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### Probing SARS-CoV-2 sequence diversity of Pakistani isolates

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To the Editor,

With the increasing spread of COVID-19 pandemic around the world, there have been 223,983 whole genome sequences of SARS-CoV-2 submitted in GISAID database as of November 2, 2020. This wealth of sequences can be useful in probing variations in the viral genome which can potentially affect its transmissibility and virulence. The SARS-CoV-2 is a RNA virus constituting six major open reading frames (ORF) that encodes structural and non-structural proteins. Sixteen non-structural proteins (nsp 1-16) are encoded by ORF1a and 1b while the accessory genes are encoded by ORF3a, ORF6, ORF7a and b, and ORF8 (Shimamoto et al., 2015; Zhou et al., 2020). From a comparative standpoint, RNA viruses (like influenza and HIV) tend to incorporate nucleotide variations due to the lack of proof reading activity of RNA polymerase enzyme. Logically, this can bring about high mutation rate, however SARS-CoV viruses has evolved with a proof reading region, the nsp14, that keeps a check on rapid mutational changes in its genome (Denison et al., 2011). Numerically, this has been exemplified from reports which have suggested 12,000 mutations in SARS CoV2 mutations till September 2020 (Callaway, 2020).

Pakistan is a populous country with 403,311 positive cases and 166 deaths as of December 1, 2020 (https://covid.gov.pk/stats/pakistan). Therefore, following genomic surveillance for SARS-CoV-2 is imperative. As of October 16, 2020, only 14 whole genome sequences of SARS-CoV-2 has been reported from Pakistan. Nevertheless, studying these sequences with respect to its divergence from worldwide sequences, is crucial to get a lead on the possible genetic variants of SARS-CoV-2 which might be circulating in Pakistani population.

All the 14 whole genome sequences of SARS-CoV-2 reported from Pakistan till October 16, 2020 were downloaded from GISAID database (https://www.epicov.org/epi3/frontend#efa72), and the first sequence SARS-CoV-2 from Wuhan was used as reference sequence (Accession number: NC\_045512.2). The multiple sequence alignment was performed using Clustal X (Larkin et al., 2007). Visualization of alignment followed by mutational analysis was performed using Jalview (Waterhouse et al., 2009). Phylogenetic analysis of 14 Pakistani sequence isolates of SARS-CoV-2, was performed using Galaxy server. For phylogenetic analysis multiple sequence alignment was performed using MAFFT followed by Maximum Likelihood tree construction using IQTree available on Galaxy server (Nguyen et al., 2015). The

visualization and editing of tree was performed using FigTree (http://tree.bio.ed.ac.uk/software/figtree/). The variations in the amino acid sequences has been compared with the SARS-CoV-2 sequences reported around the world on COVIDCG database (Chen et al., 2020) as of October 16, 2020. The effect of mutations on stability of protein was studied through I-Mutant 3.0. It predicts the effect of mutation on stability of protein by estimation of Gibbs free change ( $\Delta G$ ) (difference of energy (DDG) between native and mutated protein). The effect of mutations on the stability of protein is classified either increasing the stability (DDG > 0.5Kcal/mol), decreasing the stability (DDG < -0.5 Kcal/mol) or neutral impact (-0.5  $\leq$ DDG  $\leq$ 0.5 Kcal/mol) on protein structure (Capriotti et al., 2005).

Phylogenetic analysis revealed that the SARS-CoV-2 sequences correspond to GH, S, O, GR and L clades circulating in Pakistan. Initial sequences (March 2020) from Pakistan revealed the presence of L and O clades. The L clade sequences appeared to be closely related to strains reported from United Kingdom and United Arab Emirates, while O clade sequences were closely related to SARS-CoV-2 sequences from Japan. The samples collected in May 2020 belongs to GR clade that is closely related to isolates from USA, Sweden and Germany. The GH and S clade were observed in June, 2020 sequences that appear to be closely related to strains reported from United Arab Emirates (Fig. S1).

In total, 28 amino acid variations in the structural and nonstructural proteins of SARS-CoV-2 have been identified from patient isolates across Pakistan. There are 07 non-structural genes (nsp1, 7–11, 14–16) in ORF1ab that have been found to be conserved in Pakistani isolates. The amino acid changes have been observed in nsp2, 3, 4, 5,6, 12, and 13 (Fig. 1).

In nsp2, three changes (R207C, V378I, D448N) have been observed in the sequences collected from March and one change (L450F) have been observed in the samples from June. Interestingly, these changes have not been observed in any of the sequences reported around the world. Nsp2 is an important viral protein that along with nsp8, is involved in viral replication (Angeletti et al., 2020). Hence, any change in this gene may impair viral replication and therefore requires further investigation.

In nsp3, three changes (L944S, T1246I, K1305N, and Q2702H) have been detected in the isolates from May 2020. The T1246I variant has been reported in only 0.2% sequences from different countries (Table 1). The K1305N has been observed in only 0.1% of the Asian sequences and

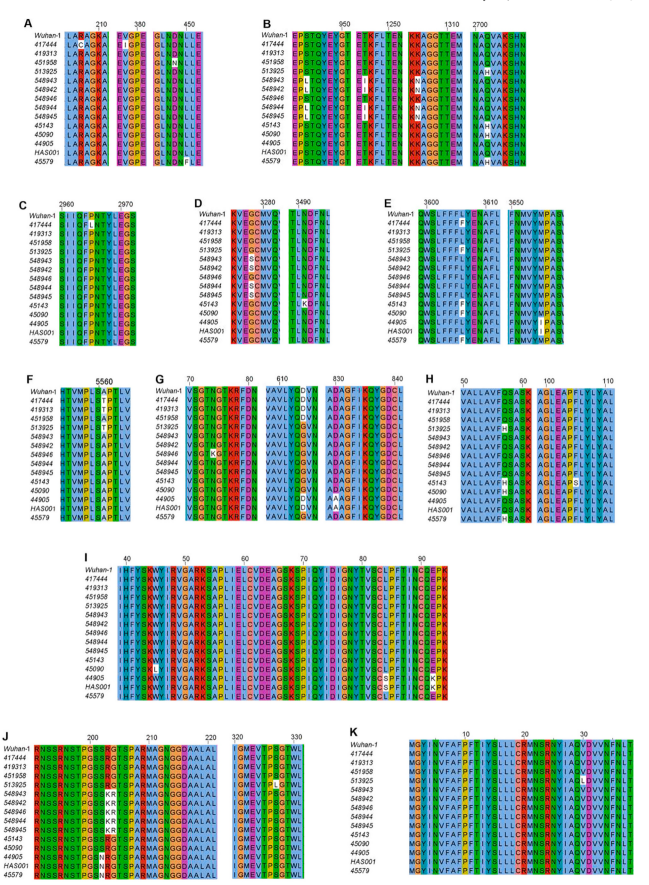


Fig. 1. Multiple sequence alignment of Pakistani SARA-CoV-2 sequences with the reference Wuhan-1 strain. (A) nsp2; (B) nsp3; (C) nsp4; (D) nsp5; (E) nsp6; (F) nsp13; (G) Spike; (H) ORF3; (I) ORF8; (J) Nucleocapsid; (K) ORF10. The amino acid variations are shown with white color.

Table 1
Details of mutations reported in Pakistani Sequences in comparison with the worldwide reported sequences and effect of mutations on stability of protein. DDG > 0.5Kcal/mol = protein stability increases, DDG < -0.5 Kcal/mol = protein stability decreases.

Sample ID	Gene	Mutation	Worldwide Prevalence	DDG (KJ/mol)	Effect on Protein Stability
EPI_ISL_417444	nsp2	R207C V378I		-0.82654688 -0.47956396	Decrease
EPI ISL 451958	nsp2	D448N		-0.91305103	Decrease
EPI_ISL_45579	nsp2	L450F		-1.6603629	Decrease
EPI ISL 548942	nsp3	L944S		1,0000023	Decrease
EPI ISL 548943	поро	T1246I	Asia = $0.2\%$	-0.61244224	Decrease
EPI_ISL_548944			Europe = 2.5%	-0.16539386	
EPI_ISL_548945			South America = 7.6%		
El 1,602,540,745			USA = 0.1%		
			Canada = $0.2\%$		
			Africa = 9.5%		
		K1305N	Asia = $0.1\%$		
		11100011	Europe = $0.6\%$		
			Africa = $0.04\%$		
EPI ISL 513925	nsp3	Q2702H	Asia = $0.5\%$	-1.4611661	Decrease
EPI ISL 45143	поро	Q27 0211	Europe = 0.9%	11/011001	Decrease
EPI ISL 45090			Canada = $0.2\%$		
EPI_ISL_45579			Africa = $1.7\%$		
EPI_ISL_417444	nsp4	P2965L	7111Ca = 1.7 70	0.433947	Increase
EPI ISL 548942	nsp5	G3278S	Asia=0.2%	-0.39907465	Decrease
EPI ISL 548943	поро	G027 00	Europe = 6.4%	0.05507 100	Decrease
EPI ISL 548944			South America = 7.6%		
EPI ISL 548945			Canada = $0.4\%$		
EF1_I3L_346943			Africa = $9.6\%$		
EDI 101 460160		NO 401 IZ	Airica = 9.0%	1 2556076	Decrease
EPI_ISL_468160	nsp5	N3491K	Asia 21.00/	-1.3556076	
EPI_ISL_417444	nsp6	L3606F	Asia = $21.0\%$	-1.2923268	Decrease
EPI_ISL_513925			Europe = 9.6%		
EPI_ISL_45143			South America = 2.6%		
EPI_ISL_45090			USA = 3.3%		
EPI_ISL_45579			Canada = 6.2%		
			Africa = 5.1%		_
EPI_ISL_468159	nsp6	M3655I	Asia = 0.6%	-0.74222727	Decrease
EPI_ISL_468163			Europe = $0.8\%$		
			Africa = 1.1%		
EPI_ISL_548942	nsp12	P4715L	Asia=62.7%		
EPI_ISL_548943			Europe = 90%		
EPI_ISL_548944			South America = 94%		
EPI_ISL_548945			USA = 89%		
EPI_ISL_513925			Canada = 85%		
EPI_ISL_468161			Africa = 91%		
EPI_ISL_548946					
EPI_ISL_468160					
EPI_ISL_468162					
EPI_ISL_513925	nsp13	A5561T		-1.8,647,808	Decrease
EPI_ISL_417444					
EPI_ISL_419313					
EPI_ISL_548946	S	N74K		0.82838446	Increase
EPI_ISL_548946	S	D614G	Asia = 62.4%	-1.4867818	
EPI_ISL_468160			Europe = 87.9%		
EPI_ISL_468161			South America = 94.3%		
EPI_ISL_548942			USA = 89.2%		
EPI_ISL_548943			Canada = 82.2%		
EPI ISL 548944			Africa = 95%		
EPI_ISL_548945					
EPI ISL 513925					
EPI ISL 468162					
EPI_ISL_468159	S	D830A		-0.59789075	Decrease
EPI ISL 468163					
EPI ISL 513925	ORF3a	Q57H	Asia = $25.2\%$	-0.91817829	Decrease
EPI_ISL_468161		Ç	Europe = 10.8%	***************************************	
EPI ISL 468160 EPI ISL 468162			South America = 12.0%		
El 1_102_100100 El 1_103_100102			USA = 62.0%		
			Canada = 36.6%		
			Africa = 9.3%		
EPI ISL 468160	ORF3a	F105S		-0.94734895	Decrease
	ORF8	W45L	Europe = $0.1\%$	-0.57599897	Decrease
EPI_ISL_468161	OKF8	vv43L	-	-u.5/59989/	Decrease
EDI ICI 4601E0	ODEO	1040	USA = 0.2%	1.0052577	Dograssa
EPI_ISL_468159	ORF8	L84S	Asia = $8.3\%$	-1.9053577	Decrease
EPI_ISL_468163			Europe = 2.2%		
			South America = 3.7%		
			USA = 8.8%		
			Canada = 12.8%		
			Africa = $4.2\%$		

(continued on next page)

Table 1 (continued)

Sample ID	Gene	Mutation	Worldwide Prevalence	DDG (KJ/mol)	Effect on Protein Stability
		E92K	Asia = 0.4%	-0.75628271	
			Africa $= 1.7\%$		
EPI_ISL_468159	N	S202N	Asia=2.7%	0.6542265	Increase
EPI_ISL_468163			Europe = 0.1%		
EPI_ISL_468162			South America = 0.0%		
			USA = 0.3%		
			Africa $= 4.2\%$		
EPI_ISL_548942	N	R203K	Asia=29.3%	-1.0587625	Decrease
EPI_ISL_548943		G204R	Europe = 46.7%		
EPI_ISL_548944			South America = 62.7%		
EPI_ISL_548945			USA = 12.9%		
EPI_ISL_548946			Canada = 16.9%		
			Africa $= 53.7\%$		
EPI_ISL_513925	N	S327L	Asia $= 0.1\%$	0.55558303	Increase
			Europe = 0.1%		
			South America $= 0.1\%$		
			USA = 0.1%		
			Africa $= 0.3\%$		
EPI_ISL_513925	ORF10	V30L	Asia=0.2%	-1.4376902	Decrease
			Europe $= 6.9\%$		
			USA = 0.1%		
			Africa $= 0.2\%$		

0.5% in European isolates. The Q2702H have been observed from 5 sequences recovered in May and June 2020. Comparatively worldwide, this mutation has been reported in less than 1% of the isolates. Another mutation that has been observed in nsp3 is T2016K which has been reported earlier by Ghanchi et al. 2020 (Ghanchi et al., 2020). This mutation has been reported in 15% of the isolates from Asia and found to be prevailing in October 2020 isolates as well. Functionally, the nsp3 plays its part in immunosuppression of innate immune responses of host (Lei et al., 2018). Hence, the changes in nsp3 can result in enhanced viral capability to evade innate immune defenses.

In nsp4, only one novel change (P2965L) has been observed in one isolate. This change has not been found in any of the worldwide reported sequences to our knowledge.

In nsp5, the G3278S and N3491K changes have been observed from the isolates of May and June, respectively. The N3491K appears to be a novel mutation while G3278 has been reported with high prevalence rate (9.6%) in the initial months of pandemic and has not been observed after August 2020. The nsp5 is also known as 3C like protease, is involved in viral replication (Macchiagodena et al., 2020) and also interferes with interferon signaling (Zhu et al., 2017a; Zhu et al., 2017b). Hence, potentially any mutation can impact viral replication capability.

In nsp6, the L3606F change has been present in 5 isolates from March, and June 2020. This mutation has been shown in 21% of isolates from Asia while in 9.6% of isolates from Europe. The nsp6 protein of other coronaviruses has been reported to interfere cellular autophagy signaling by affecting the PI3K3C3 and ATG5 proteins thereby inducing autophagosome formation but blocking its maturation (Benvenuto et al., 2020; Gassen et al., 2019; Yang and Shen, 2020).

Among the structural proteins of SARS-CoV-2, the spike protein (S) is the outermost protein that is involved in entry of virus into the host cell. The D614G change in the S protein is observed in 9 Pakistani isolates from May and June 2020. The D614G mutation has been considered to increase SARS-CoV-2 infectious capability (Korber et al., 2020) by increasing the transmissibility of virus. The D830A is another novel change observed in 2 Pakistani isolates from June 2020. This change is important as D830 is present near the TMPRSS2 binding site and may have an effect on viral fusion.

In ORF3a, the important change has been observed at position Q57H from the isolates of May and June 2020. It has been reported that this mutation is prevalent worldwide and it has an impact on protein structure (Banoun, 2020). The Q57H is present in 62% of the isolates from USA, 25% of isolates from Asia and 10.8% of isolates from Europe. ORF3a and ORF8 may have a role in host immune responses (Banoun,

2020).

Three changes (W45L, L84S, and E92K) has been observed in ORF8 from the isolates of May and June 2020. The L84S has been present in Canada (12.8% of isolates), USA (8.8% of isolates) and Asia (8.3% of isolates). The E92K has been reported in less than 0.5% of worldwide isolates and not observed after July 2020. The ORF8 is important in downregulating the MHC-I molecules thus protecting the affected cells from cytotoxic T-cell killing (Banoun, 2020).

In the N gene the important mutations have been S202N, R203K, G204R, S327L observed in Pakistani isolates. In comparison with the worldwide reported sequences, it is observed that in Asian and African sequences the S202N is present in 2.7% and 4.2% of isolates, respectively but it has disappeared after July 2020. The R203K and G204R are the changes that co-exist in the isolates while observing the worldwide reported sequences. These two mutations have also been among one of the prevalent changes observed in SARS-CoV-2 genome present in 62% of isolates in America, 53% of isolates in Africa, 46% of isolates in Europe and 29% of isolates in Asia. The N protein -contribute to viral genome assembly and serve as viral suppressor of RNAi to antagonize the host immune defense system (Liu et al., 2020).

In ORF10, only one change the V30L have been observed in one of the isolate from Pakistan. The V30L have not been observed in sequences from USA and Asia after August, 2020 while in European sequences this change is still appearing. Since ORF10 is a novel protein to SARS-CoV-2, not much data available about the role of this protein in viral pathogenesis (Cagliani et al., 2020).

By analyzing the impact of mutations on the stability of protein structure through I-Mutant, all the reported variations have been suggesting a decreasing stability of protein structure with the exception of P2965L, N75K, S202N, and S327L variations which suggest an impression of increasing the stability of nsp4, S, and N protein respectively (Table1).

Pakistan is now experiencing second wave of COVID-19 insurgence. Therefore, more SARS-CoV-2 sequences are required for effective genomic surveillance of SARS-CoV-2 and for identifying sequence divergence from Pakistan. The novel mutations in the nsp2, nsp4, nsp5, nsp13, S, and ORF3a should be further evaluated in more SARS-CoV-2 genomes from Pakistan. The novel mutation in nsp4 (P2965L) and similar variants that have been reported around the world should be further investigated for its impact on protein stability as it may affect the viral counter measures in developing efficient vaccines and therapeutic solutions.

#### Data availability

The annotated genomes of SARS-CoV-2 from Pakistan and the sequences used for phylogenetic analysis has been retrieved from the global initiative on sharing all influenza data (GISAID) (https://www.gisaid.org/). A full list of accession number along with the acknowledgment table is provided as supplementary file 1.

#### **Declaration of Competing Interest**

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

The authors declare the following financial interests/personal relationships which may be considered as potential competing interests:

## Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.meegid.2021.104752.

#### References

- Angeletti, S., et al., 2020. COVID-2019: the role of the nsp2 and nsp3 in its pathogenesis. J. Med. Virol. 92 (6), 584–588.
- Banoun, H., 2020. Evolution of Sars-Cov-2: Update September 2020. SSRN.
- Benvenuto, D., et al., 2020. Evolutionary analysis of SARS-CoV-2: how mutation of nonstructural protein 6 (NSP6) could affect viral autophagy. J. Inf. Secur. 81 (1), e24–e27.
- Cagliani, R., et al., 2020. Coding potential and sequence conservation of SARS-CoV-2 and related animal viruses. Infect. Genet. Evol. 83, 104353.
- Callaway, E., 2020. The coronavirus is mutating does it matter? Nature 585 (7824), 174–177.
- Capriotti, E., Fariselli, P., Casadio, R., 2005. I-Mutant2.0: predicting stability changes upon mutation from the protein sequence or structure. Nucleic Acids Res 33 (Web Server issue): p. W306–10.

- Chen, A.T., et al., 2020. COVID-19 CG: Tracking SARS-CoV-2 mutations by locations and dates of interest. bioRxiv.
- Denison, M.R., et al., 2011. Coronaviruses: an RNA proofreading machine regulates replication fidelity and diversity. RNA Biol. 8 (2), 270–279.
- Gassen, N.C., et al., 2019. SKP2 attenuates autophagy through Beclin1-ubiquitination and its inhibition reduces MERS-coronavirus infection. Nat. Commun. 10 (1), 5770.
- Ghanchi, N.K., et al., 2020. SARS-CoV-2 genome analysis of strains in Pakistan reveals GH, S and L clade strains at the start of the pandemic, 2020.08.04.234153.
- Korber, B., et al., 2020. Tracking changes in SARS-CoV-2 spike: evidence that D614G increases infectivity of the COVID-19 virus. Cell 182 (4), 812–827 (e19).
- Larkin, M.A., et al., 2007. Clustal W and Clustal X version 2.0. Bioinformatics 23 (21), 2947–2948.
  Lei, J., Kusov, Y., Hilgenfeld, R., 2018. Nsp3 of coronaviruses: structures and functions of
- a large multi-domain protein. Antivir. Res. 149, 58–74. Liu, Q., et al., 2020. Ongoing natural selection drives the evolution of SARS-CoV-2 ge-
- nomes, 2020.09.07.20189860.
- Macchiagodena, M., Pagliai, M., Procacci, P., 2020. Identification of potential binders of the main protease 3CL(pro) of the COVID-19 via structure-based ligand design and molecular modeling. Chem. Phys. Lett. 750, 137489.
- Nguyen, L.T., et al., 2015. IQ-TREE: a fast and effective stochastic algorithm for estimating maximum-likelihood phylogenies. Mol. Biol. Evol. 32 (1), 268–274.
- Shimamoto, Y., et al., 2015. Fused-ring structure of decahydroisoquinolin as a novel scaffold for SARS 3CL protease inhibitors. Bioorg. Med. Chem. 23 (4), 876–890.
- Waterhouse, A.M., et al., 2009. Jalview version 2-a multiple sequence alignment editor and analysis workbench. Bioinformatics 25 (9), 1189-1191.
- Yang, N., Shen, H.M., 2020. Targeting the endocytic pathway and autophagy process as a novel therapeutic strategy in COVID-19. Int. J. Biol. Sci. 16 (10), 1724–1731.
- Zhou, P., et al., 2020. A pneumonia outbreak associated with a new coronavirus of probable bat origin. Nature 579 (7798), 270–273.
- Zhu, X., et al., 2017a. Porcine deltacoronavirus nsp5 inhibits interferon-beta production through the cleavage of NEMO. Virology 502, 33–38.
- Zhu, X., et al., Porcine Deltacoronavirus nsp5 Antagonizes Type I Interferon Signaling by Cleaving STAT2. J. Virol., 2017b. 91(10).

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