SARS-CoV-2 mutations on diagnostic gene targets in the second wave in Zimbabwe: A retrospective genomics analysis

C Nyagupe,1,2 BSc, MSc; L de Oliveira Martins,3 MSc, PhD; H Gumbo,4 Grad Cert, MSc; T Mashe,5 MSc, PhD; T Takawira,2 MSc,PhD; K K Maeka,4 MSc, MComPM; A Juru,1 HBMLS, MPH; L K Chikanda,1 BSc; MSc; A R Tuya,1 BSc, MSc; A J Page,6 MSc, PhD; R A Kingsley,4 MSc, PhD; R Simbi,1 MSc, PhD; J Chirenda,1 MSc, PhD; J Manasa,1 MPhil, PhD; V Ruhanya,1 MSc, PhD; R T Mavenyengwa,1 MSc, PhD

1 Microbiology Unit, Faculty of Medicine and Health Sciences, University of Zimbabwe, Harare, Zimbabwe
2 National Microbiology Reference Laboratory, Ministry of Health and Child Care, Harare, Zimbabwe
3 Quadram Institute Bioscience, Norwich, UK
4 Directorate of Laboratory, Ministry of Health and Child Care, Harare, Zimbabwe

Corresponding author: C Nyagupe (nyagupecharles@gmail.com)

Background. SARS-CoV-2 continues to be a major issue in resource-limited settings, particularly owing to the limited supply of vaccines caused by inequitable distribution.

Objective. To monitor diagnostic gene targets to identify potential test failures caused by mutations, which is important for public health.

Methods. Here we analysed the genome sequence of SARS-CoV-2 from the second wave in Zimbabwe. A total of 377 samples were sequenced at Quadram Institute Bioscience. After quality control, 192 sequences passed and were analysed.

Results. The Beta variant was dominant during this period, contributing 77.6% (149) of the genomes sequenced and having a total of 2994 mutations in diagnostic polymerase chain reaction target genes. Many single nucleotide polymorphism mutations resulted in amino acid substitution that had the potential to impact viral fitness by increasing the rate of transmission or evading the immune response to previous infection or vaccination.

Conclusion. There were nine lineages circulating in Zimbabwe during the second wave. The B.1.351 was dominant, accounting for >75%. There were over 3 000 mutations on the diagnostic genes and lineage B.1.351, contributing almost two-thirds of the mutations. The S-gene had the most mutations and the E-gene was the least mutated.

frames (ORFs) include the S (spike) protein, ORF3a protein, E (envelope) protein, M (membrane) protein, and N (nucleocapsid) protein, of 3 822, 828, 228, 669, and 1260 nt, respectively. The 5’ and 3’ terminal sequences of Wuhan-Hu-1 are also typical of Beta coronaviruses with lengths of 265 nt and 229 nt, respectively.[39]

In Zimbabwe, the second wave began in October 2020 with cases rising significantly toward December 2020 and early January 2021. PCR tests were conducted at the National Microbiology Reference Laboratory (NMRL) between October 2020 and January 2021 and had a test positivity rate of 18% (3 299 of 18 000) compared with 2.2% (236 of 10 500) in the first wave between March and June 2020. Genomic surveillance revealed that the Beta variant was driving the rise in case numbers. The SARS-CoV-2 virus mutates at a rate of approximately two nucleotides per genome replication cycle,[39] which is lower than that of influenza (4 nt per genome replication cycle) or HIV (8 nt per genome replication cycle),[40] nonetheless presenting a risk that detection strategies do not accommodate this genomic change. There are many examples of this in public health diagnostics, such as a drop in the detection of Chlamydia trachomatis in Sweden,[13] which was hailed as a public health success until genome sequencing identified a mutation that had disrupted the PCR diagnostic target and cases had increased, in Belgium a single nucleotide polymorphism (SNP) in the E gene caused a diagnostic dropout on the Roche Cobas platform,[41] and an S-gene target failure (SGTF) on the TaqPath assay following the deletion of two amino acids was used as a proxy for the Alpha variant in the UK.[35] Also in countries including Australia, Denmark, Saudi Arabia, the UK and the USA, isolates containing the SNP C29200T abolished N2 detection on the Gene Xpert assay.[13] Though PCR remains the gold standard for SARS-CoV-2 diagnosis, the mutations on the diagnostic genes that include ORF1ab, S, N, and E may compromise the sensitivity of the kits used in its diagnosis. The test kit used during the second wave in Zimbabwe was the DaAn gene (DaAn Gene Co, Ltd, China) targeting the ORF1ab and the N gene.

Genomic surveillance of SARS-CoV-2 helps to track the changes and identify mutations that may compromise the sensitivity of diagnostic targets. Moreover, it also helps to detect variants with the ability to spread more quickly, cause mild or severe disease, decreased susceptibility to therapeutics that employ monoclonal antibodies, and those with the ability to evade natural or vaccine-induced immunity. There is a paucity of data on the variants and mutations in SARS-CoV-2 that circulated in the second wave in Zimbabwe. We need answers to this question to understand the evolution of the virus and its clinical and public health impact.

What were the major variants and mutations and possible impacts on molecular diagnosis focusing on genes targeted in PCR assays during the second wave in Zimbabwe? In this study, we aimed to use genomic surveillance to determine mutations and variants that may have arisen or been introduced in Zimbabwe during the second wave. The objectives were to identify the variants and mutations on diagnostic genes in PCR assays used in Zimbabwe, and how they are spread in the population.

Methods
Study design and data source
We have undertaken a retrospective study of 377 routinely collected nasopharyngeal samples processed by RT-PCR at the NMRL, Harare, Zimbabwe, and sequenced at the Quadram Institute Bioscience (QIB). The samples were collected by trained clinicians in all the 10 provinces of Zimbabwe. A PCR cycle threshold (Ct) value of ≤30 was used to select positive samples for sequencing.

Demographic and epidemiological information was obtained, such as geographical origin, case classification obtained from the NMRL's laboratory COVID-19 request forms and the electronic laboratory information management system (LIMS), which is confidential and can only be accessed by authorised personnel through the use of a private username and password.

Ethics
Ethical permission was obtained from the institution head (NMRL), the Joint Research Ethics Committee (JREC) of the University of Zimbabwe and Parirenyatwa group of hospitals (ref. no. 54/2022), and the Medical Research Council of Zimbabwe (MRCZ) (ref. no. MRCZ/B/2191). There was no specific consent obtained from the patient as the NMRL has the legal mandate to handle patient information for public health monitoring as enshrined in section 46 (notifiable diseases) of the Zimbabwe Public Health Act 2018.[44]

Specimen processing
Processing of the COVID-19 specimens was undertaken in two steps: extraction of nucleic acids, and amplification. Extraction of the nucleic acid was undertaken using the bioMérieux NucliSens Easy Mag platform, which is a semi-automated extraction machine, following the manufacturer’s COVID-19 extraction protocol.[15] The extraction process is described in Box 1. Amplification of the eluates was undertaken using Quanti Studio 3, a Thermo Fisher, USA, product which is an open system.[46] The DaAn Gene RT kit for the detection of SARS-CoV-2 was used on the Quant Studio 3 thermocycler. The DaAn Gene protocol is described in Box 1.[47]

cDNA synthesis
A total of 377 conveniently selected PCR-positive samples for SARS-CoV-2 with a Ct value ≤30 were selected for cDNA synthesis and were forwarded for genomic sequencing. The process of cDNA synthesis is outlined in Box 2.

Genome sequencing
The cDNA was frozen at –80°C before being packaged in dry ice and sent to QIB for sequencing using Illumina NextSeq 500 following the Coronavirus protocol. Good-quality sequences were uploaded on the Gisaid and Excel (Microsoft, USA) data of all the sequences with metadata were sent to NMRL for further mutation analysis.

Phylogenetic analysis
A phylogenetic tree was generated from the consensus genomes for the samples with at least 90% of the genome present using the next strain/next clade software.[49]

Data analysis
Electronically captured demographic and epidemiological data were cleaned and then imported into STATA version 12.1 (Stata Corp, USA) and Excel for analysis.

Genome analysis
Mutation rate and frequencies on the diagnostic genes were analysed by genome detective online tool version 1.137,[20] and the consensus genome was uploaded to Pangolin software[40] for lineage assignment.

Limitations of the study
The study only focused on RT-PCR SARS-CoV-2 positive samples detected by sequencing in the second wave in Zimbabwe. The
Box 1. Processing of SARS-CoV-2 nasopharyngeal sample on the bioMérieux NucliSens Easy Mag Extraction machine and amplification on Quanti Studio 3 using the DaAn gene kit at the NMRL during the second wave (October 2020 - January 2021).

**Extraction.** A sample volume of 400 μL is pipetted into a tube containing 2 mL of commercially prepared lysis buffer. It is left to stand for 30 minutes for lysis to occur. A volume of 400 μL of the sample and 65 μL of the eluate is programmed on the machine. The lysed sample is pipetted into sample processing chambers of the machine called disposables, and then 100 μL of diluted silica at a ratio of 1:1 with the commercially prepared diluent is added. Extraction takes 40 minutes and the Boom technique is employed. A volume of 65 μL of the eluate is dispensed after extraction. The machine is serviced by a local trained bioMérieux representative.

**Amplification.** A volume of 5 μL of eluate and 20 μL Da-An gene of the PCR mix solution A and B are mixed in 0.2 mL PCR tubes. The Quanti Studio 3 Thermo Fisher PCR machine is programmed following the DaAn gene RT-PCR SOP for NMRL.

Box 2. cDNA was synthesised from all the samples with a Ct value ≤30. A volume of 5.5 μL of total nucleic acids (TNA) extracts was transferred to each well on a 96-well nested plate aliquoted with 1 μL of Luna script reagent. The reaction mixture was incubated in a thermal cycler under the following set conditions: at 25°C for 2 minutes, 55°C for 10 minutes, 95°C for 1 minute then held at 4°C for 1 minute.

Results

A total of 377 samples were sequenced during the second wave (October 2020 - January 2021). However, after careful selection based on quality control pass and cycle threshold value of ≤30, we were left with 192 samples. Of the 192 good-quality sequences, 53.13% (102) were males, 44.79% (86) were females, 1.56% (3) were of unknown gender and 0.52% (1) were designated as blank as shown in Fig. 1. The 31 - 45-year age group was the most affected, with (38.02%), followed by the 46 - 60 and 16 - 30-year age groups, with 20.83% and 19.79%, respectively, as shown in Fig. 2. The Ct value for our targets was ≤30. For the screening gene N, it ranged from 9.5 to 29.6, and for the ORF1ab confirmatory gene, it ranged from 9.8 to 29.9 with averages of 21.7 and 23.0, respectively. The single nucleotide polymorphisms (SNP) positions on diagnostic genes amino acid positions affected and the nature or type of mutations for the sequenced samples during the second wave in Zimbabwe are shown in Table 1. The total percentage mutation and amino acid substitution of different gene targets for the variants circulating among the 192 sequenced genomes during the second wave in Zimbabwe are shown in Table 2.

SARS-CoV-2 lineages circulating in Zimbabwe for the sequenced samples during the second wave

The VOC B.1.351 lineage using Pangolin nomenclature, or clade 20H according to Nextstrain (https://clades.nextstrain.org) was the dominant variant circulating in Zimbabwe during the second wave, accounting for 149 (77.60%). The VOI C.2 was the second dominant variant circulating, with 15 (7.81%). All the variants circulating during the second wave are shown in Fig. 3.

Phylogeny

The phylogeny tree below in Fig. 4 was constructed by loading the Zimbabwean second-wave FASTA file extracted from GISAID onto the next strain software to generate clades circulating. The 20H clade (Beta variant), or B.1.351 lineage according to Pango, was the

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**Table 1. Common SNP positions on diagnostic genes amino acid positions affected and the nature or type of mutations for the sequenced samples during the second wave in Zimbabwe**

<table>
<thead>
<tr>
<th>SNP</th>
<th>Diagnostic gene</th>
<th>Amino acid position</th>
<th>Nature/type of mutation</th>
</tr>
</thead>
<tbody>
<tr>
<td>C14408T</td>
<td>RdRp</td>
<td>P323L</td>
<td>Non-synonymous</td>
</tr>
<tr>
<td>C28881A</td>
<td>N</td>
<td>R203K</td>
<td>Transition and non-synonymous</td>
</tr>
<tr>
<td>C28882A</td>
<td>N</td>
<td>R203K</td>
<td>Transition and non-synonymous</td>
</tr>
<tr>
<td>G22813T</td>
<td>S</td>
<td>K417N</td>
<td>Transmission</td>
</tr>
<tr>
<td>A23013C</td>
<td>S</td>
<td>E484A</td>
<td>Transmission</td>
</tr>
<tr>
<td>A23403G</td>
<td>S</td>
<td>D614G</td>
<td>Transition and non-synonymous</td>
</tr>
<tr>
<td>C23604G</td>
<td>S</td>
<td>P681P</td>
<td>Transition and non-synonymous</td>
</tr>
<tr>
<td>C23664T</td>
<td>S</td>
<td>A701V</td>
<td>Transmission</td>
</tr>
<tr>
<td>A23063T</td>
<td>S</td>
<td>N501Y</td>
<td>Transmission</td>
</tr>
</tbody>
</table>

SNP= single nucleotide polymorphism.

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**Fig. 1. Distribution of SARS-CoV-2 infection by gender during the second wave in Zimbabwe.**

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major circulating variant during this wave in Zimbabwe.

Results analysis
Of the 192 SARS-CoV-2 sequences, mutations resulted in a total of 3,452 nucleotide substitutions. The Beta variant, lineage B.1.351, had 2,994 (86.73%) of the nucleotide substitution and was the major circulating variant during the second wave period between October 2020 and January 2021. It accounted for 77.6% of the SARS-CoV-2 variants circulating in Zimbabwe. C.2, a VOI, was the second-most common circulating, with 7.81%, followed by B.1.174 with 4.69%. B.1, B.1.210, A23.1 and A lineages were also circulating in the second wave, with the corresponding percentages of 3.13, 1.56, 1.04 and 1%, respectively.

B.1.351 lineage
The Beta variant had the most mutations on the diagnostic genes. All 149 confirmed sequences had common mutations on all the diagnostic genes. On the E-gene, all the 149 sequences showed mutations on the diagnostic gene of the B1.351 variant, accounting for 39.81% of all the mutations, with all the 149 sequences showing mutations in the following positions: D80A, D614G, and H70V. Mutations on positions D215G (93.95%), L242X (92.61%), H245X (91.95%), K417N (96.64%), E484K (60.4%) and N501Y (64.42%) were also common among the sequences. These mutations confer viral fitness. They do not impact diagnostic targets. However, diagnostic impact was seen in the Alpha variant, lineage B.1.1.7, first detected in the UK in December 2020, where an SGTF was reported in TaqPath RT-PCR kit due to 69-70 amino acid deletion. This is now used as a marker of the VOC lineage B.1.1.7.

C.2 lineage
This was the second-most common circulating variant in the second wave in Zimbabwe, accounting for 15 (7.81%) sequences. The 15 sequences had a total of 224 amino acid substitutions. All the diagnostic genes had one or more mutations, except for the E-gene. Of the sequences, 14 (93.33%) had non-synonymous transition SNP substitution on G28881A, G28882A, R203K and G28883 (G204R) positions. The N-gene mutation, which includes amino acid substitution S14408T, is conserved in SARS-CoV-1 and SARS-CoV-2 variants circulating in Europe. However, they are not significant in terms of disease severity and immune evasion. They show common mutations on the diagnostic genes at position R203K on the N-gene, position P314L on the ORF1b, B.1.1.7 lineage from the E-gene, position D614G on the S-gene. Mutation at position E1311A has only been shared between the B.1.1.7 and B.1.1.210 lineages, while B.1.1.1 shows no mutation in this diagnostic gene. All three lineages show no mutation in the E-gene.
A-lineage

The original Wuhan strain was also still circulating in Zimbabwe during the second wave, accounting for 2 (1.04%) of the circulating lineages. However, there were no mutations on all the diagnostic genes except the ORF1a gene of one of the two sequences, which exhibited only one substitution at position N1776S, which was not present in the diagnostic genes of preceding variants.

Discussion

To the best of our knowledge, this is one of the studies that tried to analyse mutations on the diagnostic genes of SARS-CoV-2 in the sequenced samples in Zimbabwe during the second wave, and their impact on diagnostic kits, targets and immune escape. The SA VOC was the major circulating variant in Zimbabwe during the second wave, accounting for almost 80% of the circulating variants during that wave. The B.1.351 is said to have 21 plus mutations, 10 of which occur on the S-gene, which is one of the targets of the RT-PCR kit. Besides the D614G mutation, which occurs on the S1 subunit of the S-gene and confers viral fitness of high viral load, the viral fitness conferred by this mutation may harbour mutagenic properties and enable the virus to spread. The A23063T (N501Y), a transversion mutation, sits on the receptor-binding domain (RBD), increasing transmission of the variant by increasing the binding of the spike protein to the angiotensin converter enzyme receptor 2 (ACE2) of the host cells, facilitating the entry of the virus in human cells.\(^\text{29}\) SNP A33013C (E484K) and G22813T (K417N) are both transversions in nature, and are located in the RBD spike protein. The E484K mutation was found to reduce the neutralisation potency of some convalescent sera by >10-fold, and this can be a potential viral escape. SGTF was confirmed in the UK on the Thermo Fisher Taqpath RT-PCR kit with three targets ORF1a, N and S-gene. There were S-gene dropouts due to amino acid 69 and 70 deletion in the Alpha VOC, lineage B.1.1.7. However, the impact on molecular diagnostic tests was found to be minimal since most of the test kits target more than one gene.\(^\text{29}\) Similarly, all the 192 tests in this study passed quality control with good Ct values ranging from 9.4 to 29.4 for gene target 1 (ORF1ab) and 10.7 to 29.9 for gene target 2 (N).

The C2 lineage had non-synonymous mutations on position R203K, and G204R of the N-gene was also detected on the latest VOC omicron, lineage B.1.1.529, detected in SA.\(^\text{30}\) These mutations are believed to increase viral fitness survival, spread and adaptations
### Table 2: Total percentage mutation and amino acid substitution of different gene targets for the variants circulating among the sequenced genomes during the second wave in Zimbabwe

<table>
<thead>
<tr>
<th>Variant</th>
<th>E-gene n, %</th>
<th>M-gene n, %</th>
<th>N-gene n, %</th>
<th>ORF1a-gene n, %</th>
<th>ORF1b n, %</th>
<th>S-gene n, %</th>
<th>Amino acid substitution</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td></td>
<td></td>
<td></td>
<td>N177S 1 (0.52)</td>
<td>-</td>
<td>-</td>
<td>1</td>
</tr>
<tr>
<td>A.23.1</td>
<td></td>
<td></td>
<td></td>
<td>M575I 3 (1.56)</td>
<td>-</td>
<td>P681R 2 (1.04)</td>
<td>46</td>
</tr>
<tr>
<td>B.1</td>
<td>Q71L 1 (0.52)</td>
<td>A385 1 (0.52)</td>
<td>T361 1 (0.52)</td>
<td>K123N 2 (1.04)</td>
<td>K335R 2 (1.04)</td>
<td>P314L 4 (2.08)</td>
<td>D614G 6 (3.13)</td>
</tr>
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<td></td>
<td></td>
</tr>
<tr>
<td>B.1.1.1</td>
<td></td>
<td></td>
<td></td>
<td>R20K 1 (0.52)</td>
<td></td>
<td>P314L 1 (0.52)</td>
<td>D614G 1 (0.52)</td>
</tr>
<tr>
<td>B.1.1.74</td>
<td></td>
<td></td>
<td></td>
<td>R20K 9 (4.69)</td>
<td>E131A 9 (4.69)</td>
<td>P314L 9 (4.69)</td>
<td>D614G 9 (4.69)</td>
</tr>
<tr>
<td>B.1.1.21</td>
<td></td>
<td></td>
<td></td>
<td>S37P 1 (0.52)</td>
<td>E131A 3 (1.56)</td>
<td>P314L 3 (1.56)</td>
<td>D614G 3 (1.56)</td>
</tr>
<tr>
<td>B.1.351</td>
<td>P71L 1 (0.52)</td>
<td>A142V 1 (0.52)</td>
<td>T205I 1 (0.52)</td>
<td>T265I 149 (77.60)</td>
<td>K165N 149 (77.60)</td>
<td>V1649F 149 (77.60)</td>
<td>D614G 149 (77.60)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>T333M 32 (16.76)</td>
<td>E1843D 31 (16.15)</td>
<td>T151I 40 (20.83)</td>
<td>A701V 149 (77.60)</td>
</tr>
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<td></td>
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<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>C.1</td>
<td></td>
<td></td>
<td></td>
<td>T1246I 1 (0.52)</td>
<td>T1246I 1 (0.52)</td>
<td>T1246I 1 (0.52)</td>
<td>D614G 1 (0.52)</td>
</tr>
<tr>
<td>C.2</td>
<td>M109I 2 (1.04)</td>
<td></td>
<td></td>
<td>T1246I 14 (7.29)</td>
<td>T1246I 14 (7.29)</td>
<td>T1246I 14 (7.29)</td>
<td>D614G 15 (7.81)</td>
</tr>
</tbody>
</table>

Note: Numbers in parentheses indicate the percentage of mutations or substitutions.
to the human. However, they are associated with a disorder in the linker region of the nucleoprotein, they have no impact on the diagnostic target of the RT-PCR kits. However, studies in countries that include Australia, Denmark, Saudi Arabia, the UK and the USA is isolates using obtaining the SNP C29200T abolished N2 detection on the Gene Xpert assay. The C2 and C1 lineages shared these N-gene-non-synonymous mutations with the common mutation on positions C14408T(P314L) and A23403G(D614G) of the ORF1b and S-gene. However, phylogenetic analysis with international C2 lineages showed that they were interspersed, suggesting that Zimbabwe was a possible source.

The A23.1 lineage, which was first identified in Uganda, had the non-synonymous mutations N202S and R203K on the N-gene and a unique P681R substitution on the S-gene, which was also found in the Delta variant in India's second wave and the new VOC B1.1.529 in SA. It is located in the furin cleavage site and could increase the rate of S1-S2 cleavage, resulting in better fusogenecity and pathogenicity. The mutations exhibited by the sequenced samples indicated the evolution of SARS-CoV-2 into an efficient super bug regarding diagnostic target dropouts, viral fitness, spread and immune escape.

Conclusion

The Zimbabwe COVID-19 second wave was characterised by nine lineages, with the SA B.1.351 lineage dominating the sequenced samples, constituting >75% of the circulating lineages. There were over 3,000 mutations on the diagnostic genes, with the B.1.351 lineages contributing almost two-thirds of the mutations and the S-gene being the diagnostic gene with the most mutations. The E-gene of all the nine lineages was the least mutated diagnostic gene, with only 2/9 of the lineages possessing a common mutation P71L on B.1.351 and B.1. A different amino acid glutamine is replaced by the same amino acid leucine. These mutations confer different attributes to the virus evolution process, ranging from a high viral load, immune evasion and spreading efficiency to antibody neutralisation. However, they had no impact on the diagnostic kit used during the second wave in Zimbabwe. Regular and efficient surveillance systems to detect circulating variants and mutations need to be put in place by the country to detect any potential harmful mutation so that public and social health measures (PShM) are put in place to arrest the pandemic.

Declaration. This article partially fulfils the requirements of CN’s MPhil degree.

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Conflicts of interest. None.