID	G Score WT (-8.83)*	G score G158 (-10.75)*	G score K90R (-9.63)*	2D Structure	Drug Name
			Cephalosp	porins	
DB01329	-7.82	-7.05	-7.12	THE S	Cefoperazone
DB00430	-7.66	-7.45	-6.16	july the	Cefpiramide
DB01415	-6.92	-8.09	-5.68	printy.	Ceftibuten
DB09050	-7.91	-7.05	-6.96	$ \begin{array}{c} \begin{array}{c} \begin{array}{c} \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\$	Ceftolozane
			Peptide deri	ivatives	
DB03632	-7.29	-7.39	-7.22	H H H H H H H H H H H H H H H H H H H	Argifin
DB00224	-7.47	-8.87	-6.98	at in the	Indinavir

Figure 6. Binding affinity of the different drug candidates for WT and mutated 3CL-protease. This figure reports the absolute best G-score value for each analysed target, expressed as kcal/mol. The molecular recognition studies of 3CL-protease were carried out by structure-based virtual screening techniques using the DrugBank database as library. This figure appears in colour in the online version of *JAC* and in black and white in the print version of *JAC*.

Among the identified pairs of mutations, D614G_{Spike} + P323L_{RdRp} occurred in all continents, although with a frequency ranging from 13.6% in Asia to 71.9% in Europe and 83.3% in Africa. In particular, in Europe, D614G_{Spike} was first noted in a viral strain isolated from Germany.¹⁸ Notably, D614G_{Spike} can introduce a novel protease cleavage site capable of enhancing the fusion between the viral envelope and cell membrane, thus increasing viral infectivity and, in turn, interhuman transmission potential.^{19–21} Furthermore, D614G_{Spike} lies in a recently identified B cell epitope encompassing amino acids 592–620. Molecular dynamics simulations show that D614G_{Spike} profoundly alters the stability of this epitope, supporting an altered conformation and consequently an

impaired recognition by humoral responses. This is in line with a recent finding showing the association of D614G_{Spike} with a reduced antigenicity compared with the SARS-CoV-2 strains isolated in the early epidemic phase.²¹ On this basis, D614G_{Spike} could also pose concerns for the full effectiveness of vaccine strategies under development and thus deserves further investigation in immunological studies.

 $P323L_{RdRp}$ resides in the interface domain (amino acids 250– 365) known to connect N- and C-terminal RdRp domains.²² By molecular dynamics simulations, $P323L_{RdRp}$ increased the number of HBs between RdRp and NSP-8, a cofactor which is part of the replication complex along with RdRp, suggesting a stabilized

DB00722	-6.85	-5.80	-6.15		Lisinopril			
DB02629	-8.12	-10.75	-7.98		N,N-[2,5-O-di-2- fluoro-benzyl- glucaryl]-di-[1- amino-indan-2- ol]			
			Purine ana	logues				
DB00157	-7.79	-8.05	-7.71	Jury Ho	NADH			
DB04421	-7.92	-8.77	-9.63		Nicotinamide adenine dinucleotide 3- pentanone adduct			
DB14761	-6.96	-6.40	-6.86	y xy	Remdesivir			
DB12846	-7.31	-7.83	-7.66		Reproterol			
			Flavone der	ivatives				
DB08995	-6.85	-6.36	-7.45	ng the second	Diosmin			
DB04703	-8.51	-6.76	-7.63		Hesperidin			
DB01698	-7.81	-8.89	-8.94		Rutin			
	Pyrimidine analogues							

Figure 6. Continued

DE	314906	-8.63	-8.75	-7.16		Adafosbuvir		
DE	312165	-6.72	-5.28	-7.45		Presatovir		
DE	308934	-8.83	-7.52	-7.35		Sofosbuvir		
	Triazole derivatives							
DE	311633	-6.85	-5.83	-5.51		Isavuconazole		
DE	312138	-7.20	-7.65	-8.37	4-9-9-9 19-9-9-9 19-9-9-9	PF-03715455		
				Benzeneacetamid	e derivatives			
DE	311871	-7.72	-7.98	-7.79	ografit	PF-00610355		

Figure 6. Continued



Figure 7. 3D representation of the best docking pose of (a) sofosbuvir (yellow carbon sticks), (b) *N*,*N*-[2,5-*O*-di-2-fluoro-benzyl-glucaryl]-di-[1-amino-indan-2-ol] (green carbon sticks) and (c) NAD 3-pentanone adduct (cyan carbon sticks) in the WT 3CL-Pr, G15S and K90R mutated models, respective-ly. The enzyme is shown as a slate carbon and the residues involved in crucial contacts with the compounds are reported as slate carbon sticks. This figure appears in colour in the online version of *JAC* and in black and white in the print version of *JAC*. In the colour version, H bond, stacking and salt bridge interactions are indicated, respectively, as yellow, light blue and violet dashed lines.

interaction between these two proteins. Furthermore, by *in silico* prediction, this mutation can reduce the binding affinity for specific HLAs, suggesting that P323L_{RdRp} along with D614G_{Spike} can favour viral evasion from immune responses.

Notably, by SBVS on FDA-approved drugs, $P323L_{RdRp}$ determined a reduced binding affinity for specific compounds including remdesivir. This suggests that $P323L_{RdRp}$ could act as a 'natural' drug resistance mutation, hampering the full effectiveness of specific antiviral treatments. This point deserves investigation in ongoing clinical trials.²² At the same time, $P323L_{RdRp}$ determined an increased binding affinity for specific purine analogues, including penciclovir and tenofovir, suggesting a potential viral hypersusceptibility.

Regarding 3CL-Pr, SBVS identified four cephalosporins among the best 3CL-Pr inhibitors, in line with a previous study suggesting the incorporation of a lactam ring in the lead optimization process of SARS-CoV 3CL-Pr inhibitors.²³ While K90R_{3CL-Pr} determined a reduced binding affinity for all of them, G15S_{3CL-Pr} was associated with an increased binding affinity for a specific cephalosporin, suggesting differential viral susceptibility to this drug class according to the observed mutational profile.

Among the Pr inhibitors already approved for other viral infections, lopinavir was not included in the list of the best 3CL-Pr inhibitors, due to its low binding affinity towards both WT and mutated 3CL-Pr. This is in line with recent clinical data showing no significant benefits in overall mortality and reduction of SARS-CoV-2 load.²⁴

The overall findings highlight the role of viral genetic variability in modulating, either positively or negatively, drug-binding affinity, a concept that should be taken into account in the current drugrepurposing approach.

The above-mentioned pair D614G_{Spike} + P323L_{RdRp} formed a tight cluster with R203K_N + G204R_N. This mutational pair resides in the Ser/Arg-rich motif of the N protein, known to be the target of phosphorylation by cellular kinases, to regulate the equilibrium between viral genome replication and morphogenesis²⁵ and to be involved in several intracellular signalling pathways.^{14,22}

Furthermore, R203K_N + G204R_N along with S193I_N, S194L_N, S197L_N and S202N_N reside in a region spanning amino acids 177-215 identified as a B cell epitope.¹² Thus, these mutations could alter the antigenicity of the nucleocapsid, potentially affecting its capability to elicit the production of antibodies in infected subjects.²⁶ Similarly, the peculiar enrichment of mutations in the N protein could have important implications in the serological SARS-CoV-2 diagnosis, based on the detection of antibodies against the nucleocapsid, and poses some concerns on the use of this region as a target of molecular diagnostic assays. Indeed, these mutations could contribute to the variations in the sensitivity observed among the assays used for SARS-CoV-2 diagnosis.²⁷ Further comparative studies on the performance of molecular and serological assays for SARS-CoV-2 in the presence of these mutations will be useful to clarify this issue.

Our analysis also showed that the cluster D614G_{Spike} + P323L_{RdRp} + R203K_N + G204R_N can be linked to continent-specific mutations. Indeed, in Europe, this cluster was accompanied by T175M_M. During viral morphogenesis, the M protein interacts with the N and the S protein, acting as a central organizer of viral assembly.^{25,28} Notably, both R203K and G204R reside in a stretch of amino acids (168–208) that has been proposed to be involved in

the interaction with the M protein in SARS-CoV-1.²⁵ Thus, the overall findings suggest that the association of mutations in the abovementioned proteins can promote the packaging of the encapsidated genome, thus enhancing viral morphogenesis and explaining their emergence in the viral population.

Another widely circulating cluster, characterizing 26.7% and 8.2% of American and Australian viral strains, was made up of H504C_{Hel} + Y541C_{Hel} + L84S_{ORF-8}. Notably, H504C_{Hel} and Y541C_{Hel} are the only mutations capable of abrogating the binding affinity of CD8+ T cell epitopes for some specific HLAs, supporting that these mutations can act as immune escape mutations, favouring viral evasion from CD8+ T cell responses.

In conclusion, the identification of key geographically dependent genetic elements, single or in pairs or clusters, reflects a geographically dependent viral evolutionary adaptation to human hosts. These mutations may play a potential role in modulating viral susceptibility to drugs either positively or negatively and in favouring vaccine and diagnostic escape events. For this reason, *in vitro* studies are necessary to further confirm such *in silico*-based results. In this light, a close and continuous monitoring of SARS-CoV-2 mutational patterns across different geographical areas is crucial to ensure the effectiveness of antiviral treatments and vaccines as well as the accuracy of diagnostic assays worldwide.

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Transparency declarations

None to declare.

Supplementary data

Supplementary Information, Supplementary Material, Figures S1 to S44 and Table S1 are available as Supplementary data at JAC Online.

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