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Genomic monitoring of the SARS-CoV-2 B1.1.7 (WHO VOC Alpha) in the Sao Paulo state, Brazil

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A B S T R A C T

The SARS-CoV-2 alpha VOC (also known as lineage B.1.1.7) initially described in the autumn, 2020 in UK, rapidly became the dominant lineage across much of Europe. Despite multiple studies reporting molecular evidence suggestive of its circulation in Brazil, much is still unknown about its genomic diversity in the state of São Paulo, the main Brazilian economic and transportation hub. To get more insight regarding its transmission dynamics into the State we performed phylogenetic analysis on all alpha VOC strains obtained between February and August 2021 from the São Paulo state Network for Pandemic Alert of Emerging SARS-CoV-2 variants. The performed phylogenetic analysis showed that most of the alpha VOC genomes were interspersed with viral strains sampled from different Brazilian states and other countries suggesting that multiple independent alpha VOC introductions from Brazil and overseas have occurred in the São Paulo State over time. Nevertheless, large monophyletic clusters were also observed especially from the Central-West part of the São Paulo State (the city of São Paulo Ribeirão Preto).
1. Introduction

The SARS-CoV-2 pandemic has been a source for the rapid emergence and spread of variants of concern (VOCs) with major genomic changes (Davies et al., 2021). Those VOCs appear to have the ability to substitute the circulating ancestral strains, exerting thus enormous pressure on the healthcare systems as reported worldwide (Faria et al., 2021; Choudhary et al., 2021). SARS-CoV-2 variants are typically characterized by constellations of specific mutations which can enhance their transmissibility, immunological escape and phenotypic characteristics (Volz et al., 2021). The SARS-CoV-2 Alpha VOC (B.1.1.7/GR/501Y.V1/20I) which was initially reported in late 2020 in the United Kingdom, rapidly spread across the whole country between November 2020 and January 2021 (Volz et al., 2021) and later in continental Europe (Gaymard et al., 2021; Loconsole et al., 2021) and the USA (Washington et al., 2021). This is a result of the increased transmissibility of the Alpha VOC which is up to 90% more transmissible compared to the predecessor lineages (Davies et al., 2021).

Brazil is currently a world leader in the number of confirmed SARS-CoV-2 cases (21,849,137 confirmed cases by November, 5th, 2021, https://covid.saude.gov.br/) and represents a complex mixture of circulating variants. The first Alpha VOC cases in Brazil were accidentally identified in December 2020 due to an uncommon PCR amplification pattern observed during routine SARS-CoV-2 diagnosis in the city of São Paulo (Claro et al., 2021). The Alpha VOC molecular epidemiology in Brazil is still little examined as the only study which traces the molecular Alpha VOC profile analyzes only 25 complete genomes which were obtained from different Brazilian Federal States (Moreira et al., 2021). Currently, São Paulo State in Brazil demonstrates the highest number of confirmed SARS-CoV-2 cases (4409,404 by November, 5th, 2021; https://www.saopaulo.sp.gov.br/coronavirus/). Due to the high number of SARS-CoV-2, Sao Paulo State Network for Pandemic Alert of Emerging SARS-CoV-2 Variants was implemented with the objective to monitor the VOCs and VOIs in the State. This raised important questions regarding the dissemination and molecular epidemiology of the Alpha VOC in this Brazilian region. From a total number of 21,700 SARS-CoV-2 genomes sequenced by the São Paulo Network for Pandemic Alert of Emerging SARS-CoV-2 Variants and which passed quality control, 1.9% has been classified as alpha VOC but no systematic research regarding this VOC has been performed in the country so far. In this report, we analyzed the Alpha VOC phylogenetics using 366 strains obtained from different Health Districts of the São Paulo State during the genomic surveillance performed by the Sao Paulo State Network for Pandemic Alert of Emerging SARS-CoV-2 Variants.

2. Materials and methods

2.1. Clinical samples

From February to August 2021 the Network for Pandemic Alert of
Emerging SARS-CoV-2 Variants generated 366 full genome sequences which were classified as the Alpha VOC using the PANGO lineages (https://cov-lineages.org/lineages.html). These sequences were obtained from all 17 Regional Health Divisions of the São Paulo State (Fig. 1) as follows: Araçatuba (13 sequences), Baixada Santista (6 sequences), Barretos (5 sequences), Bauru (79 sequences), Campinas (64 sequences), Franca (16 sequences), Metropolitan São Paulo (60 sequences), Marília (17 sequences), Piracicaba (21 sequences), Presidente Prudente (11 sequence), São Jose do Rio Preto (9 sequences), São João da Boa Vista (11 sequences), Sorocaba (10 sequences), Taubaté (22 sequences), Ribeirão Preto (11 sequences) and Araçatuba (11 sequences). The sample representativity from the total number of positive cases per each Regional Health Division varied between 0.2% (10th epidemiological week) to 6.4% (26th epidemiological week). From the beginning of the 20th epidemiological week, the sampling significance was superior of 82% (G^2-Power, Statistical Power Analysis software). SARS-CoV-2 genomeic libraries were generated using the COVIDESeq kit (Illumina, San Diego, CA) following the manufacturer’s specifications. The normalized sample libraries were sequenced using Illumina MiSeq platform using the MiSeq Reagent v.2 Kit (2 × 300 cycles) (Illumina, San Diego, CA, USA).

2.2. Bioinformatic pipeline

The raw sequence data obtained were submitted to quality control analysis using the FastaQC (Andrews, 2010) software version 0.11.8. Trimming was performed using Trimmomatic (Bolger et al., 2014) version 0.39 in order to select the sequences with best quality. Sequences with quality scores > 30 were only used. We mapped the trimmed sequences against the SARS-CoV-2 reference (Genbank refseq NC_045512.2) using BWA (Li, 2013) software and samtools (Li et al., 2013) for read indexing. The mapped files were submitted to refinement with the software Pilon (Walker et al., 2014) in order to obtain the indels and insertions in the most correct way possible, afterwards the trimmed sequences were subjected to a remap against the genome refined by Pilon. Finally, we use bcftools (Narasimhan et al., 2016) for variant calling, and seqtk (Shen et al., 2016) for creation of consensus genomes. Positions covered by fewer than 10 reads (DP<10) and bases of quality lower than 30 were considered a gap in coverage and converted to Ns. Coverage values for each genome were calculated using samtools v1.12 (Li et al., 2009). We assessed genome consensus sequence quality using Nextclade v0.8.1 (https://clades.nextstrain.org).

2.3. Phylogenetic analysis

The sequences which were generated in this study were initially classified as belonging to the Alpha VOC using the Pangolin 3.1.11 and pangolLEARN 1.2.56 software (O’Toole et al., 2021). A set of representative Alpha VOC genomes (and associated metadata) across countries, dates, and lineages (curated weekly) was downloaded from GISAID (ShuklaMcCaulley, 2017), including all Brazilian samples for a set of 1103 genomes (247 genomes obtained from Brazil, 490 world genomes and 366 from our study). Only genomes > 29,000 bp and <1% of ambiguities were retrieved, low-quality genomes (> 10% of ambiguous positions) were excluded. Sequence alignment was performed using MAFFT v7.453 (Katoh & Standley, 2013) with subsequent realignment of misaligned stretches done in Aliview (Larsson, 2013). The beginning and end of the alignment were removed to avoid these regions of lower coverage (Larsson, 2013). Maximum Likelihood (ML) trees were estimated using IQ-TREE v1.6.12 (Nguyen et al., 2015) applying the maximum likelihood algorithm with statistical support of ultraraf bootstrap with 1000 replicates. We inferred time-scaled trees and rooted these with least-squares criteria and the evolutionary rate of > 1.1 × 10^{-2} substitutions/site/year estimated by S. Duchene et al., 2020 using the TreeTime v.1 software (Sagulenko et al., 2018).

3. Results

The inferred phylogenetic history of the Alpha VOC circulating in the state of São Paulo using a number of representative reference strains obtained from Brazil and worldwide showed that our isolates were randomly interspersed with other Alpha VOCs and grouped predominantly with Alpha VOC strains circulating in Brazil and Europe (Fig. 1). Additionally, we observed several large monophyletic clusters composed of Alpha VOC strains obtained from different regions of the state including the metropolitan region of São Paulo and especially Bauru. The largest observed cluster belonged to sequences obtained from the Central-West part of the state of São Paulo and principally from the city of Bauru. Interestingly from this location we obtained the largest number of strains (total number of 79 Alpha VOC, 21.6%) which may be related to sustained Alpha VOC transmission and dissemination in this region. Another 60 genomes (16.4%), which were sequenced from the metropolitan region of the state of São Paulo formed also several smaller monophyletic clusters which might be also related to a number of independent introductions of this VOC in this location which is additionally justified by the importance of the city of São Paulo as a national and international travel, economic and demographic hub (Fig. 2).

4. Discussion

The performed study uses considerable number of complete Alpha VOC genomes obtained from the state of São Paulo in order to give detailed information on the dissemination and molecular epidemiology of this important VOC in Southeast Brazil. All of the examined sequences were obtained between February-August 2021 and this roughly corresponded to the second SARS-CoV-2 wave which hit the São Paulo State between December and February 2021. The second SARS-CoV-2 pandemic wave was further related with implementation of restriction measures and lockdowns along the State municipalities and coincided with the wide dissemination of the gamma VOC (P.1) nationwide.

Although in the reconstructed phylogenetic tree the sequences analyzed in this study were randomly interspersed with other Alpha VOC strains, several large monophyletic clusters composed of sequences obtained from different geographic regions of the state were observed (observe boxes Fig. 2). Such a local monophyletic clustering might be related to multiple introductions of the Alpha VOC in specific areas of the São Paulo State. In addition to the multiple introductions, we found phylogenetic evidence for a local sustained transmission of this VOC in several geographic regions of the State and especially the Central-West part (city of Bauru) and metropolitan areas of the city of São Paulo. Interestingly the largest cluster containing the largest number of sequences belonged to the city of Bauru, which is located ~ 340 km from the city of São Paulo. We suggest that the introduction of Alpha VOC in this region led to sustained transmission and wide dissemination of this VOC, the detection of which was only possible due to the massive sequencing provided by the Butantan Network for Pandemic Alert of SARS-CoV-2 variants. Higher number of Alpha VOC sequences were also obtained from the metropolitan area of the city of São Paulo, but these were interspersed in several smaller clusters along the phylogenetic tree. Such a clustering might be related to multiple Alpha VOC introductions in the city of São Paulo which is justified with the national importance of this metropolis. São Paulo city is served by the largest international airport of the country and is the most important economical, port (port of Santos) and transport center of South America. Given the intense economic, transportation and communication characteristics it is possible that different Alpha VOC introductions have occurred along time which lead to the sustained Alpha VOC transmission in São Paulo city. This finding is similar to other studies which show the contribution of the large urban and densely populated areas as principal hubs for introduction, diversification and dissemination of SARS-CoV-2 (Alteri et al., 2021; Kraemer et al., 2021).

Despite that the alpha VOC strains in this study were obtained from
districts that encompass the whole State territory, the proportion of the alpha VOC lineages was not prominent especially compared to Europe (Davis et al., 2021). We could not explain why the Alpha VOC in Brazil did not reach epidemic proportions. We believe that probably, the most important factor was related to a background of a fully dominant P.1 VOC in Brazil. By the time when Alpha VOC was identified for the first time in São Paulo, Brazil has been already experiencing the second gamma VOC wave and a significant part of the inhabitants were already SARS-CoV-2 immunologically competent which may have not created a favorable environment for Alpha VOC dissemination. Additionally, the substitution of a given SARS-CoV-2 lineage is a complex process which depends on a series of factors including high incidence of other VOC infections, previous SARS-CoV-2 seroprevalence, vaccination rates and human mobility/regional transportation characteristics (Kraemer et al., 2021).

The dissemination profile of the alpha VOC in the São Paulo State was obtained through extensive SARS-CoV-2 molecular surveillance program, the Network for Pandemic Alert of Emerging SARS-CoV-2 Variants. This effort was fundamental for a more profound understanding of VOC and VOI circulation in the State and monitoring the emergence and distribution of novel or underrepresented SARS-CoV-2 variants which also included the alpha VOC. Therefore, SARS-CoV-2 genomic surveillance remains the most important tool which predicts the spread of SARS-CoV-2 VOCs and is a fundamental approach for pandemic response. Moreover, the unification of efforts towards SARS-CoV-2 molecular surveillance will be imperative to understand the
Nucleotide sequence accession numbers

All of the generated sequences were deposited in GISAID (www.gisaid.org). All of the identification numbers are given in a Supplementary File 1.

Ethics approval

This study was approved by the Institutional Ethics Committee of the Faculty of Medicine of Ribeirão Preto (Process CAAE: 50,367,721.7.1001.5440).

Credit statement

ERS, EVS, ME, LPOL, RAB, PDSCM, RLRCCL, JPK, DS, PAA, FASC, MDP, JSLP, VL, VF, SNS, MCE, SCS, SK, ARUJ, GR analyzed the data and wrote the article. MB, PMMG, NNF, RTC, and DTC evaluated the clinical/epidemiological data and reviewed the article. MDP, JCCL, ECM, CAB, and LS wrote the article. MG, LCJA, LLC, RMTG, MJP, CRB, and CRSB evaluated the epidemiological survey. MG, LCJA, LLC, RMTG, MJP, CRB, and CRSB wrote the article. MB, PMMG, NNF, RTC, and DTC supervised this work. All the authors agreed on the content of the final manuscript.

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Supplementary material

Supplementary table 3

Declaration of Competing Interest

The authors declare no competing interests.

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Supplementary material

Supplementary material associated with this article can be found, in the online version, at doi:10.1016/j.ejps.2020.105216.

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