Molecular Analysis and Genome Sequencing of SARS-CoV-2 during Second Wave 2021 Revealed Variant Diversity in India

Rupinder Bakshi*, Satinder Kaur, Karashdeep Kaur, Ramanpreet Kaur, Jaspreet Kaur Boparai, Ritika Ghai, Tanveer Kaur, Amritpal Kaur, Jaspreet Kaur, Kajal Verma, Palika Sharma, Gagandeep Singh, Sweety, Vikram Thakur, Kulwinder Singh, Savita Rani, Paramjeet Kaur, Sukhvir Kaur, Suman, Navdeep Kaur, Naina Rai, Dhavni Singla, Akshdeep Singh, Balwant Singh, Sukhpal Singh and Harbhajan Singh

Viral Research and Diagnostic Laboratory, Government Medical College, Patiala - 147 001, Punjab, India.

Abstract

SARS-CoV-2 variants rapid emergence has posed critical challenge of higher transmission and immune escape causing serious threats to control the pandemic. The present study was carried out in confirmed cases of SARS-CoV-2 patients to elucidate the prevalence of SARS-CoV-2 variant strain. We performed RT-PCR using extracted RNA from the nasopharyngeal swabs of suspected Covid-19 patients. Confirmed positive cases with CT<25 were subjected to whole-genome sequencing to track the prevalence of the virus in the Malwa region of Punjab. The presence of B.1.1.7, B.1.351, B.1.617.1, B.1.617.2, AY.1 and other unidentified variants of SARS-CoV-2 was found in the studied population. Among all the variants, B.1.1.7 (UK variant) and B.1.617.2 (delta-Indian variant) was found to be the most dominant variant in the population and was found majorly in Patiala followed by Ludhiana, SBS Nagar, Mansa and Sangrur. In addition to this, sequencing results also observed that the dominant trait was more prevalent in male population and age group 21-40 years. The B.1.1.7 and B.1.617.2 variant of SARS-CoV-2 is replacing the wild type (Wuhan Strain) and emerging as the dominant variant in Punjab.

Keywords: SARS-CoV-2, RT-PCR, whole genome sequencing, UK variant, Delta variant, Punjab

*Correspondence: rupindergill1@yahoo.co.in; +91-9815320300

(Received: September 04, 2021; accepted: September 23, 2021)
INTRODUCTION
Severe acute respiratory syndrome coronavirus-2 (SARS-CoV-2), earlier known as novel CoV/ nCoV lead to coronavirus pandemic in 2019. SARS-CoV-2 belongs to Coronaviridae family having single stranded ‘+’ sense RNA causing infection in mainly humans and animals resulting in respiratory congestions. The first reported case of this virus was identified in Wuhan Hubei, China during December 2019 and spread hastily all around the world. The beginning of the outbreak in India began in January 2020, one of the medical student carrying Wuhan strain returned from Wuhan University reported positive in Kerala on 30th January 2020. In Punjab it appeared during the month of March 2020 with 42 active cases. Later Indian health ministry initiated the surveillance of SARS-CoV-2 with ICMR (Indian Council of Medical Research) through VRDL (Viral Research Diagnostics Laboratories).

Initially, there were seven reported species of coronavirus such as HCoV-229E, HKU1, NL63, OC43, SARS-CoV, MERS-CoV and SARS-CoV-2 of the coronavirus leading to mild and severe respiratory problems. Among these, HCoV-229E, HKU1, NL63, OC43 causes mild symptoms while these species SARS-CoV, MERS-CoV and SARS-CoV-2 are reported to cause fatal respiratory problems. The viral RNA has the potential of getting mutated repeatedly so it is very crucial to understand the nature of mutation in different strains of SARS-CoV-2. Scientists observed various variants of SARS-CoV-2 from different geographical regions of the world. These variants have reported to show insertions, deletions or substitutions which might be beneficial or detrimental to the organism due to changes in the viral structural and pathogenic properties. Understanding of the mutation will help to estimate viral transmission, immune escape, efficiency of replication, and virus virulence. As India is the second largest infected country of SARS-CoV-2, the current study focused on Indian isolates (Punjab) to explore the variations occurring in the population. Therefore, taking this into consideration we analyzed the current trend of SARS-CoV-2 emergence in India.

METHODOLOGY
Study Area
Punjab state is geographically located in North-West India. Post-partitioned Punjab is divided into three main regions: Majha, Malwa and Doaba. The study area of our research was Malwa region, which is situated in south of river Sutlej. The present work was carried out by studying 10,22,820 symptomatic and asymptomatic individuals during second wave of Covid-19 surge from January to May 2021. The research was carried out from 5 districts of Malwa region in Punjab such as Sangrur, Mansa, Ludhiana, Patiala, and SBS Nagar. Personal demographic data such as name, age, gender, geographical location, identity proof, contact details etc. was gathered from individuals infected by Covid-19 as per the guidelines given by ICMR.

Sample Handling
Respiratory specimens i.e., nasopharyngeal swabs in universal viral transport media were received from 5 different districts of Malwa region at Viral Research and Diagnostic Laboratory (VRDL), tertiary care hospital, Punjab. The received samples were further processed for RNA extraction and RT PCR. Then the remaining samples were stored at -80°C.

RNA Extraction
RNA was extracted from the nasopharyngeal swabs using 200 µL of the sample. Nucleic acid extraction was conducted in 96 well plates using MagMAX™ viral pathogen nucleic acid extraction kit. RNA was eluted in 50 µL of elution buffer. The extracted genetic material was downstream for molecular detection.

RT-PCR and Sequencing
The Real Time RT-PCR Test was used for the qualitative recognition of SARS-CoV-2 nucleic acid from samples collected during January 2021 to May 2021. The Real Time-PCR test incorporates reverse transcription of viral RNA into DNA for the easy detection of the virus. A total of 9µL of the extracted RNA was subjected to RT PCR for the qualitative detection of SARS-CoV-2 RNA using Genes2Me (VIRALDTECT-II) Multiples real Time PCR kit. The PCR reaction was conducted using following thermal conditions heated to 55°C for
10 minutes for reverse transcription, denatured at 95°C for 3 minutes and then 40 cycles of amplification were carried out at 95°C for 15 seconds and 60°C for 60 seconds using reporter dyes Cy5 for N gene, Rox for RdRp gene and FAM for E gene. The results were analyzed and few positive samples with cycle threshold of <25 were packed in dry ice with triple-layer packaging and sent to National Centre for Disease Control; New Delhi (NCDC) for whole genome sequencing. We analyzed lineage according to age, gender and geographic spread in five districts using data provided by NCDC, New Delhi.

RESULTS

A serial cross-sectional study was conducted by Tertiary Care Hospital, Malwa region. In the present study 10,22,820 suspected individuals were tested for SARS-CoV-2 during the period from January 2021 to May 2021. Among these 63,293 individuals were confirmed positive by RT-PCR. Trends in the distribution of positivity as compared to total sample in second wave during January 2021 to May 2021 have been depicted in the Fig. 1. Out of these positive samples 1800 sample with cycle threshold <25 were randomly selected for whole genome sequencing as per directions given by NCDC. Among these sequencing results revealed the variants in 1762 samples and 38 samples were rejected due to poor sample quantity.

The average age of the covid-19 patients was 37.4 ± 0.5 years with age range 2–80 years with median age of 24 years and mode age of 32

Table 1. Different Variants of SARS-CoV-2 and their mutations in the spike protein

| Variant Identified | Variation Identified in spike Protein | Name of the Variant |
|--------------------|--------------------------------------|---------------------|
| B.1                | D614G                                | UK variant          |
| B.1.1.7            | N501Y, P681H, 69/70 Del               | South Africa variant|
| B.1.351            | N501Y, E484K, K417N, 243/245 Del      | Kappa-Indian variant|
| B.1.617.1          | L452R, E484Q and P681R                | Delta- Indian variant |
| B.1.617.2          | L452R, P681R and T478K                | Delta plus- Indian variant |
| AY.1               | L452R, P681R, T478K and K417N         | Unknown              |
| Unidentified       | (N440K), (P681H, 69/70, 144/145 DEL), (E484Q, L452R, P681R, Q1071H), (P681H), (K417N, N501Y), (E484K), (N501Y), (K417N) |                        |

Fig. 1. Trends in the distribution of positivity as compared to total sample during the second wave in time period of January 2021 to May 2021. New cases in 5 districts of Punjab are rapidly on the rise since February reaching a peak of approx. about 35,0000 among which there is upsurge of positive cases every month.
years. The overall gender ratio of female-to-male was 713:1048. The median age of female covid-19 patients were 37 years with mode age of 50 years. The median age of male covid-19 patients were 24 years with mode age of 18 years.

SARS-COV-2 variant: In the present study various variants of SARS-COV-2 identified using whole genome sequencing in the studied population groups were B.1, B.1.1.7, B.1.351, B.1.617.1, B.1.617.2, B.1.167.2.1/AY.1 and other unidentified substitutions and deletions. The presence of different combinations of mutations in amino acids has been found in the studied population groups, many variants share similar mutations as depicted in Table 1. A total of 11 amino acid replacements /SNPs (Single nucleotide polymorphisms) and 3 deletions/ substitution were recognized. Frequency of the amino acid replacement has been shown in Fig. 2.

Prevalence of SARS-COV-2 variant: In the present study it was observed that till December 2020 wild type (Wuhan strain) was predominant as no variant was detected at that time but from January 2021 to May 2021. B.1.1.7 (UK variant)
was dominant in February which further modified to B.1.617.2 (delta- Indian Variant) by starting of April. Another interesting variant AY.1 (Delta- plus Indian variant) was observed in the study by the end of April and starting of May with other unidentified variants as shown in Fig. 3. Among the five studied districts of Malwa region the dominant variant (B.1.1.7/UK variant) was found majorly in Patiala followed by Ludhiana, SBS Nagar, Mansa and Sangrur. Similarly B.1.617.2 (delta- Indian variant) was dominant in Patiala, Ludhiana followed by Mansa and Sangrur (Table 2). In addition to this, sequencing results also observed that dominant trait was more prevalent in male population and age group 21-40 years.

Gender and Age wise distribution of SARS-CoV-2 variant: The SARS-CoV-2 whole genome sequencing of both the gender revealed that the B.1.1.7 (UK variant) dominates in both the gender categories (Fig. 4a). Similarly, in other variants B.1, B.1.351, B.1.617.1, wild (Wuhan strain) and unidentified variants, there is no major difference in the distribution between male and female population. The SARS-CoV-2 whole genome sequencing was also analyzed among different age groups ranging from 0-20 years, 21-40 years, 41-60 years, 61-80 years and more than 80 years. Highest numbers of confirmed positive cases were observed in younger age group 21-40 years followed by age group 41-60 years (Fig. 4b).

Table 2. Distribution of SARS-CoV-2 variants during the time period of January 2021 to May 2021 in different geographic regions of Punjab (Data provided in supplementary sheets)

| Variant/District | Ludhiana | Mansa | Patiala | Sangrur | SBS Nagar |
|------------------|----------|-------|---------|---------|-----------|
| B.1              | 2 (14.3%)| 0     | 2 (14.3%)| 10 (71.4%)| 0         |
| B.1.1.7          | 223 (24.8%)| 30 (3.3%)| 512 (56.9%)| 23 (2.6%)| 112 (12.4%)|
| B.1.167.1        | 12 (41.4%)| 0     | 17 (58.6%)| 0       | 0         |
| B.1.167.2        | 116 (17.5%)| 30 (3%)| 508 (76.5%)| 20 (3%)| 0         |
| B.1.167.2.1/AY.1 | 1 (50%)| 0     | 1 (50%)| 0       | 0         |
| B.1.351          | 1 (100%)| 0     | 0       | 0       | 0         |
| Wild (Wuhan strain) | 16 (20%)| 0     | 44 (55%)| 0       | 20 (25%) |
| Unidentified     | 22 (46.8%)| 0     | 22 (46.8%)| 0       | 3 (6.4%) |
| Total            | 393      | 50    | 1106    | 53      | 135       |

Fig. 4. Bar graphs showing frequency of SARS-CoV-2 variants B.1, B.1.1.7, B.1.351, B.1.617.1, B.1.617.2, Wild/Wuhan strain, unidentified variant. a) Among different genders b) Among different age group of Malwa region in Punjab, India. The occurrence of B.1.1.7 variant in male and female is 46.5% and 52% respectively. The variant B.1.617.2 (Delta Indian variant) is the second highest among the population studied as the percent occurrence of this variant among male and female is 39.5% and 35% respectively. In all the age groups the dominant variant was B.1.1.7 (UK variant) followed by B.1.617.2 (Delta Indian variant) and the other variants were equally distributed in all the age groups.
DISCUSSION

Covid-19 cases have continued to surge in India during second wave as new records have been made by current outbreak of SARS-CoV-2 in India. Prior to second wave less than 0.7% of population was infected but by the end of April 2021 new cases reached to 1.5 million as per Indian government report.11 The abrupt hike in SARS-CoV-2 cases in India corresponds with high prevalence of more-transmissible variant, associated with diagnostic test failures and antibody escape.12 Population-based surveillance helped to evaluate and screen the trend of infection for SARS-CoV-2 in the studied population. Whole genome sequencing revealed the presence of B.1, B.1.1.7 (UK variant), B.1.351 (South Africa), B.1.617.1 (kappa-Indian variant), B.1.617.2 (delta-Indian variant), AY.1 (delta plus- Indian variant) and unidentified variants of SARS-CoV-2 in the studied population. In the present study, the sequencing report revealed that during the 1st wave till December and January end wild type (Wuhan strain) was more prevalent but during second wave B.1.1.7 (UK variant) was found to be the most dominant variant in the population followed by B.1.617.2 (delta-Indian variant), B.1.617.1 (kappa-Indian variant), AY.1 (delta plus- Indian variant), B.1.351 (South Africa), B.1 and other unidentified variants in the studied population.

Variant B.1.1.7 has been identified in the month of September 2020 in England and has quickly been spread over to other countries including India.13 This variant has been classified as variant of concern (VOC, 202012/1). Current study showed highest prevalence of this variant from month of January till May. In the studied population B.1.1.7 variant carrying N501Y mutation in spike protein found to have SNP (single nucleotide polymorphism) in which asparagine (N) changes to tyrosine (Y) at 501 position. This mutation leads to lower sensitivity towards immune responses as compared to the wild Wuhan variant. Vaccine trials suggest that infection with variants may offer only restricted protection from reinfection with the 501YYV2 variant.14 B.1.1.7 variant also has poor diagnosis ability as it has deletion in the spike protein.15 It has been responsible for the upsurge in the mortality rate with the 1.35 fold higher probability and estimated to be 40-80% more infectious than wild type and other variants due to the increased viral load.16-18 Reports suggest that in Delhi the distribution of B.1.1.7 (VOC) was minimal (5%) and then gradually enlarged up to (60%) by the end of April 202119 while in Karnataka the lineage was identified to be 44.4% from both imported cases and circulating cases.20

B.1.351 variant reported in the present study was also found in multiple countries but it was first identified in South Africa in October, 2020 and has the increase transmissibility because of the mutations in the spike gene.21,22 B.1.351 shares identical mutations with B.1.1.7 and also seems to show reduced sensitivity to acquired immune responses against the ‘wild-type’ Wuhan virus.23 Its prevalence has been reported in West Bengal with percent value of 16.9.23

Another major variant of concern of current study was B.1.617 of the SARS-CoV-2 consisting 3 sub-lineages such as B.1.617.1, B.1.617.2 and B.1.617.3. The emergence of a new lineage of current study B.1.617.1 and B.1.617.2 has raised concerns among public health. B.1.617.1 and B.1.617.2 lineage of the SARS-CoV-2 referred to as triple mutant and first recognized in India but now mounting in prevalence across the country as shown by genome sequencing data.24 In present study also B.1.617.2 sub-lineage was the second most dominating variant from January to April. In fact all the three lineages of B.1.617 were reported to have L452R mutation. Mutation L452R was observed in the California variant having more transmissibility and viral load.25 The mutations L452R, E484Q, and P681R found in the current study shared by lineage B.1.617.1 while the B.1.617.2 lacks the E484Q mutation.26-28 Additionally, Delta plus (AY.1) variant acquired K417N mutation in the receptor binding domain of spike protein, which was first found in Beta variant. The first two cases of Delta plus variant in the study group were found in Patiala and Ludhiana district in the month of April 2021. Scientists suggest that this variant pose various issues including rapid transmissibility, antibody neutralization and reduced effectiveness of vaccine.29 Other Q1071H mutation was also observed in the present study and also shared by B.1.617.1 lineage30 and in triple Mutant Bengal Strain (B.1.618). Interestingly, mutations P681R and Q1071H observed in the current study were present in the B.1.618 variant along with E144K mutation.31 The E484Q mutation
is similar to E484K, a mutation found in the UK variant and South Africa variants. These mutations are suggested to cause reduction in body weight, histopathological changes in lung, lung lesions. Studies have reported that the highest numbers of this variant were from India than other countries Brazil, Argentina, United States. B.1.617.2 sub-lineage has increase from 10% to about 80% in Delhi from January to April. The variant has also been recognized in several nations with higher transmissibility, pathogenicity and immune escape.

Interestingly, other unidentified replacements in amino acids have also been found in the present study. Among them N440K, P681H considered to be powerful having more virulence, infectious and have higher immune escape than the wild type strain. N440K mutation in SARS-CoV-2 became the reason behind the major destruction caused by the SARS-CoV-2 in Visakhapatnam, Karnataka, Telangana and other southern parts of India. N501Y mutation found in this study was also detected in Telangana, the southern parts of India. Occurrence of re-infections and rapid transmission is mainly due to the emergence of all these new variants, either after natural infection or after vaccination. A recent study confirmed that re-infections are already happening in India.

Mutations N501Y and E484K found in the present study are acquired by the B.1.1.7 lineage whereas N501Y mutation has also been shared by B.1.351 and P.1 lineages along with K417N/T and E484K mutations. The E484Q and L452R mutations observed in the current study were also present in the Indian variant B.1.617. These variants have been associated with increased transmissibility and are more prone to immune escape. Other unidentified mutations found in the study such as L452R, E484Q and P681R in the spike protein are possessed by the B.1.617 lineage. Additionally, these mutations are also been reported in other globally circulating lineages. Most variants of the study are identified to have increased transmissibility and neutralization of antibody.

**Limitations of the present study**

Most of the individuals in the studied population did not give information about the international travel history during second wave of covid-19 surge. Hence, the primary source of SARS Covid-19 spread could not be detected. The primary source assumed be locality, neighbourhood, their workstation or may emerge from other countries of the world. Larger number of whole genome sequencing and geographically diverse population will be help to better characterize the associations between disease severity and variations.

**CONCLUSIONS**

Altogether, we can conclude that our data highlighted the increased frequency of B.1.1.7 (UK variant) and B.1.617.2 (delta Indian variant) lineage in Patiala and Ludhiana districts of Punjab. This study also revealed several unidentified deletions/substitution in untranslated and translated regions of the SARS-CoV-2 genomes. In this wave younger population of studied area was more effected than other age groups. The mutations found in the study were globally circulated and were more virulent and dangerous than the original wild strain. Further research should emphasis on structural confirmations and phenotypic consequences of these variations. Moreover, the identification of the structural conformational changes helps to elucidate the virulence, metabolic pathway, pathogenicity, and transmission of SARS-CoV-2.

**ACKNOWLEDGMENTS**

We would like to acknowledge the staff members of the Viral Research and Diagnostic laboratory at tertiary care Hospital for dedication to providing excellent team work. We thank NCDC (National Center for Disease Control), New Delhi for whole genome sequencing. We would like to thank the frontline health care workers who remain dedicated in the battle against SARS-CoV-2. We would also like to acknowledge ICMR-DMR (Indian Council of Medical Research- Department of Medical Research), New Delhi for supporting us.

**CONFLICT OF INTEREST**

The authors declare that there is no conflict of interest.

**AUTHORS’ CONTRIBUTION**

All authors listed have made a substantial, direct and intellectual contribution to the work, and approved it for publication.

**FUNDING**

None.
DATA AVAILABILITY
All datasets generated or analyzed during this study are included in the manuscript.

ETHICS STATEMENT
Not applicable.

REFERENCES
1. Rajendrakumar AL, Narayanan Nair AT, Nangia C, et al. Epidemic Landscape and Forecasting of SARS-CoV-2 in India. *J Epidemiol Glob Health*. 2020. doi: 10.1101/2020.04.14.20065151
2. Dey JK, Dey SK. SARS-CoV-2 Pandemic, COVID-19 Case Fatality Rates and Deaths per Million Population in India. *J Bioinfomatics, Comput Syst Biol*. 2020;2(1):110.
3. Andersen KG, Rambaut A, Lipkin WI, Holmes EC, Garry RF. The proximal origin of SARS-CoV-2. *Nat Med*. 2020;26(4):450-452. doi: 10.1038/s41591-020-0820-9
4. Joshi A, Paul S. Phylogenetic Analysis of the Novel Coronavirus Reveals Important Variants in Indian Strains. *bioRxiv*. 2020;1:12. doi: 10.1101/2020.04.14.041301
5. Harvey WT, Carabelli AM, Jackson B, et al. SARS-CoV-2 variants, spike mutations and immune escape. *Nat Rev Microbiol*. 2021;19(7):409-424. doi: 10.1038/s41579-021-00573-0
6. Fujino T, Nomoto H, Kutsuna S, et al. Novel SARS-CoV-2 Variant in Travelers from Brazil to Japan. *Emerg Infect Dis*. 2021;27(4):1243-1245. doi: 10.3201/eid2704.210138
7. Srivastava S, Banu S, Singh P, Sowpati DT, Mishra SE. SARS-CoV-2 variants of concern are emerging in India. *J Epidemiol Glob Health*. 2021;46(1):22. doi: 10.1016/j.jegh.2020.05.071
8. Kaur K, Kaur R. Axiology of DNA damage by KF, XPG and ERCC1 gene polymorphisms in pesticide-exposed agricultural workers of Punjab, North-West India. *Mutat Res - Genet Toxicol Environ Mutagen*. 2021;861-862:503302. doi: 10.1016/j.mrgentox.2020.503302
9. Novel coronavirus disease (Covid-19) situation update report-64. 2021. [https://cdn.who.int/media/docs/default-source/windia/situation-report/india-situation-report-64.pdf?sfvrsn=7bf313d6_4. Accessed April 28, 2021.
10. Singh J, Rahman SA, Ehtesham NZ, Hira S, Hasnain SE. SARS-CoV-2 variants of concern are emerging in India. *Nat Med*. 2021;27(7):1131-1133. doi: 10.1038/s41591-021-01397-4
11. Galloway SE, Paul P, MacCannell DR, et al. Emergence of SARS-CoV-2 B.1.1.7 lineage. *Morbidity and Mortality Weekly Report*. 2021;70(3):95-99. doi: 10.15585/mmwr.mm7003e2
12. Akkiz H. Implications of the Novel Mutations in the SARS-CoV-2 Genome for Transmission, Disease Severity, and the Vaccine Development. *Front Med*. 2021;8:636532. doi: 10.3389/fmed.2021.636532
13. Challen AR, Dyson L, Overton CE, Guzman-rincon LM. Early epidemiological signatures of novel SARS-CoV-2 variants : establishment of B.1.617.2 in England. *medRxiv*. 2021. doi: 10.1101/2021.06.06.21215365
14. Davies NG, Jarvis CI, van Zandvoort K, et al. Increased mortality in community-tested cases of SARS-CoV-2 lineage B.1.1.7. *Nature*. 2021;593(7858):270-274. doi: 10.1038/s41586-021-03426-1
15. Volz E, Mishra S, Chand M, et al. Assessing transmissibility of SARS-CoV-2 lineage B.1.1.7 in England. *Nature*. 2021;593(7858):266-269. doi: 10.1038/s41586-021-03470-x
16. Brown KA, Gubbay J, Hopkins J, et al. Rapid Rise of S-Gene Target Failure and the UK variant B.1.1.7 among COVID-19 isolates in the Greater Toronto Area, Canada. *medRxiv*. 2021. doi: 10.1101/2021.02.09.21215225
17. Dhar MS, Marwal R, Radhakrishnan VS, et al. Genomic characterization and Epidemiology of an emerging SARS-CoV-2 variant in Delhi, India. *medRxiv*. 2021. doi: 10.1101/2021.06.02.21215807
18. Pattabiraman C, Prasad P, George AK, et al. Importation, circulation, and emergence of variants of SARS-CoV-2 in the South Indian state of Karnataka. *Wellcome Open Res*. 2021;6:110. doi: 10.12688/wellcomeopenres.16768.1
19. Tegally H, Wilkinson E, Giovannetti M, et al. Detection of a SARS-CoV-2 variant of concern in South Africa. *Nature*. 2021;592(7854):438-443. doi: 10.1038/s41586-021-03402-9
20. Wang P, Nair MS, Liu L, et al. Antibody resistance of SARS-CoV-2 variants B.1.351 and B.1.617.2 with sera of COVID-19 recovered cases and vaccinees of BBV152. *bioRxiv*. 2021. doi: 10.1101/2021.05.08.21243632
21. Yadav PD, Nyayanit DA, Sahay RR, et al. Isolation and characterization of the new SARS-CoV-2 variant in travellers from the United Kingdom to India: VUI-202012/01 of the B.1.1.7 lineage. *J Travel Med*. 2021;28(2):1-3. doi: 10.1093/jtm/taab009
22. Yadav PD, Sapkai GN, Ella R, et al. Neutralization and characterization of the new SARS-CoV-2 variant in travellers from the United Kingdom to India: VUI-202012/01 of the B.1.1.7 lineage. *J Travel Med*. 2021;28(2):1-3. doi: 10.1093/jtm/taab009
23. Ferreira I, Datir R, Papa G, et al. SARS-CoV-2 B.1.617.2 in England. *Nature*. 2021;593(7858):266-269. doi: 10.1038/s41586-021-03470-x
24. Yadvay PD, Nyayanit DA, Sahay RR, et al. Isolation and characterization of the new SARS-CoV-2 variant in travellers from the United Kingdom to India: VUI-202012/01 of the B.1.1.7 lineage. *J Travel Med*. 2021;28(2):1-3. doi: 10.1093/jtm/taab009
of Spike protein triggers recent massive expansion of SARS-CoV-2 variants. *bioRxiv Prepr Serv Biol.* 2021. doi: 10.1101/2021.02.22.432189

29. Biswas S. Delta plus India: Scientists say too early to tell risk of Covid-19 variant. 2021. https://www.bbc.com/news/world-asia-india-57564560. Accessed April 29, 2021.

30. Shrestha UK. B.1.617. 2 variant and increasing surge of COVID-19 in Nepal. *Nepal Mediciti Medical Journal.* 2021;2(1):1-4. doi: 10.3126/nmmj.v2i1.37212

31. Cherian S, Potdar V, Jadhav S, et al. Convergent evolution of SARS-CoV-2 spike mutations, L452R, E484Q and P681R, in the second wave of COVID-19 in Maharashtra, India. *bioRxiv.* 2021:. doi: 10.1101/2021.04.22.440932

32. B.1.167 Covid-19 variant found in at least 53 territories:WHO. 2021. https://www.deccanherald.com/international/world-news-politics/b1167-covid-19-variant-found-in-at-least-53-countries-who-990141.html. Accessed June 29, 2021.

33. Ahmed SM, Juvvadi SR, Kalapala R, Sreemanthula JB. Detection of SARS-CoV-2 N501Y mutation by RT-PCR to identify the UK and the South African strains in the population of South Indian state of Telangana. *medRxiv.* 2021. doi: 10.1101/2021.03.27.21254107

34. Yuan M, Huang D, Lee C-CD, et al. Structural and functional ramifications of antigenic drift in recent SARS-CoV-2 variants. *bioRxiv Prepr Serv Biol.* 2021. doi: 10.1101/2021.02.16.430500

35. Xie X, Liu Y, Liu J, et al. Neutralization of SARS-CoV-2 spike 69/70 deletion, E484K and N501Y variants by BNT162b2 vaccine-elicited sera. *Nat Med.* 2021;27(4):620-621. doi: 10.1038/s41591-021-01270-4

36. Noh JY, Jeong HW, Shin E-C. SARS-CoV-2 mutations, vaccines, and immunity: implication of variants of concern. *Signal Transduct Target Ther.* 2021;6(1):203. doi: 10.1038/s41392-021-00623-2