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Genomic and epidemiologic surveillance of SARS-CoV-2 in Southern Brazil and identification of a new Omicron-L452R sublineage

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A B S T R A C T
Recently, SARS-CoV-2 Omicron variant (B.1.1.529) was first identified in Botswana in November 2021. In a short period of time, this highly mutated variant replaced the previous dominant Delta variant, causing an exponential increase in the number of COVID-19 cases, resulting in a new wave of pandemic. This current research article aims to analyze and summarize information about the genetic characteristics, amino acid mutations and epidemiological data providing scientific findings to enrich the SARS-CoV-2 knowledge. More importantly, we describe here, for the first time, the identification of a new Omicron variant of concern: Omicron-L452R in Brazil.

1. Introduction

In November 2021, researchers in Botswana and South Africa identified a new SARS-CoV-2 variant through whole-genome sequencing (WGS), posteriorly announced by WHO as a Variant of Concern (on November 26, 2021) and named as Omicron (B.1.1.529) (WHO, 2021). Omicron has spread rapidly, increasing COVID-19 cases throughout the world (Mohapatra et al., 2022). On December 25, 2021, over 108 countries reported confirmed cases of Omicron variant isolate. In Brazil, the first description was reported a few weeks after the first case in South Africa, from an airplane passenger that arrived in São Paulo State from South Africa, in late November.

Actually, the Omicron (21M) VOC is divided into six subvariants: or BA.1 (21K), BA.2 (21L), BA.4 (22A), BA.5 (22B), BA.2.12.1 (22C) and BA.2.75 (22D). Since the beginning of the pandemic, the SARS-CoV-2 genome has been rapidly developing, mostly due to the inherent polymerase mistakes and host immune selection factors (Harvey et al., 2021). Different mutations in the spike protein raises concerns and Omicron variant is the most mutated SARS-CoV-2 variant, presenting more than 60 mutations in its genome; of which, 32 were in the receptor binding domain (RBD) of the spike protein (Tsang et al., 2022; Wang and Cheng, 2022).

Compared with previous variants, Omicron is known for decreased hospitalization rates and less severe disease (Maslo et al., 2022). The reduced viral fusogenicity, associated with low pathogenicity in SARS-CoV-2 patients, is probably one of the major reasons but the mechanism is still unclear (Motozono et al., 2021; Rajah et al., 2022). A possible cause of the reduced fusogenicity is related to the S:L452R mutation, present in the Delta variant but absent in Omicron. A recent study reported that a Omicron-L452R mutant generated; displayed

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increased fusogenicity and infectivity mediated by enhancing the cleavage of the spike protein (Zhang et al., 2022).

About transmissibility, Omicron presents a high rate when compared to Delta, which was linked to its immune evasion properties (Rossler et al., 2021). This resulted in Omicron overtaking Delta in areas where community transmission occurs (Gularte et al., 2022; Torjesen, 2021). Tracking SARS-CoV-2 variants through sequencing becomes a key part of a better understanding of viral evolution, especially after the beginning of the vaccination programs. Therefore, the main goal of this work was to describe the replacement of Delta by Omicron variant in Rio Grande do Sul (RS) state, in southern Brazil. Furthermore, through the genomic viral surveillance, we found, for the first time to our knowledge, a new Omicron-related lineage that is circulating in south Brazil, the Omicron-L452R variant.

2. Material and methods

A total of 152 SARS-CoV-2 complete genomes were sequenced through Illumina MiSeq platform at Laboratório de Microbiologia Molecular - Fe芙ale University, one of the institutions linked to the Corona-omics.BR-MCTI Network, a relevant genomic surveillance initiative in Brazil. Naso-oropharyngeal swab samples were received from suspected patients of COVID-19 at Laboratório de Microbiologia Molecular of Universidade Fe芙ale. After the diagnosis confirmation, a pre-selection was carried out, choosing samples that presented cycle threshold value (Ct) below 25. The commercial MagMAX™ CORE Nucleic Acid Purification Kit (Applied biosystems™, Thermo Fisher Scientific, Wal-tham, MA, USA) kit was used to perform viral RNA extraction using the automated equipment KingFisher™ Duo Prime (Thermo Fisher Scientific™). As previously described (da Silva et al., 2021), viral genome sequencing and phylogenetic analysis were carried out. Briefly, reverse transcription reaction was carried out in RNA extracted by SuperScript IV reverse transcriptase kit (Thermo Fisher Scientific, Waltham, MA, USA). Following the manufacturer instructions (QIAGEN, Hilden, Germany), viral genome library preparation was carried out using the QIAseq SARS-CoV-2 Primer Panel paired for library enrichment and QIAseq FX DNA Library UDI Kit. Also, Illumina MiSeq platform (Foster City, CA, USA), using MiSeq Reagent Kit v3 (600-cycle) was used.

FASTQ files were input into the Illumina BaseSpace and consensus sequences were assembled using DRAGEN software with alignment to the reference SARS-CoV-2 sequence (NC_045512.2), minimum depth 10, minimum allele frequency 0.5, and genome at 5X coverage. The Geneious Prime™ suite was used for genome annotation, editing, and mapping the sequences against the reference sequence hCoV-19/ Wuhan/WIV04/2019 (EPI_ISL_402124) available in the Epicov database of Global Initiative on Sharing Avian Influenza Data (GISAID) (Shu and Mccaeula, 2017). PANGO and Nextstrain lineage assignments were applied to characterize the consensus sequences. Phylogenetic tree was constructed including all SARS-CoV-2 complete genomes available from GISAID through the Nextclade tool on Nextstrain server (Hadfield et al., 2018). Mutational profile sequences and signatures were also analyzed in order to better understand the lineage sequences. RDP v.4.101 software was used to evaluate the presence of recombinants among the sequences obtained (Martin et al., 2015).

3. Results

3.1. Epidemiological data

Performing an epidemiological analysis of the sequences generated by our laboratory Delta variant represented 100% of frequency (between October and November 2021). In December of the same year, the first Omicron variant detections were performed and less than a month later (January 2022), the scenario reversed. Delta variant detection considerably decreased (to 6% of all samples analyzed) while Omicron represented 94% of the samples. In February 2022, Delta was not detected anymore, and Omicron represented 100% of the sequences until now (March, 2022) (Fig. 1a).

Considering these variant fluctuations, we compared the number of cases and deaths according to the same epidemiological weeks. Between October and December 2021, when the Delta variant was dominant, the number of cases was low and stable with decreasing the number of deaths. From the introduction of the Omicron variant (in the end of December), the number of cases and deaths started to exponentially increase, and a major COVID-19 wave was observed in RS state. After the peak, the number of the cases and deaths started to decline in January and February 2022, respectively (Fig. 1b).

3.2. Genetic analysis

A total of 152 SARS-CoV-2 complete genomes were retrieved through Illumina MiSeq. The samples were collected between late November, 2021 and mid-March, 2022 from patients in RS state, Brazil. Regarding genetic characterization, the sequences were aligned with complete SARS-CoV-2 genomes of different lineages through the NextClade online tool (https://clades.nextstrain.org/) and the phylogenetic tree was inferred (Fig. 1a).

3.3. Mutational profile

A complete mutational profile was performed, describing the amino acid mutations found in complete SARS-CoV-2 coding regions and comparing the Omicron sublineages. Mutation analysis at the Spike (S) protein revealed the presence of the Omicron signatures. The most frequent S protein mutations shared between the two Omicron major clades 21K (BA.1, BA.1.1, BA.1.14, BA.1.15, BA.1.17, BA.1.19 and BA.2) and 21L (BA.2) includes: G339D, K417N, N440K, S477N, T478K, E484A, Q493R, Q498R, N501Y, S494C, D614G, H655Y, N676K, P681H, N764K, D794Y, Q954H and N696K. Analyzing only 21K cluster, this group shared additional mutations not present in 21L clade: A67V, T95I, Y145D, L212I, S371L, S373P, S375F, G446S, G496S, T547K, N856K and L981F (Fig. 3). Mutations in other viral proteins and their prevalence are also shown in Fig. 3. The major difference between BA.1 and BA.1.1, is that the BA.1.1 carried an additional S:R346K mutation, also described in our sequences. Additional acquired mutations among the sequences were restricted to few sequences and complete analysis could be observed in Table 1.

3.4. Omicron-L452R

Two sequences, classified as BA.1 Omicron sub lineage and named as
hCoV-19/Brazil/LMM69880 (Gisaid accession number EPI_ISL_11514450; collection date: 2022/01/18) and hCoV-19/Brazil/LMM71052 (Gisaid accession number EPI_ISL_11514436; collection date 2022/02/07), draw attention to the presence of a Delta signature S:L452R mutation. Both sequences presented Omicron signatures mutations sharing ORF1a:L2084I, ORF1a:I3758V, ORF1b:P314L, ORF1b: I1566V, ORF9b:P10S, N:P13L, N:R203K, N:G204R, S:L452R, S:T478K, S:D614G, S:H655Y, S:N856K, S:Q954H and S:N764K. Additional uncommon mutations were also found in hCoV-19/Brazil/LMM69880/2022 (ORF1b:P1570S and ORF7a:S83Y) and hCoV-19/ Brazil/LMM71052/2022 (ORF1a:F1501S, ORF1a:V1535A, ORF1a: F3624, ORF1a:M3626T, ORF1a:F3628P and ORF1b:F2283L). No evidence of recombination was found for the analyzed sequences.

These patients presented different clinical presentations. The patient LMM69880, 60 years old, body mass index (BMI) 34 with incomplete vaccine protocol (single Oxford/AstraZeneca Covid-19 vaccine in 2021/06/18) dose presented mild symptoms with no complications. The patient LMM71052, 43 years old, BMI 37 and two Pfizer vaccine doses (last dose: 2021/10/18) had severe symptoms and was hospitalized requiring mechanical ventilation. These patients were respectively infected approximately six and four months after vaccination. Our virus genome surveillance effort found a new variant of concern (VOC), the Omicron-L452R, which should be closely monitored.

4. Discussion and conclusion

Brazil faced the fourth wave of COVID-19 cases, with the highest peak since the beginning of the pandemic. Analyzing data from the RS health authorities, comparing December 2021 and January 2022, the confirmed case numbers increased 2.572.4% (SES-RS, 2022). It’s possible to observe that the increase COVID-19 cases and deaths from the beginning of December 2021 is synchronous with the entry and predominance of the Omicron variant. Omicron has triggered a massive wave of new infections and re-emergent outbreaks worldwide. This
variant spreads more easily than the original SARS-CoV-2 virus and other VOCs, with an expressively higher transmissibility; and also can evade natural and vaccine-induced immunity better than its predecessor variants (Mohapatra et al., 2022; Rössler et al., 2021; Torjesen, 2021). The variant established in a short period of time, which was also observed in other countries, contaminating an expressive number of individuals in a short period of time. Even the apparently milder severity, especially in fully vaccinated individuals, the public health impact should be strongly considerable since the cases increased exponentially.

The SARS-CoV-2 variant Omicron sublineages has spread rapidly in RS state, and a diversification was reported herein, with BA.1 and BA.1.1 (21K) being the most frequent ones. As observed in our study, according to covariants.org, Omicron has overtaken the Delta and has spread rapidly to in Europe, Asia, North and South America. The ability of 21K Omicron sublineages to replace the previously predominant Delta variant has been credited to immune escape rather than a higher intrinsic transmissibility (Eggtink et al., 2022; Lyngse et al., 2021), but BA.2 has been shown to be even more transmissible than BA.1 (Lyngse et al., 2022). Despite this, the presence of BA.2 was very low in the samples analyzed in this study, maybe due to the still recent entrance into the RS state.

Mutational SARS-CoV-2 analysis have been important since the beginning of the pandemic, especially for the S glycoprotein, which mediates virus attachment to ACE2 receptor, membrane fusion, and entry into the host cell, and also acts as a primary target for neutralizing antibodies elicited by the host immune response (Walls et al., 2020). Omicron sublineages have mutations in common and also specific signatures, as shown in the results section. The following mutations S:K417N, S:T478K, S:N501Y, S:D614G, S:H655Y and S:P681H, detected in all Omicron sublineages analyzed herein, were also described in other VOCs (Alpha, Beta, Gamma and Delta) and have been previously related to increase viral binding, immune evasion and high transmissibility (Arora et al., 2021; Greaney et al., 2021; Harvey et al., 2021). L452R is related to increased SARS-CoV-2 fusogenicity and infectivity (Motozono et al., 2021). In our study, we report a natural Omicron-L452R in two patients. Through
mild clinical signs. Despite this, it is important to note that the severity of the disease is multifactorial, and, to link L452R with specific clinical them presented serious COVID-19 complications, and the other, only infected Omicron-L52R reported in this study were divergent, one of

this could be the reason that the big COVID-19 wave showed enhanced fusion activity and increased ability to infect lung tissues of humanized mice (Zhang et al., 2022) and SARS-CoV-2 path genetic engineering, a recent reported that an Omicron-L452R mutant associated with Omicron infections, less hospitalization rates and clin

Table 1

| Region | Common Omicron mutations | Unique mutations on BA.1 | Unique mutations on BA.1.1 | Unique mutations on BA.1.1.14 | Unique mutations on BA.1.1.15 | Unique mutations on BA.1.1.17 | Unique mutations on BA.1.9 | Unique mutations on BA.2 |
|--------|--------------------------|--------------------------|---------------------------|-------------------------------|-------------------------------|-------------------------------|----------------------------|-------------------------|
| S      | A67V, T95L, Y145D, L121I, G339D, S371L, S373P, S375F, K417N, N440K, G446S, S477N, T478K, E484A, Q493R, G496S, Q498R, N501Y, V506H, T547K, D614G, H655Y, N679K, P681H, N764K, D796Y, N856K, Q954H, N969K, L981F | K856R, L2084I, A2710T, T3255I, P3395H, I3758V | R24C, T233M, V675A, G1100S, H1113Y, G1307V, L1334V, F15015, V1535A, T1682I, T1854I, E2904D, L3027F, T3090I, L3116F, L3201F, L3919I | G379E, Y621C, E743K, T1605S, T1788M, S1857L, P2079S, R3066K, L3606F, L3612S, A3615V, F3624L, M3626F, F3628P | T4174I*, E622D, R1628S*, L3116F*, R1638S*, T2906I, T2153I, E1245K, E932G, I3456V | L3124S, P1803S, L3606F | L3125R, T842I*, G1307S*, L3027F*, T3090I*, L3201F* | |
| ORF1a  | K856R, L2084I, A2710T, T3255I, P3395H, I3758V | P1055*, A1302Y, K2421N, P1570S, F2283L | T11A | C130S, D1735Y, G1014S, R1315C* |
| ORF7a  | S83Y | L106F*, S165F*, T32I | Y154N* |
| ORF3a  | E191 | A85S* | I2R2T |
| ORF9b  | P10S | | |
| E      | E791 | | |
| M      | MQ19E, MD3G, MA63T | | |
| N      | NP131, NR203K, NG204R | P67S, E378D* | R68P, T135A, D402Y | Q163K, D343G | S413R |

genetic engineering, a recent reported that an Omicron-L452R mutant showed enhanced fusion activity and increased ability to infect lung tissues of humanized mice (Zhang et al., 2022) and SARS-CoV-2 pathogenicity in patients is strongly related to fusogenic capability (Motzono et al., 2021). This could be the reason that the big COVID-19 wave associated with Omicron infections, less hospitalization rates and clinical severity were observed (Maslo et al., 2022). The clinical signs of the infected Omicron-L52R reported in this study were divergent, one of them presented serious COVID-19 complications, and the other, only mild clinical signs. Despite this, it is important to note that the severity of the disease is multifactorial, and, to link L452R with specific clinical signs it’s not possible without further studies as only two Omicron-L453R variants were identified in this study.

The BA.4 and BA.5 Omicron sublineages, that emerged after the sequences reported herein, also presented the S:L452R mutation (reported as BA.4 and BA.5 signature), which was not observed in our sequences. Thus, our sequences probably represented a transition point between lineages, with a mutation that later stabilized in new Omicron BA.4 and BA.5.

SARS-CoV-2 overall rate of evolution, estimated by preliminary molecular clock in 2020 was 8 × 10^-4 substitutions/site/year, which equates to 24 substitutions per year. The current global estimate including multiple variants of concern/interest suggests a similar rate of approximately 25.941 substitutions per year (derived from a global Nextstrain build, https://nextstrain.org/ncov/gisaid/global, accessed April 1st, 2022). Our Omicron analysis presented a higher clock rate of 28.125 substitutions per year, showing a higher substitution rate than the majority of other sequences.

In summary, we described the replacement of the Delta lineage by Omicron in a short period of time and the presence of seven sublineages (BA.1, BA.1.1, BA.1.1.14, BA.1.1.15, BA.1.1.17 and BA.2), mapping their mutational profile. Furthermore, we report the first natural infections with Omicron-L452R variant, an important S protein mutation previously characteristic only of the Delta VOC. Our study has provided a comprehensive investigation concerning the epidemiological and genetic characteristics of the major wave caused by Omicron in RS state, Southern Brazil, and contributed to important scientific findings enriching the knowledge on SARS-CoV-2.

Ethical aspects

Project approved by the Research Ethics Committee (CEP) at Feevale University. Process number: CAAE: 33202820.7.1001.5348.

CRediT authorship contribution statement

Mariana Soares da Silva: Conceptualization, Formal analysis, Investigation, Writing – original draft. Juliana Schons Gularte: Conceptualization, Formal analysis, Investigation. Micheli Filippi: Conceptualization, Formal analysis, Investigation. Meriane Demoliner: Conceptualization, Formal analysis, Investigation. Viviane Girardi: . Ana Cristina Sbaraini Mosena: Formal analysis, Investigation. Victionary Malhaycha de Abreu Goes Pereira: Formal analysis, Investigation. Alana Witt Hansen: Formal analysis, Investigation. Frederico Fleck: Funding acquisition, Resources. Andrea Gurgel Rodrigues de Almeida: Formal analysis, Investigation. Juliana Deise Nunes Weber: Formal analysis, Investigation. Ana Cristina Vieira Vilela: Formal analysis, Investigation. Luiz Amorim Filho: Formal analysis, Investigation. Fernanda Spotto: Formal analysis, Investigation. Viviane Girardi: . Ana Cristina Sbaraini Mosena: Formal analysis, Investigation. Victionary Malhaycha de Abreu Goes Pereira: Formal analysis, Investigation. Alana Witt Hansen: Formal analysis, Investigation. Frederico Fleck: Funding acquisition, Resources. Andrea Gurgel Rodrigues de Almeida: Formal analysis, Investigation. Juliana Deise Nunes Weber: Formal analysis, Investigation. Ana Cristina Vieira Vilela: Formal analysis, Investigation. Luiz Amorim Filho: Formal analysis, Investigation. Fernanda Spotto: Formal analysis, Investigation.
Supervision.

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.

Data availability

Data will be made available on request.

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Supplementary materials

Supplementary material associated with this article can be found, in the online version, at doi: 10.1016/j.virusres.2022.198597.

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