RESEARCH ARTICLE SUMMARY

CORONAVIRUS

The evolving SARS-CoV-2 epidemic in Africa: Insights from rapidly expanding genomic surveillance

Houriiyah Tegally et al.*+

INTRODUCTION: Investment in Africa over the past year with regard to severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) sequencing has led to a massive increase in the number of sequences, which, to date, exceeds 100,000 sequences generated to track the pandemic on the continent. These sequences have profoundly affected how public health officials in Africa have navigated the COVID-19 pandemic.

RATIONALE: We demonstrate how the first 100,000 SARS-CoV-2 sequences from Africa

have helped monitor the epidemic on the continent, how genomic surveillance expanded over the course of the pandemic, and how we adapted our sequencing methods to deal with an evolving virus. Finally, we also examine how viral lineages have spread across the continent in a phylogeographic framework to gain insights into the underlying temporal and spatial transmission dynamics for several variants of concern (VOCs).

RESULTS: Our results indicate that the number of countries in Africa that can sequence the

A Local sequencing facilities **B** Sequencing strategies Sequencing Africa Local Outside Africa 10.000 1000 100 C Number of African SARS-CoV-2 genomes 100.000 Thousands of genomes 30 40 50 60 70 80 90 100 Time to sequence 100,000 genomes (per 10,000) 375 days 47 41 43 36 24 24 29

Expanse of SARS-CoV-2 sequencing capacity in Africa. (A) African countries (shaded in gray) and institutions (red circles) with on-site sequencing facilities that are capable of producing SARS-CoV-2 whole genomes locally. (B) The number of SARS-CoV-2 genomes produced per country and the proportion of those genomes that were produced locally, regionally within Africa, or abroad. (C) Decreased turnaround time of sequencing output in Africa to an almost real-time release of genomic data.

virus within their own borders is growing and that this is coupled with a shorter turnaround time from the time of sampling to sequence submission. Ongoing evolution necessitated the continual updating of primer sets, and, as a result, eight primer sets were designed in tandem with viral evolution and used to ensure effective sequencing of the virus. The pandemic unfolded through multiple waves of infection that were each driven by distinct genetic lineages, with B.1-like ancestral strains associated with the first pandemic wave of infections in 2020. Successive waves on the continent were fueled by different VOCs, with Alpha and Beta cocirculating in distinct spatial patterns during the second wave and Delta and Omicron affecting the whole continent during the third and fourth waves, respectively. Phylogeographic reconstruction points toward distinct differences in viral importation and exportation patterns associated with the Alpha, Beta, Delta, and Omicron variants and subvariants, when considering both Africa versus the rest of the world and viral dissemination within the continent. Our epidemiological and phylogenetic inferences therefore underscore the heterogeneous nature of the pandemic on the continent and highlight key insights and challenges, for instance, recognizing the limitations of low testing proportions. We also highlight the early warning capacity that genomic surveillance in Africa has had for the rest of the world with the detection of new lineages and variants, the most recent being the characterization of various Omicron subvariants.

CONCLUSION: Sustained investment for diagnostics and genomic surveillance in Africa is needed as the virus continues to evolve. This is important not only to help combat SARS-CoV-2 on the continent but also because it can be used as a platform to help address the many emerging and reemerging infectious disease threats in Africa. In particular, capacity building for local sequencing within countries or within the continent should be prioritized because this is generally associated with shorter turnaround times, providing the most benefit to local public health authorities tasked with pandemic response and mitigation and allowing for the fastest reaction to localized outbreaks. These investments are crucial for pandemic preparedness and response and will serve the health of the continent well into the 21st century. ■

*Corresponding author. Email: Tulio de Oliveira (tulio@sun.ac.za); Eduan Wilkinson (ewilkinson@sun.ac.za) †All authors and affiliations appear in the full article online. Cite this article as H. Tegally et al., Science 378, eabq5358 (2022). DOI: 10.1126/science.abq5358



Tegally et al., Science **378**, 42 (2022) 7 October 2022

RESEARCH ARTICLE

CORONAVIRUS

The evolving SARS-CoV-2 epidemic in Africa: Insights from rapidly expanding genomic surveillance

All authors and their affiliations appear at the end of this paper.

Investment in severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) sequencing in Africa over the past year has led to a major increase in the number of sequences that have been generated and used to track the pandemic on the continent, a number that now exceeds 100,000 genomes. Our results show an increase in the number of African countries that are able to sequence domestically and highlight that local sequencing enables faster turnaround times and more-regular routine surveillance. Despite limitations of low testing proportions, findings from this genomic surveillance study underscore the heterogeneous nature of the pandemic and illuminate the distinct dispersal dynamics of variants of concern—particularly Alpha, Beta, Delta, and Omicron—on the continent. Sustained investment for diagnostics and genomic surveillance in Africa is needed as the virus continues to evolve while the continent faces many emerging and reemerging infectious disease threats. These investments are crucial for pandemic preparedness and response and will serve the health of the continent well into the 21st century.

hat originally started as a small cluster of pneumonia cases in Wuhan, China, more than 2 years ago (1) quickly turned into a global pandemic. COVID-19 is the clinical manifestation of severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) infection, and by March 2022, there had been more than 437 million reported cases and more than 5.9 million reported deaths (2). Although Africa accounts for the lowest number of reported cases and deaths thus far, with ~11.3 million reported cases and 245,000 reported deaths as of February 2022, the continent has played an important role in shaping the scientific response to the pandemic with the implementation of genomic surveillance and the identification of two of the five variants of concern (VOCs) (3, 4).

Since it emerged in 2019, SARS-CoV-2 has continued to evolve and adapt (5). This has led to the emergence of several viral lineages that carry mutations that either confer some viral adaptive advantages that increase transmission and infection (6, 7) or counter the effect of neutralizing antibodies from vaccination (8) or previous infections (9-11). The World Health Organization (WHO) classifies certain viral lineages as VOCs or variants of interest (VOIs) based on the potential impact they may have on the pandemic, with VOCs regarded as the highest risk. To date, five VOCs have been classified by the WHO; of these, two were first detected on the African continent (Beta and Omicron) (3, 4, 12) and two (Alpha and Delta) (12, 13) have spread extensively on the continent in successive waves. The remaining VOC, Gamma (14), originated in Brazil and had a limited influence in Africa, with only four recorded sequenced cases.

For genomic surveillance to be useful for public health responses, sampling for sequenc-

ing needs to be both spatially and temporally representative. In the case of SARS-CoV-2 in Africa, this means extending the geographic coverage of sequencing capacity to capture the dynamic genomic epidemiology in as many locations as possible. In a meta-analysis of the first 10,000 SARS-CoV-2 sequences generated in 2020 from Africa (15), several blind spots were identified with regard to genomic surveillance on the continent. Since then, much investment has been devoted to building capacity for genomic surveillance in Africa, coordinated mostly by the Africa Centers for Disease Control (Africa CDC) and the regional office of the WHO in Africa (or WHO AFRO) but also provided by several national and international partners, resulting in an additional 90,000 sequences shared over the past year (April 2021 to March 2022). This makes the sequencing effort for SARS-CoV-2 a phenomenal milestone. In comparison, only 12,000 whole-genome influenza sequences (16) and only ~3700 whole-genome HIV sequences (17) from Africa have been shared publicly, even though HIV has plagued the continent for decades.

Here, we describe how the first 100,000 SARS-CoV-2 sequences from Africa have helped describe the pandemic on the continent, how this genomic surveillance in Africa has expanded, and how we adapted our sequencing methods to deal with an evolving virus. We also highlight the impact that genomic sequencing in Africa has had on the global public health response, particularly through the identification and early analysis of new variants. Finally, we also describe here how the Delta and Omicron variants have spread across the continent and how their transmission dynamics were distinct from the Alpha and Beta variants that preceded them.

Results

Epidemic waves driven by variant dynamics and geography

Scaling up sequencing in Africa has provided a wealth of information on how the pandemic unfolded on the continent. The epidemic has largely been spatially heterogeneous across Africa, but most countries have experienced multiple waves of infection (18–29), with substantial local and regional diversity in the first wave and to a lesser extent in the second wave, followed by successive sweeps of the continent with Delta and Omicron (Fig. 1A). In all regions of the continent, different lineages and VOIs evolved and cocirculated with VOCs and, in some cases, contributed considerably to epidemic waves.

In North Africa (Fig. 1B and fig. S1A), B.1 lineages and Alpha dominated in the first and second waves of the pandemic and were replaced by Delta and Omicron in the third and fourth waves, respectively. Interestingly, the C.36 and C.36.3 sublineages dominated the epidemic in Egypt (~40% of reported infections) before July 2021 when they were replaced by Delta (30). Similarly, in Tunisia, the first and second waves were associated with the B.1.160 lineage and were replaced by Delta during the country's third wave of infections. In southern Africa (Fig. 1C and fig. S1C), we see a similar pandemic profile, with B.1 dominating the first wave; however, instead of Alpha, Beta was responsible for the second wave, followed by Delta and Omicron. Another lineage that was flagged for close monitoring in the region was C.1.2 because of its mutational profile and predicted capacity for immune escape (31). However, the C.1.2 lineage did not cause many infections in the region because it was circulating at a time when Delta was dominant. In West Africa (Fig. 1D and fig. S1B), the B.1.525 lineage caused a large proportion of infections in the second and third waves, where it shared the pandemic landscape with the Alpha variant. As with other regions on the continent, these variants were later replaced by the Delta and then the Omicron VOCs in successive waves. In Central Africa (Fig. 1E and fig. S1D), the B.1.620 lineage caused most of the infections between January and June 2021 (32) before systematically being replaced by Delta and then Omicron. Lastly, in East Africa (Fig. 1F and fig. S1E), the A.23.1 lineage dominated the second wave of infections in Uganda (33) and much of East Africa. In all of these regions, minor lineages such as B.1.525, C.36, and A.23.1 were eventually replaced by VOCs that emerged in later waves.

Finally, we directly compared the official recorded cases in Africa with the ongoing SARS-CoV-2 genomic surveillance data (GISAID date of access: 31 March 2022) for a crude estimation of the variants' contributions to cases. We observe that Delta was responsible for an epidemic wave between May and October 2021

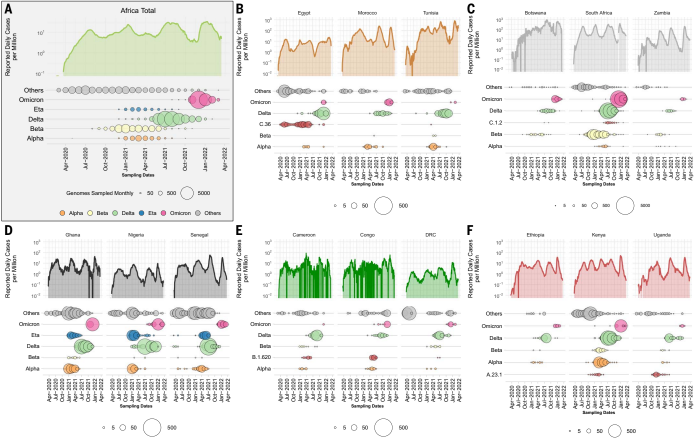


Fig. 1. Epidemiological progression of the COVID-19 pandemic on the African continent. (**A**) Total reported new case counts per million inhabitants in Africa (data source: Our World in Data; log-transformed) along with the distribution of VOCs, the Eta VOI, and other lineages through time (the size of each circle is proportional to the number of genomes sampled per month for each category). (**B** to **F**) Breakdown of reported new cases

per million (data source: Our World in Data; log-transformed) and monthly sampling of VOCs, regional variant, or lineage of interest and other lineages for three selected countries for North, southern, West, Central, and East Africa, respectively. For each region, a different variant or lineage of interest is shown, relevant to that region (C.36, C.1.2, Eta, B.1.620, and A.23.1, respectively).

(Fig. 1A) and had the greatest impact on the continent, with almost 34.2% of overall infections in Africa possibly attributed to it. Beta was responsible for an epidemic wave at the end of 2020 and beginning of 2021 (Fig. 1A), with 13.3% of infections overall attributed to it. Notably, Alpha, despite being predominant in other parts of the world at the beginning of 2021, had only minimal importance in Africa, accounting for just 4.3% of infections. At the time of writing, the Omicron VOC had contributed to 21.6% of the overall number of sequenced infections. At this time, the Omicron wave was still unfolding globally and in Africa with the expansion of several sublineages (34), such that its full impact is vet to be determined. However, because of increased population immunity (35) from SARS-CoV-2 infection and vaccination (fig. S2), the impact of Omicron on mortality has been less in comparison to the other VOCs, as can be observed by the relatively low death rate in South Africa during the Omicron wave (36). The findings from mapping epidemiological numbers onto genomic surveillance data are reliable as far as the proportional scaling of genomic sampling across Africa with the size and timing of epidemic waves [fig. S3; model estimate (b) = 0.011, standard error (SE) = 0.001, $p < 2 \times 10^{-16}$].

This comes with the obvious caveats that testing and reporting practices have varied widely across the continent along with genomic surveillance volumes throughout the pandemic. Countries in Africa with reported data have tested in proportions from as little as 0.1 daily tests per million population to more than 1000 tests per million (fig. S4). Some countries have consistently tested at high proportions, for example, South Africa, Botswana, Morocco, and Tunisia. Incidentally, these countries have also generally reported more cases per million population, providing an indication that recorded low incidences in other parts of the continent have been underestimates due to low testing rates. However, even for these countries, epidemic numbers are certainly underrepresented and underdetected, given that in several time frames, test positivity rates were still on the higher end, approaching or exceeding 20% (fig. S4), and as concluded by seroprevalence surveys and estimates of true infection burdens in Africa (37, 38). Findings of attributing case numbers of variants must therefore be interpreted in the context of this limitation but can nevertheless provide a qualitative overview of the spatial and temporal dynamics of VOCs in relation to epidemic progression in Africa.

The African regional (table S1) and country-specific (table S2) NextStrain builds also clearly support the changing nature of the pandemic over time. From these builds, we observe a strong association of B.1-like viruses circulating on the continent during the first wave. These "ancestral" lineages were subsequently replaced by the Alpha and Beta variants, which dominated the pandemic landscape during the second wave and were later replaced by the Delta and Omicron variants during the third and fourth waves.

Optimizing surveillance coverage in Africa

By mapping and comparing the locations of specimen sampling laboratories to the sequencing

laboratories, a number of aspects regarding the expansion of genomic surveillance on the continent became clear. First, even though several countries in Africa started sequencing SARS-CoV-2 in the first months of the pandemic, local sequencing capacity was initially limited. However, local sequencing capabilities slowly expanded over time, particularly after the emergence of VOCs (Fig. 2A). The fact that almost half of all SARS-CoV-2 sequencing in Africa was performed using the Oxford Nanopore Technology (ONT), which is relatively low-cost compared with other sequencing technologies and better adapted to modest

laboratory infrastructures, illustrates one component of how this rapid scale-up of local sequencing was achieved (fig. S5). Yet, to rely only on local sequencing would have thwarted the continent's chance at a reliable genomic surveillance program. At the time of writing, 52 of 55 countries in Africa had SARS-CoV-2

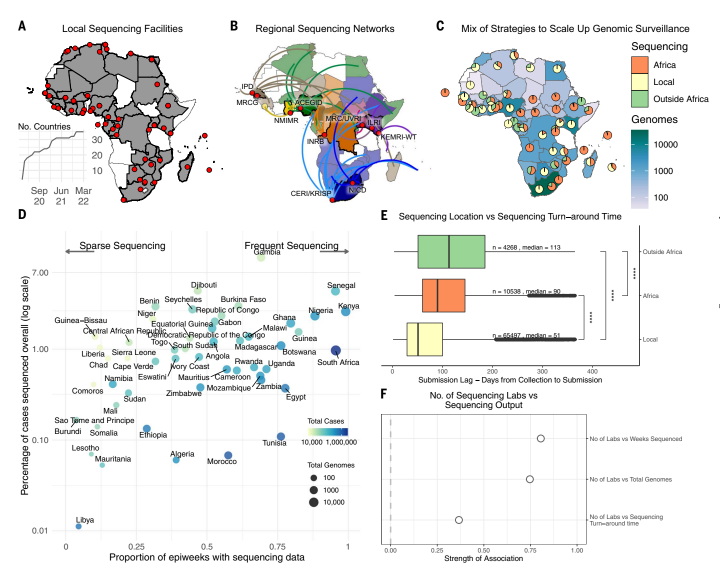


Fig. 2. Sequencing strategies and outputs in Africa. (A) Geographical representation of all countries (shaded in gray) and institutions (red dots) in Africa with their own on-site sequencing facilities. The inset graph shows the number of countries in Africa that are able to carry out sequencing locally over time. (B) Key regional sequencing hubs and networks in Africa showing countries (shaded in bright colors) and institutions (red dots) that have sequenced for other countries (shaded in corresponding light colors and linking curves) on the continent. ACEGID, African Centre of Excellence for Genomics of Infectious Diseases; CERI, Centre for Epidemic Response and Innovation; KEMRI-WT, Kenya Medical Research Institute—Wellcome Trust; KRISP, KwaZulu-Natal Research Innovation and Sequencing Platform; ILRI, International Livestock Research Institute; INRB, Institut National de Recherche Biomédicale; IPD, Institut Pasteur de Dakar; MRC/UVRI, Medical Research Council/Uganda Virus Research Institute; MRCG, Medical Research Council Unit—The Gambia; NICD, National Institute for Communicable Diseases; NMIMR, Noguchi Memorial

Institute for Medical Research. (**C**) Geographical representation of the total number of SARS-CoV-2 whole genomes produced over the course of the pandemic in each country, as well as the proportion of those sequences that were produced locally, regionally, or abroad. (**D**) Correlation of the proportion of COVID-19 positive cases that have been sequenced and the corresponding number of epidemiological weeks since the start of the pandemic that are represented with genomes for each African country. The color of each circle represents the number of cases and its size the number of genomes. (**E**) Comparison of sequencing turnaround times (lag times from sample collection to sequence submission) for the three strategies of sequencing in Africa, showing a significant difference in the means (****p < 0.0001). The box and whisker plot denotes the lower quartile, the median and upper quartiles (box), the minimum and maximum values (whiskers), and the outliers (black dots). (**F**) Pearson correlations of the total number of sequencing laboratories per country against key sequencing outputs.

genomes deposited in GISAID; however, there were still 16 countries with no reported local sequencing capacity (Fig. 2A) and undoubtedly many with limited capacity to meet demand during pandemic waves.

To tackle this, three centers of excellence and various regional sequencing hubs were established to maximize the resources available in a few countries to assist in genomic surveillance across the continent. This sequencing is done either as the sole source of viral genomes for those countries (e.g., Angola, South Sudan, and Namibia) or concurrently with local efforts to increase capacity during resurgences (Fig. 2B). Sequencing is further supplemented by a number of countries that use facilities outside of Africa. Ultimately, a mix of strategies from local sequencing, collaborative resource sharing among African countries, and sequencing with academic collaborators outside the continent helped close surveillance blind spots (Fig. 2C). Countries in sub-Saharan Africa, particularly in southern and East Africa, most benefited from the regional sequencing networks, whereas countries in West and North Africa often partnered with collaborators outside of Africa.

The success of pathogen genomic surveillance programs relies on how representative it is of the epidemic under investigation. For SARS-CoV-2, this is often measured in terms of the percentage of reported cases sequenced and the regularity of sampling. African countries were positioned across a range of different combinations of overall proportion and frequency of genomic sampling (Fig. 2D), Although the ultimate goal would be to optimize both of these parameters, a lower proportion of sampling can also be useful if the frequency of sampling is maintained at as high a level as possible. For instance, South Africa and Nigeria, which have both sequenced ~1% of cases overall, can be considered to have successful genomic surveillance programs based on the fact that sampling is representative over time and has enabled the timely detection of variants (Beta, Eta, Omicron).

Additionally, for genomic surveillance to be most useful for rapid public health response during a pandemic, sequencing would ideally be done in real time or in a framework as close as possible to that. We show a general trend of decreasing sequencing turnaround time in Africa (fig. S6), particularly from a mean of 182 days between October and December 2020 to a mean of 50 days over the same period a vear later, although this does come with several caveats. First, we measure sequencing turnaround time in the most accessible manner, which is by comparing the date of sampling of a specimen to the date its sequence was deposited in GISAID. Generally, the genomic data potentially informs the public health response more rapidly than reflected here, particularly

when it comes to local outbreak investigations or variant detection. This analysis is also confounded by various factors such as countryto-country variation in these trends (fig. S7), delays in data sharing, and potential retrospective sequencing, particularly by countries that joined sequencing efforts at later stages of the pandemic. The most critical caveat is the fact that sequencing from the most recently collected samples (e.g., over the past 6 months) may still be ongoing. The shortening duration between sampling and genomic data sharing is nevertheless a positive takeaway, given that these data also feed into continental and global genomic monitoring networks. Overall, the continental average delay from specimen collection to sequencing submission is 87 days, with 10 countries having an average turnaround time of less than 60 days and Botswana of less than 30 days (fig. S8).

Most importantly, in the context of optimizing genomic surveillance, we found that the route taken to sequencing affects the speed of data generation. Of the three frameworks we investigated, local sequencing has statistically faster sequencing turnaround times (median of 51 days), followed by sequencing within regional sequencing networks in Africa (median of 93 days) and finally outsourced sequencing to countries outside Africa (median of 113 days) (Fig. 2E). This finding strongly supports the investments in local genomic surveillance to generate timely and regular data for local and regional decision-making. Finally, we show that it is beneficial in several ways for countries to undertake genomic surveillance through several sequencing laboratories rather than by centralizing efforts. For instance, we estimate strong correlations between the numbers of sequencing laboratories per country and the total number of genomes produced by that country (Pearson correlation, 0.75), the total number of epiweeks for which sequencing data was produced (Pearson correlation, 0.81), and, importantly, sequencing turnaround time (Pearson correlation, -0.37) (Fig. 2F).

With the increase in sequencing capacity on the continent, a decrease in the time taken to detect new variants was observed. For example, the Beta variant was identified in December 2020 in South Africa (4), but sampling and molecular clock analyses suggest that the variant originated in September 2020. This 3-month lag in detection means that a new variant, like Beta, has ample time to spread over a large geographic region before its detection. However, by the end of 2021, the time to detect a new variant was substantially improved. Phylogenetic and molecular clock analyses suggest that the Omicron variant originated around 9 October 2021 (95% highest posterior density: 30 September to 20 October 2021), and the variant was described on 23 November 2021 (3). Thus, Omicron was detected within ~5 weeks from origin compared with the Beta variant (~16 weeks) and the Alpha variant, which was detected in the United Kingdom (~10 weeks). More importantly, the time from sequence deposition to the WHO declaring the new variant a VOC was substantially shortened to 72 hours for the Omicron variant.

To interpret insights from the described genomic surveillance in Africa, it is important to understand the context of epidemiological reporting and sampling strategies used for sequencing on the continent (table S3). Most countries provided daily reports of newly recorded cases, whereas a few provided weekly and monthly reports. For most countries, surveillance was mainly focused on the major cities, suggesting potential cryptic circulation in rural areas. We find that at the onset of the pandemic, surveillance was focused on identification of imported cases from incoming travelers or local residents returning from various countries. As community transmissions began to emerge, the focus shifted toward regular surveillance and outbreak investigations. Together, these three strategies account for the vast majority of samples generated on the continent and analyzed here. As the pandemic progressed and vaccines were made available, some countries on the continent began to explore other sampling strategies such as reinfections, environmental samples such as wastewater samples, and vaccine breakthrough cases to gain new insights into the evolutionary dynamics of SARS-CoV-2. The utility of sequencing for viral evolution tracking and VOC detection in the way described above is obviously also dependent on sampling proportions, especially within sampling for regular surveillance.

The speed of SARS-CoV-2 evolution has complicated sequencing efforts. Common methods of RNA sequencing include reverse transcription followed by double-stranded DNA amplification using sequence-specific primer sets (39). Ongoing SARS-CoV-2 evolution has necessitated the continual evaluation and updating of these primer sets to ensure their sustained utility during genomic surveillance efforts. Here, we examined the current set of genomes to determine aspects of the sequencing process that might be improved in the future. Many of the primer sets that were used were designed using viral sequences from the start of the pandemic and may require updating to keep pace with evolution. Indeed, the ARTIC primer sets are now in version 4.1 (40). The Entebbe primer set was designed mid-2020, well into the first year of the epidemic, and used an algorithm and design that accommodates evolution (41).

The effects of viral evolution on sequencing patterns can be seen with low median unspecified nucleotide (N) values (a consequence of primer dropout or low coverage at that site) that were observed for the first 12 months of the epidemic, with an increase from October 2020

(Fig. 3A). Additional challenges appear (as indicated by increasing median N values) as the virus further evolved into the Delta and Omicron lineages from January 2021 onward (Fig. 3A). By examining the role of sequencing technology, it appears that the two major technologies used (Illumina and ONT) have similar gap profiles (as measured by mean N count per genome), whereas Ion Torrent, MGI, and Sanger show a reduced mean N count per genome (Fig. 3B). Likely factors for this pattern are the primers used in sequencing, with primer choice playing a key role in the quantity of gaps (Fig. 3C). The mean N count per genome varied with viral lineage (Fig. 3D). There was a modest difference in mean N count per genome across the lineages. Lineages that returned no classification with Pangolin ("none") showed the highest mean N count, suggesting that high mean N count per genome was probably the basis for failed classification. The more recent lineages, Delta (e.g., AY.39, AY.75) and Omicron (BA.1.1, BA.2), also showed higher mean N count per genome, consistent with virus evolution impairing primer function. This pattern is further explored in fig. S9, where the position of gaps shows an enrichment in the genome regions after position 19,000, with frequent gaps disrupting the spike coding region.

Phylogenetic insights into the rise and spread of VOCs in Africa

During the first wave of infections in 2020 in Africa, as was the case globally, most corresponding genomes were classified as PANGO B.1 (n =2456) or B.1.1 viruses (n = 1329). Toward the end of 2020, more-distinct viral lineages started to appear. Of these, the most important ones that affected the African continent are B.1.525 (n = 797), B.1.1.318 (n = 398) (42), B.1.1.418 (n = 398)395), A.23.1 (n = 358) (15, 29, 31, 33), C.1 (n = 446) (29), C.1.2 (n = 300) (31), C.36 (n = 305) (30, 43), B.1.1.54 (n = 287) (15, 29, 31, 33), B.1.416 (n = 272), B.1.177 (n = 203), B.1.620 (n =138), and B.1.160 (n = 61) (32) (fig. S10, A and B). Our discrete state phylogeographic inference from phylogenetic reconstruction of non-VOC African sequences and an equal number of external references revealed that African countries were primarily seeded by multiple introductions of viral lineages from abroad (mainly Europe) at the beginning of the pandemic. The observed pattern of non-VOC viral lineage movement then consistently shifted toward more intercontinental exchanges (fig. S10C). Mapping out the spatial routes of dissemination shows that various countries in all subregions of the continent acted as sources of these viral lineages at one point or another (fig. S10D). Although uneven testing rates and proportions of samples sequenced on the continent may have influenced these inferences (discussed later), the results presented here are in line with the fact that these most predominant

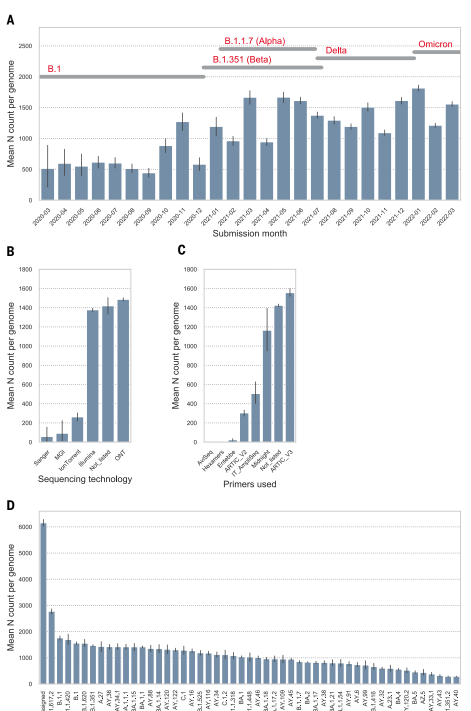


Fig. 3. Genome gap analysis. (A) The mean N count per genome by month of submission to GISAID. The time periods corresponding to the detection of important SARS-CoV-2 lineages are indicated at the top of the figure. (B) Illustration of the mean N count per genome stratified by sequencing technology. (C) The mean N count per genome stratified by the sequencing primers sets used. (D) Mean N count per genome by lineage. The mean N data were stratified by SARS-CoV-2 lineages to investigate the lineage-specific frequency of genome gaps, an indirect measure of primer mismatch. All lineages that were present at least 100 times in the genome data are presented. For (A) to (D), error bars indicate 95% confidence intervals.

SARS-CoV-2 lineage

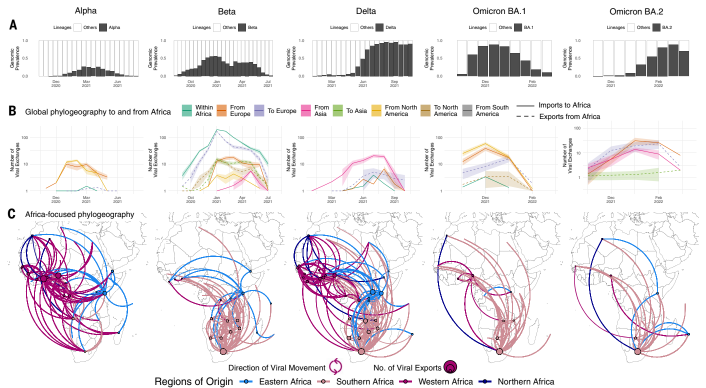


Fig. 4. Inferred viral dissemination patterns of VOCs within Africa. (A) Genomic prevalence of VOCs Alpha, Beta, Delta, and Omicron in Africa over time. (B) Inferred viral exchange patterns to, from, and within the continent of Africa for the four VOCs (Omicron as BA.1 and BA.2) based on case-sensitive phylogeographic inference. Introductions and viral transitions within Africa are shown as solid lines, and exports from Africa are shown as dotted lines; the lines are colored by continent.

The shaded areas around the lines represent the uncertainty of this analysis from 10 replicates (±SD). (**C**) Dissemination patterns of the VOCs within Africa obtained from inferred ancestral-state reconstructions performed on Africa-enriched datasets, annotated and colored by region in Africa. The countries of origin of viral exchange routes are also shown with dots, and the curves go from country of origin to destination country in a counterclockwise direction.

non-VOC lineages in Africa, except B.1.177, emerged and circulated widely in different subregions (Fig. 1).

Similar to the pandemic globally, VOCs became increasingly important in Africa toward the end of 2020. The Alpha, Beta, Delta, and Omicron variants demonstrate many similarities as well as differences in the way that they spread on the continent. For all these VOCs, we observe large regional monophyletic transmission clusters in each of their phylogenetic reconstructions in Africa (fig. S11). This suggests an important extent of continental dissemination within Africa. Alpha and Beta were epidemiologically important in distinct regions of the continent, with Alpha primarily circulating in West Africa, North Africa, and most of Central Africa: Beta circulating in southern and most of East Africa; and both only substantially cocirculating in a few countries such as Angola, Kenva, Comoros, Burundi, and Ghana (Fig. 1 and fig. S12). However, we may not have enough resolution in the geospatial data to know whether and to what extent they were truly cocirculating throughout these countries or whether there were regional outbreaks of Alpha and Beta within these countries. In Kenya, for example, Beta was detected more frequently in coastal regions and Alpha more frequently inland (26, 44). By contrast, the Delta and Omicron variants sequentially dominated most infections on the entire continent shortly after their emergence (Fig. 4A and fig. S12).

The Alpha variant was first identified in December 2020 in the United Kingdom and has since spread globally. In Africa, Alpha was detected in 43 countries, with evidence of community transmission based on phylogenetic clustering in many countries, including Ghana, Nigeria, Kenya, Gabon, and Angola (fig. S11). Discrete state maximum likelihood reconstruction from a globally case-sensitive genomic subsampling inferred at least 80 introductions [95% confidence interval (CI): 78 to 82] into Africa, with the bulk of imports attributed to the United States (>47%) and the United Kingdom (>25%) (Fig. 4B). Only 1% of imports into any particular African country were attributed to another African nation. Phylogeographic reconstruction enriched in African sequences revealed that of those, >85% of the intercontinental Alpha exchanges in Africa originated from West African countries (Fig. 4C). This occurred in spite of initial importations of the Alpha variant from Europe into all regions of the continent (fig. S13B) but is in line with Alpha having dominated circulation mostly in West Africa (fig. S12). In countries where Alpha was introduced but did not grow and cause an expansion of cases, this can be explained by competition with the already established Beta variant, which simultaneously circulated. The characteristics of multiple introductions of Alpha into Africa and between African countries is similar to the spread of Alpha that has been documented in the United Kingdom, Scotland, and Ireland (45–47).

The second VOC, Beta, was identified in December 2020 in South Africa (4). However, sampling and molecular clock analyses suggest that the variant originated around September 2020 (fig. S11). At the end of 2020 and beginning of 2021, Beta was driving a second wave of infection in South Africa and quickly spread to other countries within the region. The concurrent introductions and spread of Alpha and other variants (Eta, A.23.1) in other regions of the continent may have reduced the Beta variant's initial growth, limiting its spread largely to southern Africa and, to a lesser extent, the East Africa region. Beta spread to at least 114 countries globally, including 37 countries and territories in Africa. For this

variant, viral circulation and geographical exchanges occurred predominantly within the continent. Indeed, phylogeographic reconstruction from a globally case-sensitive sampling revealed that of the 810 (95% CI: 803 to 818) inferred introductions of the Beta variant into African countries, only 110 (95% CI: 105 to 115; 13%) were attributed to sources outside the continent (fig. S13C), whereas more than half of the introductions were attributed to South Africa (63%) (Fig. 4C). This is in line with expectations because the variant originated in South Africa. Beyond southern Africa, most of the introductions back into the continent were attributed to France and other European Union countries into the French overseas territories, Mayotte and Reunion, and other Francophone African countries. Africafocused phylogeographic analysis revealed a similar spatial pattern that showed southern countries as substantial sources of the variant, followed in small numbers by countries in East Africa (Fig. 4C).

The fourth VOC observed was Delta (13), which rose to prominence in April 2021 in India, where it fueled an explosive second wave. Since its emergence, Delta has been detected in >170 countries, including 37 African countries and territories (fig. S11). Our global case-sensitive subsampled analysis infers at least 100 (95% CI: 93 to 106) introductions of the Delta variant into Africa, with the bulk attributed to India (~72%), mainland Europe (~8%), the United Kingdom (\sim 5%), and the United States (\sim 2.5%). Viral introductions of Delta also occurred from one African country to others in 7% of inferred introductions. From our Africa-focused phylogeographic inferences, we infer that unlike Alpha and Beta, viral dissemination of Delta within Africa was not restricted to or dominated by any particular region but rather spread across the entire continent (Fig. 4C). After introductions from Asia in the middle of 2021, Delta rapidly replaced the other circulating variants (Fig. 4A). For example, in southern African countries, the Delta variant rapidly displaced Beta and, by June 2021, was circulating at very high (>90%) frequencies (48).

The latest VOC, Omicron, was identified and characterized in November 2021 in southern Africa (3). At the time of writing, the variant had been detected and caused waves of infections in >160 countries, including 39 African countries and two overseas territories (fig. S11). Because of the genetic distance between them and their sequential (rather than simultaneous) epidemic expansion globally, phylogenies were reconstructed separately for Omicron BA.1 and BA.2. Our discrete ancestral-state reconstruction from a global case-sensitive sampling for Omicron BA.1 infers at least 55 (95% CI: 47 to 62) viral exports of BA.1 out of various African countries, of which 31 (95% CI: 25 to 36) were toward Europe and 8 (95% CI:

6 to 10) were toward North America (Fig. 4B). After explosive expansion of Omicron around the world, we inferred even more reintroductions of the variant back into Africa, at least 69 (95% CI: 60 to 78) from Europe and 102 (95% CI: 92 to 112) from North America (Fig. 4B). From our Africa-focused phylogeographic reconstructions, we determine that, as with Delta, routes of dissemination of this variant involved all regions of the continent spatially (Fig. 4C). Yet ~75% of all BA.1 viral movement volume in Africa happened between southern African countries, likely because of rapid epidemic expansion in the region soon after its detection (3). Omicron BA.2's reach in Africa was limited at the time of writing, with only 3260 sequences from 19 countries attributed to BA.2 on GISAID (date of access: 31 March 2022) (15% of all Omicron sequences from Africa). Our discrete ancestral-state reconstruction from a global case-sensitive sampling for Omicron BA.2 infers at least 68 (95% CI: 53 to 84) viral exports out of African countries, of which most were toward Europe (~88%) (Fig. 4B). We also infer at least 99 (95% CI: 87 to 109) separate introduction or reintroduction events of BA.2 back into African countries, of which ~65% are from Europe and ~30% from Asia, primarily from India (Fig. 4B). This is consistent with India having experienced one of the earliest large BA.2 waves globally. In the context of global incidence of BA.2, this case-sensitive phylogeographic analysis revealed that only 0.01% of viral movements of this lineage globally happened from one African country to another. Our Africa-focused analysis inferred a similar pattern of BA.2 spatial diffusion within African to that of BA.1 (Fig. 4C). However, given that this accounted for such a small percentage of global BA.2 movements, BA.2 diffusion from one African country to another is unlikely to have had a substantial impact on epidemiological expansion, compared with introductions from Asia, Europe, or North America.

Globally, dissemination of the SARS-CoV-2 virus throughout the pandemic was intricately linked with human mobility patterns (49-53). To determine the validity of the VOC movement patterns that we infer into and within the Africa continent in this study, we compared viral import and export events to and from South Africa with travel to the country. In December 2020, the United Kingdom accounted for the fifth-highest number of passengers entering South Africa, whereas other countries with the top-nine sources of travelers were all neighboring countries in southern Africa (fig. S14A). Considering that incidence of the Alpha variant was not meaningful in the region, this supports our inference of the United Kingdom contributing 60% of Alpha introductions to South Africa (fig. S15A). In March 2021, the United States, Germany, the United Kingdom, and India were among the top-12 sources of travelers to South Africa after eight African countries (fig. S14B). During this time of Delta dissemination globally, we infer that ~90% of introductions of Delta into South Africa originated in the United Kingdom, the United States, and India (fig. S15B). At the end of 2021, most introductions or reintroductions of Omicron to the country came from the United Kingdom, the United States, or Botswana, corresponding to locations of both high Omicron incidence at the time and high numbers of passengers to South Africa (figs. S14C and S15C). These travel patterns also fit the findings that ~89, ~70, and ~75% of Beta, Delta, and Omicron exports, respectively, from South Africa to other African countries were directed to locations in southern Africa (figs. S14, D and E, and S15, D and E).

Discussion, limitations, and conclusions

By April 2020, a total of 20 African countries were able to sequence the virus within their own borders. This was largely made possible by other preexisting sequencing efforts on the continent that were focused on other human pathogens (e.g., HIV, tuberculosis, Ebola, and H1N1). However, these efforts were quickly limited by global supply chain issues, and, in many countries, sequencing efforts substantially slowed down or stopped toward the end of 2020. To facilitate more sequencing on the continent over the course of the past year (April 2021 to March 2022), the Africa CDC and partners invested heavily to support genomic surveillance on the continent. This included the transfer of 24 new sequencing platforms (including MinIon, GridIon, MiSeq, and NextSeq), the distribution of reagents and flow cells to support the sequencing of 100,000 positive samples, the training of >230 students and technicians in wet laboratory and bioinformatic techniques, and additional grants to support 10 regional sequencing hubs. This investment has started bearing fruit and should be intensified as the virus continues to evolve, requiring the adaptation of methodologies locally on the continent to keep pace with the emergence of variants. The continued development of sequencing protocols in Africa is of crucial importance (41, 54, 55) given the number of variants and lineages that emerged in, and were introduced to, the continent. In North Africa, the SARS-CoV-2 pandemic was caused by waves of infections that were similar to those seen in Europe (first wave attributed to B.1 descendants, second wave to Alpha, third wave to Delta, and fourth wave to Omicron): in southern Africa, the pattern was similar but with a Beta wave instead of an Alpha one. In East Africa, the pandemic was more complex, involving both Alpha and Beta as well as its own lineage A.23.1 before the arrival of Delta and Omicron. Central Africa experienced epidemic patterns that sometimes mirrored those of East Africa and other times those of southern Africa. In West Africa, Eta made a considerable contribution to both a second wave (together with Alpha) and a third wave (together with Delta). The factors that resulted in these regional differences are not clear but could be due to differences in human mobility, founder effects, competition between lineages, or the immunity induced by earlier waves in a region.

Public health benefits of such broadly inclusive genomic surveillance are manifold. The most prominent insight from this expanded genomic surveillance in Africa has been an early warning capacity for the world after the detection of new lineages and variants, most recently relevant in the detection of Omicron BA.1, BA.2, BA.3, BA.4, and BA.5 subvariants (3, 4, 34). Furthermore, the reporting of local SARS-CoV-2 sequences made the epidemic more immediate to the Ministries of Health from the reporting African countries. It became clear early on that the viral evolution is global and that the transmission of the virus is extremely rapid, which guided mitigation strategies. The generation and availability of local sequences also validated local diagnostics and allowed investigators to determine whether nucleic acid-based diagnostics that were in use could still detect local variants. The detection of SARS-CoV-2 in returning travelers and truck drivers indicated routes that the virus might be using to enter a country and guided early efforts to slow virus entry and gain time to establish vaccination plans. Later, the difficulty of stopping the virus at borders combined with data showing that the variants were already in community circulation allowed public health officials to focus efforts and limited resources on vaccination rather than on border controls. The detection and reporting of the more-recent lineages with enhanced transmission (i.e., Omicron) and the ability to bypass existing immunity is important information and an early alert to public health officials globally that the epidemic is still proceeding. As the pandemic progresses in an evolving global context, we provide evidence that with each new variant, transmission dynamics are changing and the use of sequencing with phylogenetics could potentially alter decisions of public health measures. For example, the demonstrated shift away from regional dynamics of Alpha and Beta toward more global patterns with Delta and Omicron can provide insights to public health officials as they anticipate epidemic developments locally. With Omicron, it became clear that although the variant expanded first in Africa, the continent ultimately had a minimal role in global dissemination and that continental expansion beyond southern Africa was most influenced by external introductions, in contrast to the Beta variant. All of these public health benefits to sequencing SARS-CoV-2 are primarily amplified, as we show in this study, if the sequencing can be conducted locally within a country, which strongly supports the continued investment into pathogen sequencing on the continent.

Despite the recent successful expansion of genomics surveillance in Africa, additional work is necessary. Even with investments from the Africa CDC-Africa Pathogen Genomics Initiative and other investments, there are still 16 countries with no sequencing capacity within their own borders. The only option for these countries is to send samples to continental sequencing hubs or to centers outside of the continent, which increases turnaround times and limits the utility of genomic surveillance for public health decision-making. Secondly, not all countries are willing to share data openly in a timely fashion for fear of being subject to travel bans or restrictions that could bring substantial economic harm. Such hesitancy has obvious potential ramifications for the future of genomic surveillance on the continent. Furthermore, with the expansion of sequencing on the continent, there is a growing need for more bioinformatics support and knowledge to allow investigators to analyze and report their data in a reasonable time frame that makes it useful for a public health response. It is also clear that the SARS-CoV-2 sequencing primers are not a static development and may require updating as the virus evolves. A number of research groups have been addressing the SARS-CoV-2 sequencing primer questions. Issues of gaps in the genomes due to missing amplicons have been discussed (56, 57). The ARTIC primer set has gone through a number of revisions to accommodate virus evolution (39, 40). Additional longer amplicon methods have been published (58-60), including methods to use a subset of ARTIC primers (61).

The patterns we describe here are of course limited to reported cases and apply to both the phylogeographic as well as the epidemiology inferences. As such, the results need to be interpreted with these limitations in mind. Our primary phylogeographic inference relied on a sampling strategy that considered all highquality African sequences and an equal number of external references. Though this strategy has the advantage of placing all African sequences in a phylogenetic context, it introduces a bias when applied to discrete ancestral-state reconstruction because more internal nodes are inferred to be from Africa. To address this, we performed an even sampling of global cases, based on reported case counts through time, to compare against our oversampled inference. The even-sampling approach has the benefit that the discrete ancestral-state reconstruction is not biased by uneven sampling. After comparing the two, there are obvious differences, most notably that the number of inferred introductions into Africa is proportional to sampling proportions (fig. S16) because we no longer consider all African sequences but rather just a small subset against a global sample. However, inferences from the two approaches correspond well with one another. For example, considering Alpha, we still observed that the vast majority of introductions into Africa originated from Western Europe. Patterns of dissemination within Africa are more robustly comparable between the two, for instance, that countries in West Africa were the biggest source of Alpha within the continent. High concordance between the two inference methods was also observed for other VOCs for dispersal routes within Africa, which gives us confidence in the inferred patterns we observe here. Although we represent an inference based on oversampling and case-sensitive sampling, it is, at present, not possible to explore how undersampling affects the phylogeographic reconstruction because of uneven testing rates. Additionally, the robustness of the phylogeographic inference can also be affected by the underlying methodology that is used. Broad consensus would favor the use of Bayesian methods for phylogeographic reconstruction, which is often considered to be the "gold standard" in the field. The main drawbacks of Bayesian methods are that they can only be applied to a relatively small number of sequences at a time (<1000) and they are extremely computationally and time intensive. Given the explosion of sequence data over the past 2 years, the scientific community will have to adapt or put forth new analytical methods to fully capitalize on the global sequencing efforts for SARS-CoV-2.

Despite our best attempts to consider and minimize genomic sampling bias, the accuracy of the resulting phylogenetic inferences is limited by the available epidemiological and genomic data, leading to unaccounted biases in the estimates of viral movements. This includes limited testing and subsequent sequencing in many African countries. Although the percentage of reported cases sequenced in African countries (0.01 to 10%, mean = 1.27%) is not far from global figures (0.01 to 16%, mean = 1.31%), testing rates and infection-to-detection ratios in Africa were some of the lowest globally (38, 62). Together with estimates of excess mortality being as much as 20-fold greater than the reported numbers in African countries (63), these are strong indications of undetected and underreported epidemic sizes in Africa, leading to undersampling of genomic data (62) and thus underestimates of viral exchange inferences in our study. Some countries with no publicly available SARS-CoV-2 sequences are, by definition, completely missing in our inference. This in turn means that inferred routes of viral transmission within Africa could be missing important intermediate locations, although this is potentially true around the world. Nevertheless, we believe that the viral movement inferences that we discuss in this

study provide a likely qualitative description of the patterns of SARS-CoV-2 migration into, out of, and within Africa.

Finally, we should also mention uneven sequencing and reporting standards across the different laboratories on the continent-and globally, for that matter. Different groups use different measures for what constitutes a highquality sequence (e.g., 70 versus 80% sequence coverage) or use different sequencing depth coverage. This lack of global standardization complicates the direct comparison of sequences that may have been submitted to GISIAD using different criteria, further biasing any inference. Given the sheer size of SARS-CoV-2 sequencing, with ~10 million whole-genome sequences shared on the GISAID database (date of access: 31 March 2022), there is an urgent need for global standards with regard to sequence quality and associated metadata.

Africa needs to continue expanding genomic sequencing technologies on the continent in conjunction with diagnostic capabilities. This holds true not just for SARS-CoV-2 but also for other emerging or reemerging pathogens on the continent. For example, in February 2022, the WHO announced the reemergence of wild polio in Africa, and sporadic influenza H1N1, measles, and Ebola outbreaks continue to occur on the continent. The Africa CDC has estimated that more than 100 pathogen outbreaks are reported across the continent every year. Beyond the current pandemic, continued investment in diagnostic and sequencing capacity for these pathogens could serve the public health of the continent well into the 21st century.

Methods and methods Ethics statement

This project relied on sequence data and associated metadata that are publicly shared by the GISAID data repository and adhere to the terms and conditions laid out by GISAID (16). The African samples processed in this study were obtained anonymously from material exceeding the routine diagnosis of SARS-CoV-2 in African public and private health laboratories. Individual institutional review board references or material transfer agreements (MTAs) for countries are as follows: Angola (MTA - CON8260); Botswana-genomic surveillance in Botswana was approved by the Health Research and Development Committee (protocol HPDME 13/18/1); Egypt-surveillance in Egypt was approved by the Research Ethics Committee of the National Research Centre (Egypt) (protocol number 14 155, dated 22 March 2020); Kenya-samples were collected under the Ministry of Health protocols as part of the national COVID-19 public health response, and the whole-genome sequencing study protocol was reviewed and approved by the Scientific and Ethics Review Committee (SERU) at Kenya Medical Research Institute (KEMRI), Nairobi,

Kenya (SERU protocol #4035); Nigeria (NHREC/ 01/01/2007), Mali-study of the sequence of SARS-CoV-2 isolates in Mali, Letter of Ethical Committee (N0-2020 /201/CE/FMPOS/FAPH of 09/17/ 2020); Mozambique (MTA - CON7800); Malawi (MTA - CON8265); South Africa-the use of South African samples for sequencing and genomic surveillance was approved by University of KwaZulu-Natal Biomedical Research Ethics Committee (ref. BREC/00001510/2020), the University of the Witwatersrand Human Research Ethics Committee (HREC) (ref. M180832), Stellenbosch University HREC (ref. N20/04/008_ COVID-19), the University of the Free State Research Ethics Committee (ref. UFS-HSD2020/ 1860/2710), and the University of Cape Town HREC (ref. 383/2020); Tunisia-for sequences derived from sampling in Tunisia, all patients provided their informed consent to use their samples for sequencing of the viral genomes, and the ethical agreement was provided to the research project ADAGE (PRFCOVID19GP2) by the Committee of Protection of Persons (Tunisian Ministry of Health) under the reference CPP SUD N 0265/2020; Uganda-the use of samples and sequences from Uganda was approved by the Uganda Virus Research Institute, Research and Ethics Committee UVRI-REC Federalwide Assurance (FWA) no. 00001354, study reference GC/127/20/04/771, and by the Uganda National Council for Science and Technology, reference number HS936ES; and Zimbabwe (MTA - CON8271).

Epidemiological and genomic data dynamics

We analyzed trends in daily numbers of cases of SARS-CoV-2 in Africa up to 31 March 2022 from publicly released data provided by the Our World in Data repository for the continent of Africa (https://github.com/owid/covid-19-data/ tree/master/public/data) as a whole and for individual countries (2). To provide a comparable view of epidemiological dynamics over time in various countries, the variable under primary consideration for Fig. 1 was "new cases per million (smoothed)." To calculate the genomic sampling proportion and frequency for each country for Fig. 2, the total number of recorded cases as of 31 March 2022 was considered, as well as the total length of time for which each country had recorded cases of SARS-CoV-2.

Genomic metadata was downloaded for all African entries on GISAID for the same time period (date of access: 31 March 2022). From this, information extracted from all entries for this study included the date of sampling, country of sampling, viral lineage and clade, originating laboratory, sequencing laboratory, and date of submission to the GISAID database. The geographical locations of the originating and sequencing laboratories were manually curated. Sequences originating and sequenced in the same country were defined as locally sequenced, irrespective of specific laboratory or finer location. Sequences originating in one African country and sequenced in another were defined as sequenced within regional sequencing networks. Sequences sequenced in a location not within Africa were labeled as sequenced outside Africa. Sequencing turnaround time was defined as the number of days that had elapsed from specimen collection to sequence submission to GISAID. Sequencing technology information for all African entries was also downloaded from GISAID on 31 March 2022.

Primer choice and sequencing outcomes

All SARS-CoV-2 genomes from African countries were retrieved from GISAID (16) for submission dates from 1 December 2019 to 31 March 2022, yielding 100,470 entries. Associated metadata for the entries were also retrieved, including collection date, submission date, country, viral strain, and sequencing technology. Data on the primers used for the sequencing were requested from investigators and yielded primer data for 13,973 of the entries (~13%). The total N (bases with low sequence depth) per genome were counted, the results of which were then used for genome quality analysis and visualization. Gap locations in the genomes were mapped and visualized with respect to the original Wuhan strain (64).

Phylogenetic investigation

All African sequences on the GISAID sequence database (16) were downloaded on 31 March 2022 (n = 100,470). Of these, Alpha accounted for 3851 sequences. Beta accounted for 14.548 sequences, Delta accounted for 35,027 sequences, Omicron accounted for 21,708 sequences, and 25,336 sequences were classified as non-VOCs. Before any phylogenetic inference, we performed some quality assessment on the sequences to exclude incomplete or problematic sequences as well as sequences lacking complete metadata. Briefly, all African sequences were passed through the NextClade analysis pipeline (65) to identify and exclude (i) sequences missing >10% of the SARS-CoV-2 genome, (ii) sequences that deviate by >70 nucleotides from the Wuhan reference strain, (iii) sequences with >10 ambiguous bases, (iv) clustered mutations, and (v) sequences flagged with private mutations by NextClade. Additionally, Omicron variants were screened for traces of viral recombination with RDP5.23 (66) using default settings and a p value of ≤ 0.05 as evidence of recombination. A large number of sequences were removed (n = 57,421), with incomplete sequences (<90% genome coverage) being the biggest contributor. This produced a final African dataset of 43,049 high-quality African sequences. Because of the sheer size of the dataset, we opted to perform independent phylogenetic inferences on the main VOCs (Alpha, Beta, Delta, and Omicron BA.1 and BA.2) that have spread on the African continent, as well as a separate inference for all non-VOC SARS-CoV-2 sequences.

To evaluate the spread of the virus on the African continent, we aligned the African datasets against a large number of globally representative sequences from around the world. Because of the oversampling of some variants or lineages, we performed a random down sampling while retaining the oldest two known variants from each country. Reference sequences were respectively aligned with their African counterparts independently with NextAlign (65). Each of the alignments was then used to infer maximum likelihood (ML) tree topologies in FastTree v 2.0 (67) using the general time reversible model of nucleotide substitution and a total of 100 bootstrap replicates (68). The resulting ML tree topologies were first inspected in TempEst (69) to identify any sequences that deviate more than 0.0001 from the residual mean. After the removal of potential outliers in R with the ape package (70), the resulting ML trees were then transformed into timecalibrated phylogenies in TreeTime (71) by applying a rate of 8×10^{-4} substitutions per site per year (72) to transform the branches into units of calendar time. Time-calibrated trees were then visualized, along with associated metadata, in R using ggtree (73) and other packages.

We performed a basic viral dispersal analysis for each of the VOCs (excluding Gamma) as well as for the non-VOC dataset. Briefly, a migration model was fitted to each of the timecalibrated tree topologies in TreeTime, mapping the country location of sampled sequences to the external tips of the trees. The mugration model of TreeTime also infers the most likely location for internal nodes in the trees. Using a custom python script, we could then count the number of state changes by iterating over each phylogeny from the root to the external tips. We count state changes when an internal node transitions from one country to a different country in the resulting child node or tip(s). The timing of transition events is then recorded, which serves as the estimated import or export event. To infer some confidence around these estimates, we performed 10 replicates for each of the datasets by random selection from the 100 bootstrap trees. Because of the high uncertainty in the inferred locations for deep internal nodes in the trees, we truncated state changes to the earliest date of sampling in each dataset. All data analytics were performed using custom python and R scripts, and the results were visualized using the ggplot libraries (74). Such phylogeographic methods are always subject to uneven sampling through time (i.e., over the course of the pandemic) and through space (by sampling location). To address this, we have performed a case-sensitive analysis to investigate the effects of oversampling African locations on the inferred number of viral introductions. Furthermore, in a previous analysis (15), we performed a sensitivity analysis to address some of these issues and found no substantial variations in estimates.

Case-sensitive phylogeographic inference

To address the potential oversampling of African sequences relative to global reference in the above-mentioned analyses, we performed another phylogeographic inference on subsamples based on global case counts to try to eliminate oversampling bias in our inference. To this end, we considered all high-quality sequences for each of the VOCs (Alpha, Beta, Delta, and Omicron BA.1 and BA.2) globally over the same sampling period (until 31 March 2022). We used subsampler (https://github.com/andersonbrito/ subsampler) to generate subsamples for each variant based on globally reported cases. In short, subsampler uses a case-count matrix of daily cases, along with the fasta sequences and GISAID associated metadata, to sample a user-defined number of sequences. For each VOC and for BA.1 and BA.2, we performed 10 samplings using different number seeds to sample datasets of ~20,000. Once again, sampled sequences were screened for viral recombination as described above and sequences with signs of recombination were removed. Subsampler has the added advantage that it disregards poor quality sequences (e.g., <90% coverage) and sequences with missing metadata (e.g., exact date of sampling). Each dataset was then subjected to the same analytical pipeline as mentioned above to infer the viral transitions between Africa and the rest of

Regional and country-specific NextStrain builds

To investigate more-granular changes in lineage dynamics within a specific country or region in Africa, we used the NextStrain pipeline (https://github.com/nextstrain/ncov) to generate the regional and country-specific builds for African countries (75). First, all sequence data and metadata were retrieved from the GISAID sequence database and filtered for Africa based on the "region" tab for inclusion in regional and country-specific African builds. For country-specific builds, ~4000 sequences from a given country were randomly selected and analyzed against ~1000 randomly selected sequences from the Africa "nextregions" records that do not match the focal country of interest. For regional (e.g., West Africa) builds, ~4000 sequences from the focal region were selected at random and analyzed against ~1000 randomly selected sequences from the Africa "nextregions" records that do not match the focal region of interest. The methodological pipeline for NextStrain is well documented and performs all analyses within one workflow, including filtering of sequences, alignment, tree inference, molecular clock, and ancestralstate reconstruction. For more information, please visit https://docs.nextstrain.org/en/latest/index.html.

All regional and country-specific builds are regularly updated to keep track of the evolving pandemic on the continent. All builds are publicly available under the links provided in tables S1 and S2 as well as on the NextStrain web page (https://nextstrain.org/sars-cov-2/#datasets).

REFERENCES AND NOTES

- Q. Li et al., Early transmission dynamics in Wuhan, China, of novel coronavirus-infected pneumonia. N. Engl. J. Med. 382, 1199–1207 (2020). doi: 10.1056/NEJMoa2001316; pmid: 31995857
- J. Hasell et al., A cross-country database of COVID-19 testing. Sci. Data 7, 345 (2020). doi: 10.1038/s41597-020-00688-8; pmid: 33033256
- R. Viana et al., Rapid epidemic expansion of the SARS-CoV-2 Omicron variant in southern Africa. Nature 603, 679–686 (2022). doi: 10.1038/s41586-022-04411-y; pmid: 35042229
- H. Tegally et al., Detection of a SARS-CoV-2 variant of concern in South Africa. Nature 592, 438–443 (2021). doi: 10.1038/ s41586-021-03402-9; pmid: 33690265
- D. P. Martin et al., The emergence and ongoing convergent evolution of the SARS-CoV-2 N501Y lineages. Cell 184, 5189–5200.e7 (2021). doi: 10.1016/j.cell.2021.09.003; pmid: 34537136
- F. Campbell et al., Increased transmissibility and global spread of SARS-CoV-2 variants of concern as at June 2021. Euro Surveill. 26, (2021). doi: 10.2807/1560-7917.ES.2021.26.24.2100509; pmid: 34142653
- B. Korber et al., Tracking changes in SARS-CoV-2 spike: Evidence that D614G increases infectivity of the COVID-19 virus. Cell 182, 812–827.e19 (2020). doi: 10.1016/ j.cell.2020.06.043; pmid: 32697968
- E. Hacisuleyman et al., Vaccine breakthrough infections with SARS-CoV-2 variants. N. Engl. J. Med. 384, 2212–2218 (2021). doi: 10.1056/NEJMoa2105000; pmid: 33882219
- D. Planas et al., Reduced sensitivity of SARS-CoV-2 variant Delta to antibody neutralization. Nature 596, 276–280 (2021). doi: 10.1038/s41586-021-03777-9; pmid: 34237773
- S. Yue et al., Sensitivity of SARS-CoV-2 variants to neutralization by convalescent sera and a VH3-30 monoclonal antibody. Front. Immunol. 12, 751584 (2021). doi: 10.3389/ fimmu.2021.751584; pmid: 34630430
- S. Cele et al., Escape of SARS-CoV-2 501Y.V2 from neutralization by convalescent plasma. Nature 593, 142–146 (2021). doi: 10.1038/s41586-021-03471-w; pmid: 33780970
- B. Meng et al., Recurrent emergence of SARS-CoV-2 spike deletion H69/V70 and its role in the Alpha variant B.1.1.7.
 Cell Rep. 35, 109292 (2021). doi: 10.1016/j.celrep.2021.109292; pmid: 34166617
- P. Mlcochova et al., SARS-CoV-2 B.1.617.2 Delta variant replication and immune evasion. Nature 599, 114–119 (2021) doi: 10.1038/s41586-021-03944-y; pmid: 34488225
- N. R. Faria et al., Genomics and epidemiology of the P.1 SARS-CoV-2 lineage in Manaus, Brazil. Science 372, 815–821 (2021). doi: 10.1126/science.abh2644; pmid: 33853970
- E. Wilkinson et al., A year of genomic surveillance reveals how the SARS-CoV-2 pandemic unfolded in Africa. Science 374, 423–431 (2021). doi: 10.1126/science.abj4336; pmid: 34672751
- Y. Shu, J. McCauley, GISAID: Global initiative on sharing all influenza data - from vision to reality. *Euro Surveill.* 22, 30494 (2017). doi: 10.2807/1560-7917.ES.2017.22.13.30494; pmid: 28382917
- C. Kuiken, B. Korber, R. W. Shafer, HIV sequence databases. *AIDS Rev.* 5, 52–61 (2003). pmid: 12875108
- D. L. Bugembe et al., Main routes of entry and genomic diversity of SARS-CoV-2, Uganda. Emerg. Infect. Dis. 26, 2411–2415 (2020). doi: 10.3201/eid2610.202575; pmid: 32614767
- T. Mashe et al., Genomic epidemiology and the role of international and regional travel in the SARS-CoV-2 epidemic in Zimbabwe: A retrospective study of routinely collected surveillance data. Lancet Glob. Health 9, e1658–e1666 (2021). doi: 10.1016/ S2214-109X(21)00434-4; pmid: 34695371
- A. Chouikha et al., Molecular epidemiology of SARS-CoV-2 in Tunisia (North Africa) through several successive waves of COVID-19. Viruses 14, 624 (2022). doi: 10.3390/v14030624; pmid: 35337031

- F. Ntoumi et al., Genomic surveillance of SARS-CoV-2 in the Republic of Congo. Int. J. Infect. Dis. 105, 735–738 (2021). doi: 10.1016/j.ijid.2021.03.036; pmid: 33737129
- Y. Butera et al., Genomic sequencing of SARS-CoV-2 in Rwanda: Evolution and regional dynamics. medRxiv 2021.04.02.21254839 [Preprint] (2021); https://doi.org/ 10.1101/2021.04.02.21254839.
- C. N Agoti *et al.*, Detection of SARS-CoV-2 variant 501Y.V2 in Comoros Islands in January 2021. *Wellcome Open Res.* 6, 192 (2021). doi: 10.12688/wellcomeopenres.16889.1; pmid: 35071798
- J. M. Morobe et al., Genomic Epidemiology of SARS-CoV-2 in Seychelles, 2020-2021. Viruses 14, 1318 (2022). doi: 10.3390/ v14061318; pmid: 35746789
- C. M. Morang'a et al., Genetic diversity of SARS-CoV-2 infections in Ghana from 2020-2021. Nat. Commun. 13, 2494 (2022). doi: 10.1038/s41467-022-30219-5; pmid: 35523782
- C. N. Agoti et al., Transmission networks of SARS-CoV-2 in Coastal Kenya during the first two waves: A retrospective genomic study. eLife 11, e71703 (2022). doi: 10.7554/ eLife.71703; pmid: 35699426
- S. P. C. Brand et al., COVID-19 transmission dynamics underlying epidemic waves in Kenya. Science 374, 989–994 (2021). doi: 10.1126/science.abk0414; pmid: 34618602
- G. Githinji et al., Tracking the introduction and spread of SARS-CoV-2 in coastal Kenya. Nat. Commun. 12, 4809 (2021). doi: 10.1038/s41467-021-25137-x; pmid: 34376689
- H. Tegally et al., Sixteen novel lineages of SARS-CoV-2 in South Africa. Nat. Med. 27, 440–446 (2021). doi: 10.1038/s41591-021-01255-3; pmid: 33531709
- W. H. Roshdy *et al.*, SARS-CoV-2 Genetic diversity and lineage dynamics of in Egypt. medRxiv 2022.01.05.22268646 [Preprint] (2022); https://doi.org/10.1101/2022.01.05.22268646
- C. Scheepers et al., Emergence and phenotypic characterization of the global SARS-CoV-2 C.1.2 lineage. Nat. Commun. 13, 1976 (2022). doi: 10.1038/s41467-022-29579-9; pmid: 35396511
- G. Dudas et al., Emergence and spread of SARS-CoV-2 lineage B.1.620 with variant of concern-like mutations and deletions. Nat. Commun. 12, 5769 (2021). doi: 10.1038/s41467-021-26055-8; pmid: 34599175
- D. L. Bugembe et al., Emergence and spread of a SARS-CoV-2 lineage A variant (A.23.1) with altered spike protein in Uganda. Nat. Microbiol. 6, 1094–1101 (2021). doi: 10.1038/ s41564-021-00933-9; pmid: 34163035
- 34. H. Tegally et al., Emergence of SARS-CoV-2 Omicron lineages BA.4 and BA.5 in South Africa. Nat. Med. (2022). doi: 10.1038/s41591-022-01911-2; pmid: 35760080
- S. A. Madhi et al., Population immunity and Covid-19 severity with omicron variant in South Africa. N. Engl. J. Med. 386, 1314–1326 (2022). doi: 10.1056/NEJMoa2119658; pmid: 35196424
- N. Wolter et al., Early assessment of the clinical severity of the SARS-CoV-2 omicron variant in South Africa: A data linkage study. Lancet 399, 437–446 (2022). doi: 10.1016/S0140-6736 (22)00017-4; pmid: 35065011
- H. C. Lewis et al., SARS-CoV-2 infection in Africa: A systematic review and meta-analysis of standardised seroprevalence studies, from January 2020 to December 2021. BMJ Glob. Health 7, e008793 (2022). doi: 10.1136/bmjgh-2022-008793; pmid: 35998978
- R. M. Barber et al., Estimating global, regional, and national daily and cumulative infections with SARS-CoV-2 through Nov 14, 2021: A statistical analysis. Lancet 399, 2351–2380 (2022). doi: 10.1016/S0140-6736(22)00484-6; pmid: 35405084
- 39. J. Quick, nCoV-2019 sequencing protocol v3 (LoCost) (2020).
- J. R. Tyson, P. James et al., Improvements to the ARTIC multiplex PCR method for SARS-CoV-2 genome sequencing using nanopore. bioRxiv 2020.09.04.283077 [Preprint] (2020); https://doi.org/10.1101/2020.09.04.283077.
- M. Cotten, D. Lule Bugembe, P. Kaleebu, M. V T Phan, Alternate primers for whole-genome SARS-CoV-2 sequencing. Virus Evol. 7, veab006 (2021). doi: 10.1093/ve/veab006; pmid: 33841912
- H. Tegally et al., A novel and expanding SARS-CoV-2 variant, B.1.318, dominates infections in Mauritius. medRxiv 2021.06.16.21259017 [Preprint] (2021); https://doi.org/ 10.1101/2021.06.16.21259017.
- A. N. Zekri et al., Characterization of the SARS-CoV-2 genomes in Egypt in first and second waves of infection. Sci. Rep. 11, 21632 (2021). doi: 10.1038/s41598-021-99014-4; pmid: 34732835
- 44. C. Nasimiyu et al., Imported SARS-COV-2 variants of concern drove spread of infections across Kenya during the second

- year of the pandemic. *COVID* **2**, 586–598 (2022). doi: 10.3390/covid2050044; pmid: 35262086
- M. U. G. Kraemer et al., Spatiotemporal invasion dynamics of SARS-CoV-2 lineage B.1.1.7 emergence. Science 373, 889–895 (2021). doi: 10.1126/science.abj0113; pmid: 34301854
- S. J. Lycett et al., COVID-19 Genomics UK (COG-UK) Consortium, Epidemic waves of COVID-19 in Scotland: a genomic perspective on the impact of the introduction and relaxation of lockdown on SARS-CoV-2. medRxiv 2021.01.08.20248677 [Preprint] (2021); https://doi.org/ 10.1101/2021.01.08.20248677.
- P. W. G. Mallon et al., Whole-genome sequencing of SARS-CoV-2 in the Republic of Ireland during waves 1 and 2 of the pandemic. medRxiv 2021.02.09.21251402 [Preprint] (2021); https://doi.org/10.1101/2021.02.09.21251402.
- H. Tegally et al., Rapid replacement of the Beta variant by the Delta variant in South Africa. medRxiv 2021.09.23.21264018 [Preprint] (2021); https://doi.org/10.1101/2021.09.23.21264018.
- S. Chang et al., Mobility network models of COVID-19 explain inequities and inform reopening. Nature 589, 82–87 (2021). doi: 10.1038/s41586-020-2923-3; pmid: 33171481
- M. Chinazzi et al., The effect of travel restrictions on the spread of the 2019 novel coronavirus (COVID-19) outbreak. Science 368, 395–400 (2020). doi: 10.1126/science.aba9757; pmid: 32144116
- M. U. G. Kraemer et al., The effect of human mobility and control measures on the COVID-19 epidemic in China. Science 368, 493–497 (2020). doi: 10.1126/science.abb4218; pmid: 32213647
- P. Nouvellet et al., Reduction in mobility and COVID-19 transmission. Nat. Commun. 12, 1090 (2021). doi: 10.1038/ s41467-021-21358-2; pmid: 33597546
- C. Xiong, S. Hu, M. Yang, W. Luo, L. Zhang, Mobile device data reveal the dynamics in a positive relationship between human mobility and COVID-19 infections. *Proc. Natl. Acad. Sci. U.S.A.* 117, 27087–27089 (2020). doi: 10.1073/ pnas.2010836117; pmid: 33060300
- S. Pillay et al., Whole genome sequencing of SARS-CoV-2: Adapting Illumina protocols for quick and accurate outbreak investigation during a pandemic. Genes 11, 949 (2020). doi: 10.3390/genes11080949; pmid: 32824573
- L. Singh et al., Targeted Sanger sequencing to recover key mutations in SARS-CoV-2 variant genome assemblies produced by next-generation sequencing. Microb. Genom. 8, (2022). doi: 10.1099/mgen.0.000774; pmid: 35294336
- A. J. Page et al., Large-scale sequencing of SARS-CoV-2 genomes from one region allows detailed epidemiology and enables local outbreak management. Microb. Genom. 7, (2021). doi: 10.1099/mgen.0.000589; pmid: 34184982
- N. De Maio, C. Walker, R. Borges, L. Weilguny, G. Slodkowicz, N. Goldman, Issues with SARS-CoV-2 sequencing data, Virological.org (2020); https://virological.org/t/issues-withsars-cov-2-sequencing-data/473.
- N. E. Freed, M. Vlková, M. B. Faisal, O. K. Silander, Rapid and inexpensive whole-genome sequencing of SARS-CoV-2 using 1200 bp tiled amplicons and Oxford Nanopore Rapid Barcoding. *Biol. Methods Protoc.* 5, bpaa014 (2020). doi: 10.1093/biomethods/bpaa014; pmid: 33029559
- J.-S. Eden et al., An emergent clade of SARS-CoV-2 linked to returned travellers from Iran. Virus Evol. 6, veaa027 (2020). doi: 10.1093/ve/veaa027; pmid: 32296544
- A. S. Gonzalez-Reiche et al., Introductions and early spread of SARS-CoV-2 in the New York City area. Science 369, 297–301 (2020). doi: 10.1126/science.abc1917: pmid: 32471856
- K. Itokawa, T. Sekizuka, M. Hashino, R. Tanaka, M. Kuroda, Disentangling primer interactions improves SARS-CoV-2 genome sequencing by multiplex tiling PCR. PLOS ONE 15, e0239403 (2020). doi: 10.1371/journal.pone.0239403; pmid: 32946527
- A. X. Han et al., Low testing rates limit the ability of genomic surveillance programs to monitor SARS-CoV-2 variants: a mathematical modelling study. medRxiv 2022.05.20. 22275319 [Preprint] (2022):https://doi.org/10.1101/ 2022.05.20.22275319 .
- H. Wang et al., Estimating excess mortality due to the COVID-19 pandemic: A systematic analysis of COVID-19-related mortality, 2020-21. Lancet 399, 1513–1536 (2022). doi: 10.1016/S0140-6736(21)02796-3; pmid: 35279232
- F. Wu et al., A new coronavirus associated with human respiratory disease in China. Nature 579, 265–269 (2020). doi: 10.1038/s41586-020-2008-3; pmid: 32015508
- 65. I. Aksamentov, C. Roemer, E. Hodcroft, R. Neher, Nextclade: Clade assignment, mutation calling and quality control for viral

- genomes. J. Open Source Softw. **6**, 3773 (2021). doi: 10.21105/ioss.03773
- D. P. Martin et al., RDP5: A computer program for analyzing recombination in, and removing signals of recombination from, nucleotide sequence datasets. Virus Evol. 7, veaa087 (2020). doi: 10.1093/ve/veaa087; pmid: 33936774
- M. N. Price, P. S. Dehal, A. P. Arkin, FastTree 2—Approximately maximum-likelihood trees for large alignments. *PLOS ONE* 5, e9490 (2010). doi: 10.1371/journal.pone.0009490; pmid: 20224823
- J. Felsenstein, Confidence limits on phylogenies: An approach using the bootstrap. Evolution 39, 783–791 (1985). doi: 10.1111/j.1558-5646.1985.tb00420.x; pmid: 28561359
- A. Rambaut, T. T. Lam, L. Max Carvalho, O. G. Pybus, Exploring the temporal structure of heterochronous sequences using TempEst (formerly Path-O-Gen). Virus Evol. 2, vew007 (2016). doi: 10.1093/ve/vew007; pmid: 27774300
- A.-A. Popescu, K. T. Huber, E. Paradis, ape 3.0: New tools for distance-based phylogenetics and evolutionary analysis in R. *Bioinformatics* 28, 1536–1537 (2012). doi: 10.1093/ bioinformatics/bts184; pmid: 22495750
- P. Sagulenko, V. Puller, R. A. Neher, TreeTime: Maximumlikelihood phylodynamic analysis. Virus Evol. 4, vex042 (2018). doi: 10.1093/ve/vex042; pmid: 29340210
- S. Wang et al., Molecular evolutionary characteristics of SARS-CoV-2 emerging in the United States. J. Med. Virol. 94, 310–317 (2022). doi: 10.1002/jmv.27331; pmid: 34506640
- G. Yu, Using ggtree to visualize data on tree-like structures. Curr. Protoc. Bioinformatics 69, e96 (2020). doi: 10.1002/ cpbi.96; pmid: 32162851
- H. Wickham, ggplot2. Wiley Interdiscip. Rev. Comput. Stat. 3, 180–185 (2011). doi: 10.1002/wics.147
- J. Hadfield et al., Nextstrain: Real-time tracking of pathogen evolution. Bioinformatics 34, 4121–4123 (2018). doi: 10.1093/ bioinformatics/bty407; pmid: 29790939
- S. E. James, CERI-KRISP/SARS-CoV-2-epidemic-in-Africa: Expanding Africa SARS-CoV-2 sequencing capacity in a fast evolving pandemic analysis. Zenodo (2022); https://doi.org/ 10.5281/zenodo.7006806.
- T. Ward, A. Johnsen, Understanding an evolving pandemic: An analysis of the clinical time delay distributions of COVID-19 in the United Kingdom. *PLOS ONE* 16, e0257978 (2021). doi: 10.1371/journal.pone.0257978; pmid: 34669712

ACKNOWLEDGMENTS

First and foremost, we acknowledge authors in institutions in Africa and beyond who have made invaluable contributions toward specimen collection and sequencing to produce and share, via GISAID, SARS-CoV-2 genomic data. We also acknowledge the authors from the originating and submitting laboratories worldwide who generated and shared SARS-CoV-2 sequence data, via GISAID, from other regions in the world, which was used to contextualize the African genomic data. A full list of GISAID sequence IDs used in the current study is available in table S4. Funding: Sequencing efforts in the African Union Member States were supported by the Africa Centers for Disease Control (Africa CDC)-Africa Pathogen Genomics Initiative (Africa PGI) and the World Health Organization Regional Office for Africa (WHO AFRO) through the transfer of laboratory infrastructure, the provision of reagents, and training. The Africa PGI is supported by the African Union, US Centers for Disease Control and Prevention (CDC), Bill & Melinda Gates Foundation, Illumina Inc., Oxford Nanopore Technologies, and other partners. In addition, all Institut Pasteur organizations and CERMES in Niger are part of the PEPAIR COVID-19-Africa project, which is funded by the French Ministry for European and Foreign Affairs. The KwaZulu-Natal Research Innovation and Sequencing Platform (KRISP) and Centre for Epidemic Response and Innovation (CERI) are supported in part by grants from WHO, the Rockefeller Foundation (HTH 017), the Abbott Pandemic Defense Coalition (APDC), the US National Institutes of Health (NIH) (U01 Al151698) for the United World Antivirus Research Network (UWARN) and the INFORM Africa project through the Institute of Human Virology, Nigeria (IHVN) (U54 TW012041), H3BioNet Africa (grant no. 2020 HTH 062), the World Bank (TF0B8412), the South African Department of Science and Innovation (SA DSI), and the South African Medical Research Council (SAMRC) under the BRICS JAF #2020/049. The International Livestock Research Institute (ILRI) is also supported by the Ministry for Economic Cooperation and Federal Development of Germany (BMZ). Work conducted at the African Centre of Excellence for Genomics of Infectious Diseases (ACEGID) is made possible by support provided to ACEGID by a cohort of generous donors through TED's Audacious Project, including the ELMA Foundation, MacKenzie Scott, the Skoll Foundation, and Open

Philanthropy. Work at ACEGI) was also partly supported by grants from the National Institute of Allergy and Infectious Diseases (NIAID) (https://www.niaid.nih.gov) NIH-H3Africa (https://h3africa.org) (U01HG007480 and U54HG007480), the World Bank (projects ACE-019 and ACE-IMPACT), the Rockefeller Foundation (grant #2021 HTH), the Africa CDC through the African Society of Laboratory Medicine (ASLM) (grant #INV018978), the Wellcome Trust (project 216619/7/19/7), and the Science for Africa Foundation, Sequencing efforts at the National Institute for Communicable Diseases (NICD) were also supported by a conditional grant from the South African National Department of Health as part of the emergency COVID-19 response; a cooperative agreement between the NICD of the National Health Laboratory Service (NHLS) and the CDC (FAIN# U01IP001048 and NU51IP000930); the South African Medical Research Council (SAMRC) (project number 96838); the ASLM and the Bill & Melinda Gates Foundation (grant number INV-018978); the UK Foreign, Commonwealth and Development Office and the Wellcome Trust (grant no. 221003/Z/20/Z); and the UK Department of Health and Social Care and were managed by the Fleming Fund and performed under the auspices of the SEQAFRICA project. The NICD also acknowledges support from Hyrax Biosciences for the use of their Exatype platform. This was made possible through funding from the South African Medical Research Council, the Department of Science and Innovation, as well as support from the Health Equity Initiative at Amazon Web Services, Funding for sequencing efforts in Angola were supported through Projecto Bongola (N.º 11/ MESCTI/PDCT/2020) and Orçamento Geral do Estado Instituto Nacional de Investigação de Saúde (OGE INIS) (2020/2021). Botswana's sequencing efforts, which were led by the Botswana Harvard AIDS Institute Partnership, were supported by the Foundation for Innovative New Diagnostics (FINDdx), Bill & Melinda Gates Foundation H3ABioNet (U41HG006941), Sub-Saharan African Network for TB/HIV Research Excellence (SANTHE), and Fogarty International Center (grant no. 5D43TW009610). H3ABioNet is an initiative of the Human Health and Heredity in Africa Consortium (H3Africa) program of the African Academy of Science (AAS) and the US Department of Health and Human Services (HHS), NIH, and NIAID (5K24AI131928-04; 5K24AI131924-04); SANTHE is a DELTAS Africa Initiative (grant no. DEL-15-006). The DELTAS Africa Initiative is an independent funding scheme of the AAS's Alliance for Accelerating Excellence in Science in Africa (AESA) and is supported by the New Partnership for Africa's Development Planning and Coordinating Agency (NEPADAgency) with funding from the Wellcome Trust (grant #107752/Z/15/Z) and the UK government, From Brazil, J.S.X. was funded by Coordenação de Aperfeicoamento de Pessoal de Nível Superior-Brazil (CAPES)-Finance Code 001. Sequencing efforts from Côte d'Ivoire were funded by the Robert Koch Institute and the German Federal Ministry of Education and Research (BMBF). Sequencing efforts in the Democratic Republic of the Congo were funded by the Bill & Melinda Gates Foundation under grant INV-018030 awarded to C.B.P. and further supported by funding from the Africa CDC through ASLM for Accelerating SARS-CoV-2 Genomic Surveillance in Africa, the CDC, US Army Medical Research Institute of Infectious Diseases (USAMRIID), Institut de Recherche pour le Développement (IRD)/Montepellier, University of California-Los Angeles (UCLA), and SACIDS FIND. Efforts from Egypt were funded by the Egyptian Ministry of Health, the Egyptian Academy for Scientific Research and Technology (ASRT) JESOR project #3046 (Center for Genome and Microbiome Research), the Cairo University anti-COVID-19 fund, and the Science and Technology Development Fund (STDF), project ID 41907. The sequencing effort in Equatorial Guinea was supported by a publicprivate partnership, the Bioko Island Malaria Elimination Project, which is composed of the government of Equatorial Guinea Ministries of Mines and Hydrocarbons, and Health and Social Welfare, Marathon EG Production Limited, Noble Energy, Atlantic Methanol Production Company, and EG LNG. Analysis for the Gabon strains was supported by the Science and Technology Research Partnership for Sustainable Development (SATREPS), Japan International Cooperation Agency (JICA), and Japan Agency for Medical Research and Development (AMFD) (grant number JP21jm0110013) and a grant from AMED (grant number JP21wm0225003). The Centre Interdisciplinaires de Recherches Medicales de Franceville (CIRMF) (Gabon) is funded by the Gabonese Government and TOTAL Energy inc. CIRMF is a member of the Central Africa Network on Tuberculosis HIV/AIDS and Malaria (CANTAM), which is supported by the European and Developing Countries Clinical Trials Partnership (EDCTP). The work at the West African Centre for Cell Biology of Infectious Pathogens (WACCBIP) (Ghana) was funded by a grant from the Rockefeller Foundation (2021 HTH 006), an IRD grant (ARIACOV), an African Research Universities Alliance (ARUA) Vaccine Development Hubs

grant with funds from Open Society Foundation, National Institute of Health Research (NIHR) (17.63.91) grants using UK aid from the UK Government for a global health research group for genomic surveillance of malaria in West Africa (Wellcome Sanger Institute, UK), and a World Bank African Centers of Excellence Impact grant (WACCBIP-NCDs: Awandare). In addition to the funding sources from ILRI, Kenya Medical Research Institute (KEMRI) (Kenyan) contributions to sequencing efforts were supported in part by the National Institute for Health Research (NIHR (project references 17/63/82 and 16/136/33) using UK aid from the UK government to support global health research; the UK Foreign, Commonwealth and Development Office (FCDO) and the Wellcome Trust (grant no. 220985/Z/20/Z); and the Kenya Medical Research Institute (grant no. KEMRI/COV/SPE/012). Contributions from Lesotho were supported by the Africa CDC, ALSM, and South Africa NICD. Liberian efforts were funded by the Africa CDC through a subaward from the Bill & Melinda Gates Foundation, and efforts from Madagascar were funded by the French Ministry for Europe and Foreign Affairs through the REPAIR COVID-19-Africa project coordinated by the Pasteur International Network association. Sequencing from Malawi was supported by the Wellcome Trust. Contributions from Mali were supported by Fogarty International Center and NIAID sections of the NIH under Leidos-15X051, award numbers U2RTW010673 for the West African Center of Excellence for Global Health Bioinformatics Research Training and U19Al089696 and U19Al129387 for the West Africa International Center of Excellence for Malaria Research. Funding for surveillance, sampling, and testing in Madagascar was provided by the WHO, the CDC (grant no. U5/IP000812-05), the United States Agency for International Development (USAID) (Cooperation Agreement 72068719CA00001), and the Office of the Assistant Secretary for Preparedness and Response in the HHS (grant no. IDSEP190051-01-0200). Funding for sequencing was provided by the Bill & Melinda Gates Foundation (GCE/ID OPP1211841), Chan Zuckerberg Biohub, and the Innovative Genomics Institute at UC Berkeley. Mozambique acknowledges support from the Mozambican Ministry of Health and the President's Emergency Plan for AIDS Relief (PEPFAR) through the CDC under the terms of grant nos. GH002021 and GH001944 and from the Bill & Melinda Gates Foundation (#OPP1214435). Namibian efforts were supported by Africa CDC through a subaward from the Bill & Melinda Gates Foundation, Efforts from Niger were supported by the French Ministry for Europe and Foreign Affairs through the REPAIR COVID-19-Africa project coordinated by the Pasteur International Network association. In addition to the funding support for ACEGID already listed, Nigeria's contributions were made possible by support from Flu Lab and a cohort of donors through the Audacious Project, a collaborative funding initiative housed at TED, including the ELMA Foundation, MacKenzie Scott, the Skoll Foundation, and Open Philanthropy. COVID-19 genomic surveillance at the Centre for Human Virology and Genomics, Nigerian Institute of Medical Research, is supported by the government of Nigeria special funding for COVID-19 to the Nigerian Institute of Medical Research, Lagos, Nigeria. It is also supported by funding from the AIDS Healthcare Foundation (AHF) Global Public Health Institute (GESIT Study). Efforts from the Republic of the Congo were supported by the European and Developing Countries Clinical Trials Partnership (EDCTP) IDs PANDORA and CANTAM and the German Academic Exchange Service (DAAD) ID PACE-UP and DAAD project ID 5759234. Rwanda's contributions were made possible by funding from the African Network for Improved Diagnostics, Epidemiology and Management of Common Infectious Agents (ANDEMIA), which was granted by the German Federal Ministry of Education and Research (BMBF grants 01KA1606, 01KA2021, and 01KA2110B) and the NIHR Global Health Research program (16/136/ 33) using UK aid from the UK Government. In addition to the South African institutions listed above, the University of Cape Town's work was supported by the Wellcome Trust (grant no. 203135/Z/16/Z). EDCTP RADIATES (RIA2020EF-3030), the South African Department of Science and Innovation (SA DSI), and SAMRC; Stellenbosch University's contributions were supported by SAMRC; and the University of Pretoria's contributions were funded by the G7 Global Health Fund and a BMBF ANDEMIA grant. Funding from the Fleming Fund supported sequencing in Sudan. The Ministry of Higher Education and Scientific Research of Tunisia provided funding for sequencing from Tunisia. The Uganda Virus Research Institute (UVRI) (Uganda) acknowledge support from the Wellcome Trust and FCDO - Wellcome Epidemic Preparedness -Coronavirus (AFRICO19, grant agreement number 220977/Z/20/Z). the MRC (MC_UU_1201412), and ,the UK Medical Research Council (MRC/UKRI) and FCDO (DIASEQCO, grant agreement number MC_PC_20010). Research at the FredHutch Institute, which supported bioinformatics analyses of sequences in the present study, was supported by the Bill & Melinda Gates foundation (#INV-018979). Research support from Broad Institute colleagues was made possible

by support from Flu Lab and a cohort of generous donors through TED's Audacious Project, including the ELMA Foundation, MacKenzie Scott, the Skoll Foundation, Open Philanthropy, the Howard Hughes Medical Institute, and NIH (U01AI151812 and U54HG007480) (P.C.S.). Work from Quadram Institute Bioscience was funded by the Biotechnology and Biological Sciences Research Council (BBSRC) Institute Strategic Programme Microbes in the Food Chain BB/ R012504/1 and its constituent projects BBS/F/F/000PR10348, BBS/ F/F/000PR10349, BBS/F/F/000PR10351, and BBS/F/F/ 000PR10352 and by the Quadram Institute Bioscience BBSRC-funded Core Capability Grant (project number BB/CCG1860/1). Sequences generated in Zambia through PATH were funded by the Bill & Melinda Gates Foundation and Africa CDC. The content and findings reported herein are the sole deduction, view, and responsibility of the researcher(s) and do not reflect the official position and sentiments of the funding agencies. Author contributions: Conceptualization: H.Te., C.Ba., S.K.T., T.d.O., R.L., E.W.; Methodology: H.Te., J.E.S., M.C., B.Te., G.M., D.P.M., A.W.L., D.A.R., L.M.K., G.G., T.d.O., R.L., E.W.; Genomic data generation: H.Te., J.E.S., M.C., M.Moi., B.Te., G.M., D.P.M., A.W.L., A.D., D.G.A., M.M.D., A.Si., A.N.Z., A.S.G., A.K.Sa., A.O., A.Sow, A.O.M., A.K.Se., A.G.A., A.L., A.-S.K., A.E.A., A.A.J., A.Fo., A.O.O. A.A.A., A.J., A.Kan., A.Mo., A.R., A.Sa., A.Kaz., A.Ba., A.Chr., A.J.T., A.Ca., A.K.K., A.Ko., A.Bo., A.Sou., A.A., A.Na., A.V.G., A.Nk., A.J.P., A.Y., A.V., A.N.H., A.Cho., A.Ir., A.Ma., A.L.B., A.Is., A.A.Sv., A.G., A.Fe., A.E.S., B.Ma., B.L.S., B.S.O., B.B., B.D., B.L.H., B.Ts., B.L., B.Mv., B.N., B.T.M., B.A.K., B.K., B.A., B.P., B.Mc., C.Br., C.W., C.N., C.A., C.B.P., C.S., C.G.A., C.N.A., C.M.M., C.L., C.K.O., C.I., C.N.M., C.P., C.G., C.E.O., C.D.R., C.M.M., C.E., D.B.L., D.J.B., D.M., D.P., D.B., D.J.N., D.S., D.T., D.S.A., D.G., D.S.G., D.O.O., D.M., D.W.W., E.F., E.K.L., E.Si., E.M.O., E.N.N., E.O.A., E.O., E.Sh., E.Ba., E.B.A., E.A.Ah., E.L., E.Mu., E.P., E.Be., E.S.-L., E.A.An., F.L., F.M.T., F.W., F.A., F.T.T., F.D., F.V.A., F.T., F.O., F.N., F.M.M., F.E.R. F.A.D., F.I., G.K.M., G.T., G.L.K., G.O.A., G.U.v.Z., G.A.A., G.S., G.P.M., H.C.R., H.E.O., H.O., H.A., H.K., H.N., H.Tr., H.A.A.K., H.E., H.G., H.M., H.K., I.Sm., I.B.O., I.M.A., I.O., I.B.B., I.A.M., I.Ss., I.W., I.S.K., J.W.A.H., J.A., J.S., J.C.M., J.M.T., J.H., J.G.S., J.Gi., J.Mu., J.N., J.N.U., J.N.B., J.Y., J.Mo., J.K., J.D.S., J.H., J.K.O., J.M.M., J.O.G., J.T.K., J.C.O., J.S.X., J.Gy., J.F.W., J.H.B., J.N., J.E., J.N., J.M.N., J.N., J.U.O., J.C.A., J.J.L., J.J.H.M., J.O., K.J.S., K.V., K.T.A., K.A.T., K.S.C., K.S.M., K.D., K.G.M., K.O.D., L.F., L.S., L.M.K., L.B., L.d.O.M., L.C., L.O., L.D.O., L.L.D., L.I.O., L.T., M.Mi., M.R., M.Mas., M.E., M.Mai., M.I.M., M.Ke., M.D., M.Mom., M.d.L.L.M., M.V., M.F.P., M.F., M.M.N., M.Mar., M.D., M.W.M., M.G.M., M.O., M.R.W., M.Y.T., M.O.A., M.Ab., M.A.B., M.G.S., M.K.K., M.M.M., M.Ka., M.S., M.B.M., M.Mw., M.Al., M.V.P., N.Abi., N.R., N.Abr., N.Is., N.E., N.M.T., N.D., N.Ma., N.H., N.B.S., N.M.F., N.Sa., N.B., N.Mu., N.G., N.W., N.Si., N.N., N.A.A., N.T., N.Mbh., N.H.R., N.Ig., N.Mba, O.C.K., O.S., O.Fe., O.M.A., O.Te., O.A.O., O.Fak., O.E.O., O.-E.O., O.Fay., P.S., P.O., P.C., P.N., P.S., P.E.O., P.Ar., P.K.O., P.O.O., P.B., P.D., P.A.B., P.K.M., P.K., P.Ab., R.E., R.J., R.K.A., R.G.E., R.A., R.N., R.O.P., R.G., R.A.K., R.M.N.D., R.A.A., R.A.C., S.Gar., S.Ma., S.Bo., S.S., S.I.M., S.F., S.Mh., S.H., S.K.K., S.Me., S.T., S.H.A., S.W.M., S.D., S.M.-M., S.A. S.S.A., S.M.A., S.E., S.Mo., S.L., S.Gas., S.J., S.F.A., S.Og., S.Gr., S.L., S.Pr., S.Ou., S.v.W., S.F.S., S.K., S.A., S.R., S.Pi., S.N., S.Be., S.L.B., S.v.d.W., T.Ma., T.Mo., T.L., T.P.V., T.S., T.G.M., T.B., U.J.A., U.C., U.R., U.E.G., V.E., V.N., V.G., W.H.R., W.A.K., W.K.A., W.P. W.T.C., Y.A.A., Y.R., Y.Be., Y.N., Y.Bu., Z.R.d.L., A.E.O., A.v.G., G.G., M.Moe., O.To., P.C.S., A.A.Sa., S.O.O., Y.K.T., S.K.T., T.d.O., C.H., R.L., J.N., E.W.; Data analysis: H.Te., J.E.S., M.C., M.Moi., B.Te., G.M., D.P.M., A.W.L., A.L.F., D.A.R., F.M., G.S.K., S.v.W., G.G., T.d.O., R.L., E.W.; Funding acquisition: A.E.O., A.vG., G.G., M.Moe., O.To., A.A.Sa., S.O.O., Y.K.T., S.K.T., T.d.O., C.H.; Project administration: G.M., A.D., D.G.A., M.M.D., A.C., D.W.W., H.O., S.W.M., A.E.O., A.v.G., G.G., M.Moe., O.To., P.C.S., A.A.Sa., S.O.O., Y.K.T., S.K.T., T.d.O., C.H., R.L., J.N., E.W.: Supervision: A.E.O., A.v.G., G.G., M.Moe., O.To., P.C.S., A.A.Sa., S.O.O., Y.K.T., S.K.T., T.d.O., C.H., R.L., J.N., E.W.; Writing-original draft: H.Te., J.E.S., M.C., G.M., D.P.M., C.Ba., S.K.T., T.d.O., R.L., E.W.; Writing-review and editing: H.Te., J.E.S., M.C., M.Moi., B.Te., G.M., D.P.M., C.Ba., A.W.L., A.D., D.G.A., M.M.D., A.Si., A.N.Z., A.S.G., A.K.Sa., A.O., A.Sow, A.O.M., A.K.Se., A.I.E., A.L., A.-S.K., A.E.A., A.A.J., A.Fo., A.O.O., A.A.A., A.J., A.Kan., A.Mo., A.R., A.Sa., A.Kaz., A.Ba., A.Chr., A.J.T., A.Ca., A.K.K., A.Ko., A.Bo., A.Sou., A.A., A.V.G., A.J.P., A.Y., A.V., A.N.H., A.Cho., A.Ir., A.Ma., A.L.B., A.Is., A.A.Sy., A.G., A.Fe., A.E.S., B.Ma., B.L.S., B.S.O., B.B., B.D., B.L.H., B.Ts., B.L., B.Mv., B.N., B.T.M., B.A.K., B.K., B.A., B.P., B.Mc., C.Br., C.W., C.A., C.B.P., C.S., C.G.A., C.N.A., C.M.M., C.L., C.K.O., C.I., C.N.M., C.P., C.E.O., C.D.R., C.M.M., C.E., D.B.L., D.J.B., D.M., D.P., D.B., D.J.N., D.S., D.T., D.S.A., D.G., D.S.G., D.O.O., D.M., D.W.W., E.F., E.K.L., E.Si., E.M.O., E.N.N., E.O.A., E.O., E.Sh., E.Ba., E.B.A., E.L., E.Mu., E.P., E.Be., E.S.-L., E.A.An., E.Ma., F.L., F.M.T., F.W., F.A., F.T.T., F.D., F.V.A., F.T.,

F.O., F.N., F.M.M., F.E.R., F.A.D., F.I., G.K.M., G.T., G.L.K., G.O.A.,

G.U.v.Z., G.A.A., G.S.K., G.S., G.P.M., H.C.R., H.E.O., H.O., H.A., H.K., H.N., H.Tr., H.A.A.K., H.E., H.G., H.M., H.K., I.Sm., I.B.O., I.M.A., I.O., I.B.B., I.Ss., I.W., I.S.K., J.W.A.H., J.A., J.S., J.C.M., J.M.T., J.H., J.G.S., J.Gi., J.Mu., J.N.U., J.N.B., J.Y., J.Mo., J.K., J.D.S., J.H., J.K.O., J.M.M., J.O.G., J.T.K., J.C.O., J.S.X., J.Gy., J.H.B., J.N., J.E., J.N., J.M.N., J.N., J.U.O., J.C.A., J.J.L., J.O., K.J.S., K.V., K.T.A., K.A.T., K.S.C., K.S.M., K.D., K.G.M., K.O.D., L.F., L.S., L.B., L.d.O.M., L.C., L.O., L.L.D., L.I.O., M.Mi., M.R., M.Mas., M.E., M.Mai., M.I.M., M.Ke., M.D., M.Mom., M.d.L.L.M., M.V., M.F.P., M.F., M.M.N., M.Mar., M.D., M.W.M., M.G.M., M.O., M.R.W., M.Y.T., M.O.A., M.Aab., M.A.B., M.G.S., M.K.K., M.M.M., M.Ka., M.S., M.B.M., M.Mw., M.V.P., N.Abi., N.R., N.Is., N.M.T., N.D., N.Ma., N.H., N.B.S., N.M.F., N.Sa., N.B., N.Mu., N.G., N.W., N.Si., N.N., N.A.A., N.T., N.Mbh., N.H.R., N.Ig., N.Mba, O.C.K., O.S., O.Fe., O.M.A., O.Te., O.A.O., O.Fak., O.E.O., O.Fay., P.S., P.O., P.C., P.N., P.S., P.E.O., P.Ar., P.K.Q., P.O.O., P.B., P.D., P.A.B., P.K.M., P.K., P.Ab., R.E., R.J., R.K.A., R.G.E., R.A., R.N., R.O.P., R.G., R.A.K., R.A.A., R.A.C., S.Gar., S.Ma., S.S., S.I.M., S.F., S.Mh., S.H., S.K.K., S.Me., S.T., S.H.A., S.W.M., S.D., S.M.-M., S.A., S.S.A., S.M.A., S.E., S.Mo., S.L., S.Gas., S.J., S.F.A., S.Og., S.Gr., S.L., S.Pr., S.Ou., S.v.W., S.F.S., S.K., S.A., S.R., S.Pi., S.N., S.Be., S.L.B., S.v.d.W., T.Ma., T.Mo., T.L., T.P.V., T.S., T.G.M., T.B., U.J.A., U.C., U.R., U.E.G., V.E., V.N., V.G., W.H.R., W.A.K., W.K.A., W.P., W.T.C., Y.A.A., Y.R., Y.Be., Y.N., Y.Bu., Z.R.d.L., A.E.O., A.v.G., G.G., M.Moe., O.To., P.C.S., A.A.Sa., S.O.O., Y.K.T., S.K.T., T.d.O., C.H., R.L., J.N., E.W. Competing interests: With the exception of P.S., who is a co-founder of and consultant to Sherlock Biosciences and a Board Member of Danaher Corporation and who holds equity in the companies, the authors have no conflicts of interest to declare. Data and materials availability: All of the SARS-CoV-2 whole-genome sequences that were analyzed in the present study are all publicly available on the GISAID sequence database. We gratefully acknowledge the authors from the originating laboratories and the submitting laboratories, who generated and shared via GISAID genetic sequence data on which this research is based. A full list of the African sequences as well as global references are presented and acknowledged in table S4 and in our github repository (https://github.com/CERI-KRISP/SARS-CoV-2epidemic-in-Africa) (76). The repositories also contain all of the metadata, raw and time-scaled ML tree topologies, and annotated tree topologies, as well as the data analysis and visualization scripts used here, which will allow for the independent reproduction of results. Furthermore, the repositories also contain all institutional review board references and material transfer agreements. Please refer to the ethics statement in the methods section for more details. License information: This work is licensed under a Creative Commons Attribution 4.0 International (CC BY 4.0) license, which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited. To view a copy of this license, visit https:// creativecommons.org/licenses/by/4.0/. This license does not apply to figures/photos/artwork or other content included in the article that is credited to a third party; obtain authorization from the rights holder before using such material.

Authors and their affiliations

Authors and their attiliations
Hourilyah Tegally^{1,2}†, James E. San^{1,2}, Matthew Cotten^{3,4},
Monika Moir¹, Bryan Tegomoh^{5,6}, Gerald Mboowa⁷,
Darren P. Martin^{8,9}, Cheryl Baxter^{1,1,0}, Arnold W. Lambisia¹¹,
Amadou Diallo¹², Daniel G. Amoako^{13,1,4}, Moussa M. Diagne¹² Amadou Diallo¹², Daniel G. Amoako^{13,14}, Moussa M. Diagne¹², Abay Sisay^{15,16}, Abdel-Rahman N. Zekri¹⁷, Abdou Salam Gueye¹⁸, Abdoul K. Sangare¹⁹, Abdoul-Salam Ouedraogo^{20,204,205}, Abdourahmane Sow²¹, Abdualmoniem O. Musa^{22,23,24}, Abdul K. Sesay²⁵, Abe G. Abias²⁶, Adem I. Elzagheid²⁷, Adamou Lagare²⁸, Adelotun-Sulaiman Kemi²⁹, Aden Elmi Abar^{72,73}, Adenjiji A. Johnson^{31,32}, Adeola Fowotada^{33,34}, Aleyemi O. Oluwapelumi^{35,36}, Adrienne A. Amuri^{37,38}, Agnes Juru³⁹, Ahmed Kandeil⁴⁰, Ahmed Mostafa⁴⁰, Ahmed Rebai⁴¹, Ahmed Sayed⁴², Kazeem Akano^{43,44}, Aladje Balde^{45,46}, Alan Christoffels^{7,47}, Alexander I. Trotter⁴⁸, Allan Campholl⁴⁹, Alpha K. Keita^{50,51} Alexander J. Trotter⁴⁸, Allan Campbel⁴⁹, Alpha K. Keita^{50,51}, Amadou Kone⁵², Amal Bouzid^{41,53}, Amal Souissi⁴¹, Ambrose Agweyu¹¹, Amel Naguib⁵⁴, Ana V. Gutierrez⁴⁸, Anatole Nkeshimana⁵⁵, Andrew J. Page⁴⁸, Anges Yadouleton⁵⁶, Anika Vinze⁵⁷, Anise N. Happi⁴³, Anissa Chouikha^{58,59} Arash Iranzadeh^{8,9}, Arisha Maharaj¹, Armel L. Batchi-Bouyou^{60,61}, Arshad Ismail¹³, Augustina A. Sylverken^{62,63},
Augustine Goba^{64,65}, Ayoade Femi^{43,44}, Ayotunde E. Sijuwola⁴³,
Baba Marycelin^{66,67}, Babatunde L. Salako^{29,32}, Bamidele S. Oderinde⁶⁶, Bankole Bolajoko⁴³, Bassirou Diarra⁵², Belinda L. Herring¹⁸, Benjamin Tsofa¹¹, Bernard Lekana-Douki^{68,69}, Bernard Mvula⁷⁰, Berthe-Marie Njanpop-Lafourcade¹⁸, Blessing T. Marondera⁷¹,

Bouh Abdi Khaireh^{72,73}, Bourema Kouriba¹⁹, Bright Adu⁷⁴, Brigitte Pool⁷⁵, Bronwyn McInnis⁷, Cara Brook^{76,77}, Carolyn Williamson^{9,10,78}, Cassien Nduwimana⁵⁵, Catherine Anscombe^{79,80}, Catherine B. Pratt⁸¹, Cathrine Scheepers^{13,82}, Chantal G. Akoua-Kofff^{83,84}, Charles N. Agoti^{11,85}, Chastel M. Mapanguy^{60,86}, Chairles N. Agott F. Chastel M. Mapanguy F. Cheikh Loucoubar¹², Chiak K. Onwuamah⁸⁷, Chikwe Ihekweazu⁸⁸, Christian N. Malaka⁸⁹, Christophe Peyrefitte¹², Grace C. Chukwa^{43,44}, Chukwuma E. Omoruyi^{33,34}, Grace C. Chukwa — Chukwuma E. Omoruyi Clotaire D. Rafai³⁰, Collins M. Morangʻa³¹, Cyril Erameh⁹², Daniel B. Lule³, Daniel J. Bridges³³, Daniel Mukadi-Bamuleka³⁷, Danny Park⁵⁷, David A. Rasmussen^{94,95}, David Baker⁴⁸, David J. Nokes^{11,96}, Deografius Ssemwanga^{3,97}, Derek Tshiabuila², Dominic S. Y. Amuzu⁹¹ Dominique Goedhals⁹⁸, Donald S. Grant^{64,65,99}, Donwilliams O. Omuoyo¹¹, Dorcas Maruapula¹⁰⁰, Dorcas W. Wanjohi⁷, Ebenezer Foster-Nyarko⁴⁸, Eddy K. Lusamaki^{37,38,51}, Edgar Simulundu¹⁰¹, Edidah M. Ong'era¹¹, Edith N. Ngabana^{37,38}, Edward O. Abworo¹⁰², Edward O'tieno¹¹, Edwin Shumba⁷¹, Edwine Barasa¹¹, El Bara Ahmed^{103,104}, Elhadi A. Ahmed²³, Edwine Barasa , El Bara Arimed , Eliradi A. Arimed , Emmanuel Lokilo³⁷, Enatha Mukantwari¹⁰⁵, Eromon Philomena⁴³, Essia Belarhi¹⁰⁶, Etienne Simon-Loriere¹⁰⁷, Etilé A. Anoh⁸³, Eusebio Manuel¹⁰⁸, Fabian Leendertz¹⁰⁶, Fahn M. Taweh¹⁰⁹, Fares Wasfi⁵⁸, Fatma Abdelmoula^{41,110}, Faustinos T. Takawira³⁹, Fawzi Derrar¹¹¹, Fehintola V. Ajogbasile⁴³ Florette Treurnicht^{112,113}, Folarin Onikepe^{43,44} Francine Ntoumi^{60,114}, Francisca M. Muyembe^{37,38}, Frank E. Z. Ragomzingba¹¹⁵, Fred A. Dratibi^{116,117}, Fred-Akintunwa iyanu⁴³, Gabriel K. Mbunsu³⁸, Gaetan Thilliez⁴⁸, Gemma L. Kay⁴⁸, George O. Akpede⁹², Gert U. van Zyl^{118,119}, Gordon A. Awandare⁹¹, Grace S. Kpeli^{120,121}, Grit Schubert¹⁰⁶, Gugu P. Maphalala¹²², Hafaliana C. Ranaivoson⁷⁷, Hannah E. Omunakwe¹²³, Harris Onywera⁷, Haruka Abe¹²⁴, Hela Karray¹²⁵, Hellen Nansumba¹²⁶, Henda Triki⁵⁸, Herve Albéric Adje Kadjo¹²⁷, Hesham Elgahzaly¹²⁸, Hlanai Gumbo³⁹, Hota Mathieu¹²⁹, Hugo Kavunga-Membo³⁷, Ibtihel Smeti⁴¹, Idowu B. Olawoye⁴³, Ifedayo M. O. Adetifa^{88,130}, Ikpomwosa Odia⁹², Ilhem Boutiba-Ben Boubaker^{131,132}
Muhammad I. Ahmed⁴³, Isaac Ssewanyana¹²⁶, Isatta Wurie¹³³, Iyaloo S. Konstantinus¹³⁴, Jacqueline Wemboo Afiwa Halatoko^{1,35}, James Ayei²⁶, Janaki Sonoo¹³⁶, Jean-Claude C. Makangara^{37,38}, Jean-Jacques M. Tamfum^{37,38}, Jean-Michel Heraud^{12,77}, Jean-Jacques M. Tamfum^{37,38}, Jean-Michel Heraud^{12,77},
Jeffrey G. Shaffer¹³⁷, Jennifer Giandhari², Jennifer Musyoki¹¹,
Jerome Nkurunziza¹³⁸, Jessica N. Uwanibe⁴³,
Jinal N. Bhiman^{13,113}, Jiro Yasuda¹²⁴, Joana Morais^{139,140},
Jocelyn Kiconco⁹⁷, John D. Sandi^{64,65}, John Huddleston¹⁴¹,
John K. Odoom⁷⁴, John M. Morobe¹¹, John O. Gyapong¹²⁰,
John T. Kayiwa³, Johnson C. Okolie⁴³, Joicymara S. Xavier^{1,142,143},
Jones Gyamfi¹²⁰, Joseph F. Wamala¹⁴⁴, Joseph H. K. Bonney⁷⁴,
Joseph Nyandw^{151,145}, Josie Everatt¹³, Joweria Nakaseegu⁹⁷,
Joyce M. Ngoi⁹¹, Joyce Namulondo⁹⁷, Judith U. Oguzie^{83,44},
Julia C. Andeko⁶⁸, Julius J. Lutwama³, Juma J. H. Mogga¹⁴⁴,
Justin O'Grady⁴⁸, Katherine J. Siddle⁵⁷, Kathleen Victoir¹⁴⁶,
Kayode T. Adeyemi^{43,44}, Kefentse A. Tumedi¹⁴⁷,
Kevin S. Carvalho¹⁴⁸, Khadiia Said Mohammed¹¹ Kevin S. Carvalho¹⁴⁸, Khadija Said Mohammed¹¹, Koussay Dellagi¹⁴⁶, Kunda G. Musonda¹⁴⁹, Kwabena O. Duedu^{120,121}, Lamia Fki-Berrajah¹²⁵, Lavanya Singh², Lenora M. Kepler^{94,95}, Leon Biscornet⁷⁵, Leonardo de Oliveira Martins⁴⁸, Lucious Chabuka¹⁵⁰, Luicer Olubayo⁸, Lul Deng Ojok²⁶, Lul Lojok Deng² Luncur Ouwayor, Lui Deng Ojok^{co}, Lul Lojok Deng²⁶, Lynette I. Ochola-Oyier¹¹, Lynn Tyers⁹, Madisa Mine¹⁵¹, Magalutcheemee Ramuth¹³⁶, Maha Mastouri^{152,153}, Mahmoud ElHefnawi¹⁵⁴, Maimouna Mbanne¹², Maitshwarelo I. Matsheka¹⁴⁷, Malebogo Kebabonye¹⁵⁵, Mamadou Diop¹², Mambu Momoh^{64,65,156} Maria da Luz Lima Mendonça¹⁴⁸, Marietjie Venter¹⁵⁷, Maria da Luz Lima Mendonça¹⁴⁸, Marietjie Venter¹⁵⁷,
Marietou F. Paye⁵⁷, Martin Faye¹², Martin M. Nyaga¹⁵⁸,
Mathabo Mareka¹⁵⁹, Matoke-Muhia Damaris¹⁶⁰,
Maureen W. Mburu¹¹, Maximillian G. Mpina^{161,162,163},
Michael Owusu¹⁶⁴, Michael R. Wiley^{81,165}, Mirabeau Y. Tatfeng¹⁶⁶,
Mitoha Ondo'o Ayekaba¹⁶², Mohamed Abouelhoda^{167,168},
Mohamed Amine Beloufa¹¹¹, Mohamed G. Seadawy^{169,170},
Mohamed K. Khalifa¹⁷¹, Mooko Marethabile Matobo¹⁵⁹,
Muhamed K. Sane¹², Mungerou Salou¹⁷², Mohamis B. Mbulawa¹⁵⁵ Mouhamed Kane¹², Mounerou Salou¹⁷², Mphaphi B. Mbulawa¹⁵⁵, Mulenga Mwenda⁹³, Mushal Allam¹⁷³, My V. T. Phan³, Nabil Abid^{152,174}, Nadine Rujeni^{175,176}, Nadir Abuzaid¹⁷⁷. Nalia Ismael¹⁷⁸, Nancy Elguindy⁵⁴, Ndeye Marieme Top¹², Ndongo Dia¹², Nédio Mabunda¹⁷⁸, Nei-yuan Hsiao^{9,78}, Nelson Boricó Silochi¹⁶², Ngiambudulu M. Francisco¹³⁹, Ngonda Saasa¹⁷⁹, Nicholas Bbosa³, Nickson Murunga¹¹, Nicksy Gumede¹⁸, Nicole Wolter^{13,113}, Nikita Sitharam¹, Nnaemeka Ndodo⁸⁸, Nnennaya A. Ajayi¹⁸⁰, Noël Tordo¹⁸¹,

Nokuzola Mbhele9, Norosoa H. Razanajatovo77, Nosamiefan Iguosadolo⁴³, Nwando Mba⁸⁸, Ojide C. Kingsley¹⁸², Okogbenin Sylvanus⁹², Oladiji Femi¹⁸³, Olubusuyi M. Adewumi^{31,32}, Olumade Testimony^{43,44}, Olusola A. Ogunsanya⁴³, Oluwatosin Fakayode¹⁸⁴, Olwe E. Ogah¹⁸⁵, Ope-Ewe Oludayo⁴³, Ousmane Faye¹², Pamela Smith-Lawrence¹⁵⁵, Pascale Ondoa⁷¹, Patrice Combe¹⁸⁶, Patricia Nabisubi¹⁸⁷, Patrick Semanda¹²⁶, Paul E. Oluniyi⁴³, Paulo Arnaldo¹⁷⁸, Peter Kojo Quashie⁹¹, Peter O. Okokhere^{92,189}, Paulo Arnaldo¹⁷⁸, Peter Kojo Quashie⁹¹, Peter O. Okokhere^{92,189}, Paulo Arnaldo¹⁷⁸, Peter Kojo Quashie⁹¹, Peter O. Okokhere^{92,189}, Patrick Paulo Arnaldo¹⁷⁸, Peter No. Okokhere^{92,189}, Patrick Paulo P Philip Bejon¹¹, Philippe Dussart⁷⁷, Phillip A. Bester¹⁹⁰, Placide K. Mbala^{37,38}, Pontiano Kaleebu^{3,97}, Priscilla Abechi^{43,44}, Rabeh El-Shesheny^{40,19}, Rageema Joseph⁹, Ramy Karam Aziz^{192,193}, René G. Essomba^{194,195}, Reuben Ayivor-Djanie^{91,120,121}, Richard Njouom¹⁹⁶, Reuben Aywor-Ujanie^{5,13} Richard Njouomis⁵, Richard O. Phillips⁶³, Richmond Gorman⁶³, Robert A. Kingsley⁴⁸, Rosa Maria D. E. S. A. Neto Rodrigues^{197,138}
Rosemary A. Audu²⁹, Rosina A. A. Carr^{120,121}, Saba Gargouri¹²⁵, Saber Masmoudi⁴¹, Sacha Bootsma¹⁴⁴, Safietou Sankhe¹², Sahra Isse Mohamed¹⁹⁹, Saibu Femi⁴³, Salma Mhalla^{152,200}, Salome Hosch^{161,201}, Samar Kamal Kassim¹²⁸, Samar Metha⁵⁷, Sameh Trabelsi²⁰², Sarah Massan Agwa¹²⁸, Sarah Wambui Mwanqi⁷, Sarah Doumbia⁵² Sarah Wambui Mwangi⁷, Seydou Doumbia⁵², Sheila Makiala-Mandanda^{27,38}, Sherihane Aryeetey⁶³, Shehial Makala-Mahalada Janua Janua Janua Hayeetey , Shymaa S. Ahmed⁵⁴ Side Mohamed Ahmed¹⁰³, Siham Elhamoumi⁵⁷, Sikhulile Moyo^{100,203}, Silvia Lutucuta¹³⁹, Simani Gaseitsiwe^{100,203}, Simbirie Jalloh^{64,65}, Soa Fy Andriamandimby⁷⁷, Sobajo Oguntoppe⁴³, Solène Grayo¹⁸¹, Soa i y Andriamandimby", Sobajo Uguntope ", Solene Grayo". Sonia Lekana-Douki⁶⁸, Sophie Prosolek⁴⁸, Soumeya Ouangraou^{204,205}, Stephanie van Wyk¹, Stephen F. Schaffner⁵⁷, Stephen Kanyerezi^{187,188}, Steve Ahuka-Mundeke^{37,38}, Steven Rudder⁴⁸, Sureshnee Pillay², Susan Nabadda¹²⁶, Sylvie Behillil²⁰⁶, Sylvie L. Budiaki¹⁵⁹, Sylvie van der Werf²⁰⁶, Tapfumanei Mashe^{39,207}, Thabo Mohale¹³ Thanh Le-Viet⁴⁸, Thirumalaisamy P. Velavani^{14,208}, Tobias Schindler^{16,1,62,201}, Tongai G. Maponga¹¹⁸, Trevor Bedford^{14,2,09}, Ugochukwu J. Anyaneji², Ugwu Chinedu^{43,44}, Upasana Ramphal^{2,10,210}, Uwem E. George⁴³, Vincent Enour²⁰⁶, Vishvanath Nene¹⁰², Vivianne Gorova^{211,212} Wael H. Roshdy⁵⁴, Wasim Abdul Karim¹, William K. Ampofo²¹³, Wolfgang Preiser^{118,119}, Wonderful T. Choga^{100,214}, Yahaya Ali Ahmed¹⁸, Yajna Ramphal¹, Yaw Bediako^{91,215}, Yeshnee Naidoo², Yvan Butera^{175,216,217}, Zaydah R. de Laurent¹¹, Africa Pathogen Genomics Initiative (Africa PGI)‡, Ahmed E. O. Ouma⁷, Anne von Gottberg^{13,113},
George Githinij^{11,218}, Matshidiso Moeti¹⁸, Oyewale Tomori⁴³,
Pardis C. Sabeti⁵⁷, Amadou A. Sall¹², Samuel O. Oyola¹⁰²,
Yenew K. Tebeje⁷, Sofonias K. Tessema⁷,
Tulio de Oliveira^{12,10,219}*, Christian Happi^{43,44}, Richard Lessells², John Nkengasong⁷, Eduan Wilkinson^{1,2}

¹Centre for Epidemic Response and Innovation (CERI), School of Data Science and Computational Thinking, Stellenbosch University, Stellenbosch, South Africa. 2KwaZulu-Natal Research Innovation and Sequencing Platform (KRISP), Nelson R Mandela School of Medicine, University of KwaZulu-Natal, Durban, South Africa. 3MRC/ UVRI and LSHTM Uganda Research Unit, Entebbe, Uganda. ⁴MRC-University of Glasgow Centre for Virus Research, Glasgow, UK. ⁵The Biotechnology Centre of the University of Yaoundé I, Yaoundé, Cameroon. ⁶CDC Foundation, Atlanta, Georgia, Nebraska Department of Health and Human Services, Lincoln, NE, USA. ⁷Institute of Pathogen Genomics. Africa Centres for Disease Control and Prevention (Africa CDC), Addis Ababa, Ethiopia. 8Institute of Infectious Diseases and Molecular Medicine, Department of Integrative Biomedical Sciences, Computational Biology Division, University of Cape Town, Cape Town, South Africa. 9Division of Medical Virology, Wellcome Centre for Infectious Diseases in Africa, Institute of Infectious Disease and Molecular Medicine, University of Cape Town, Cape Town, South Africa. OCentre for the AIDS Programme of Research in South Africa (CAPRISA), Durban, South Africa. 11KEMRI-Wellcome Trust Research Programme, Kilifi, Kenya. ¹²Institut Pasteur de Dakar, Dakar, Senegal. ¹³National Institute for Communicable Diseases (NICD) of the National Health Laboratory Service (NHLS), Johannesburg, South Africa. ¹⁴School of Health Sciences, College of Health Sciences, University of KwaZulu-Natal, Durban, KwaZulu-Natal, South Africa. ¹⁵Department of Medical Laboratory Sciences, College of Health Sciences, Addis Ababa University, Addis Ababa, Ethiopia. 16 Department of Microbial, Cellular and Molecular Biology, College of Natural and Computational Sciences, Addis Ababa University, Addis

Ababa, Ethiopia. ¹⁷Cancer Biology Department, Virology and Immunology Unit, National Cancer Institute, Cairo University, Cairo, Egypt. 18 World Health Organization, Africa Region, Brazzaville, Republic of the Congo. ¹⁹Centre d'Infectiologie Charles Mérieux-Mali (CICM-Mali), Bamako, Mali. ²⁰Bacteriology and Virology Department Souro Sanou University Hospital, Bobo-Dioulasso, Burkina Faso. ²¹West African Health Organisation, Bobo-Dioulasso, Burkina Faso. ²²Faculty of Medicine and Health Sciences, Kassala University, Kassala City, Sudan. ²³Department of Microbiology, Faculty of Medical Laboratory Sciences, University of Gezira, Gezira, Sudan. ²⁴General Administration of Laboratories and Blood Banks, Ministry of Health, Kassala State, Sudan. 25MRC Unit The Gambia at LSHTM, Fajara, Gambia, 26National Public Health Laboratory, Ministry of Health, Juba, Republic of South Sudan. 27 Libyan Biotechnology Research Center, Tripoli, Libya. ²⁸Center for Medical and Sanitary Research (CERMES), Niamey, Niger. ²⁹The Nigerian Institute of Medical Research, Yaba, Lagos, Nigeria. 30 Laboratoire de la Caisse Nationale de Sécurité Sociale, Djibouti, Republic of Djibouti. 31Department of Virology, College of Medicine, Úniversity of Ibadan, Ibadan, Nigeria. ³²Infectious Disease Institute, College of Medicine, University of Ibadan, Ibadan, Nigeria. ³³Medical Microbiology and Parasitology Department, College of Medicine, University of Ibadan, Ibadan, Nigeria. 34Biorepository Clinical Virology Laboratory, College of Medicine, University of Ibadan, Ibadan, Nigeria. ³⁵Department of Medical Microbiology and Parasitology, Faculty of Basic Clinical Sciences, College of Health Sciences, University of Ilorin, Ilorin, Kwara State, Nigeria. ³⁶The Pirbright Institute, Woking, UK. ³⁷Pathogen Seguencing Lab. Institut National de Recherche Biomédicale (INRB), Kinshasa, the Democratic Republic of the Congo. ³⁸Université de Kinshasa (UNIKIN), Kinshasa, the Democratic Republic of the Congo. ³⁹National Microbiology Reference Laboratory, Harare, Zimbabwe. 40Center of Scientific Excellence for Influenza Viruses, National Research Centre (NRC), Cairo, Egypt. ⁴¹Laboratory of Molecular and Cellular Screening Processes, Centre of Biotechnology of Sfax, University of Sfax, Sfax, Tunisia. ⁴²Genomics and Epigenomics Program, Research Department CCHE57357, Cairo, Egypt. 43 African Centre of Excellence for Genomics of Infectious Diseases (ACEGID), Redeemer's University, Ede, Osun State, Nigeria. ⁴⁴Department of Biological Sciences, Faculty of Natural Sciences, Redeemer's University, Ede, Osun State, Nigeria. ⁴⁵Laboratório de Biologia Molecular Jean Piaget, Bissau, Guinea-Bissau. 46 University Jean Piaget in Guinea-Bissau, Bissau, Guinea-Bissau. 47SAMRC Bioinformatics Unit, SA Bioinformatics Institute, University of the Western Cape, Cape Town, South Africa. ⁴⁸Quadram Institute Bioscience, Norwich, UK. ⁴⁹Central Public Health Reference Laboratories, Freetown, Sierra Leone. 50 Centre de Recherche et de Formation en Infectiologie de Guinée (CERFIG), Université de Conakry, Conakry, Guinea. ⁵¹TransVIHMI, Institut de Recherche pour le Développement, Institut National de la Santé et de la Recherche Médicale (INSERM), Montpellier University, 34090, Montpellier, France. ⁵²University Clinical Research Center (UCRC), University of Sciences, Techniques and Technology of Bamako, Bamako, Mali. 53 Sharjah Institute for Medical Research, College of Medicine, University of Sharjah, Sharjah, United Arab Emirates. ⁵⁴Central Public Health Laboratories (CPHL), Cairo, Egypt. 55 National Institute of Public Health, Bujumbura, Burundi. ⁵⁶Laboratoire des Fièvres Hémorragiques Virales du Benin, Cotonou, Benin. ⁵⁷Infectious Disease and Microbiome Program, Broad Institute of Harvard and MIT, Cambridge, MA, USA. ⁵⁸Laboratory of Clinical Virology, WHO Reference Laboratory for Poliomyelitis and Measles in the Eastern Mediterranean Region, Pasteur Institute of Tunis, University Tunis El Manar (UTM), Tunis 1002, Tunisia. 59 Research Laboratory "Virus, Vectors and Hosts: One Health Apporach and Technological Innovation for a Better Health", LR20IPT02, Pasteur Institute, Tunis 1002, Tunisia.

60 Fondation Congolaise pour la Recherche Médicale, Brazzaville, Republic of the Congo. 61Marien Ngouabi, Brazzaville, Republic of the Congo. ⁶²Kwame Nkrumah University of Science and Technology, Department of Theoretical and Applied Biology, Kumasi, Ghana. ⁶³Kumasi Centre for Collaborative Research in Tropical Medicine, Kwame Nkrumah University of Science and Technology, Kumasi, Ghana. 64Viral Haemorrhagic Fever Laboratory, Kenema Government Hospital, Kenema, Sierra Leone. ⁶⁵Ministry of Health and Sanitation, Freetown, Sierra Leone.

⁶⁶Department of Immunology, University of Maiduguri Teaching Hospital, P.M.B. 1414, Maiduguri, Nigeria. ⁶⁷Department of Medical Laboratory Science, College of Medical Sciences, University of Maiduguri, P.M.B. 1069, Maiduguri, Borno State, Nigeria. ⁶⁸Centre Interdisciplinaires de Recherches Medicales de Franceville (CIRMF), Franceville, Gabon. ⁶⁹Département de Parasitologie-Mycologie Université des Sciences de la Santé (USS), Libreville, Gabon. 70 National HIV Reference Laboratory, Community Health Sciences Unit, Ministry of Health, Lilongwe, Malawi. ⁷¹African Society for Laboratory Medicine, Addis Ababa, Ethiopia. 72National Medical and Molecular Biology Laboratory, Ministry of Health, Djibouti, Republic of Djibouti. ⁷³Africa CDC, Rapid Responder, Team Djibouti, Djibouti, Djibouti. 74Noguchi Memorial Institute for Medical Research, University of Ghana, Legon, Ghana. 75 Seychelles Public Health Laboratory, Public Health Authority, Ministry of Health Seychelles, Victoria, Seychelles. ⁷⁶Department of Ecology and Evolution, University of Chicago, Chicago, IL, USA. ⁷⁷Virology Unit, Institut Pasteur de Madagascar, Antananarivo, Madagascar. 78 National Health Laboratory Service (NHLS), Cape Town, South Africa. ⁷⁹Malawi-Liverpool-Wellcome Trust Clinical Research Programme, Blantyre, Malawi. ⁸⁰Liverpool School of Tropical Medicine, Liverpool, UK. ⁸¹University of Nebraska Medical Center (UNMC), Omaha, NE, USA. ⁸²SAMRC Antibody Immunity Research Unit, School of Pathology, University of the Witwatersrand, Johannesburg, South Africa. 83CHU de Bouaké, Laboratoire/Unité de Diagnostic des Virus des Fièvres Hémorragiques et Virus Émergents, Bouaké, Côte d'Ivoire. 84UFR Sciences Médicales, Universite Alassane Ouattara, Bouaké, Côte d'Ivoire. 85School of Public Health, Pwani University, Kilifi, Kenya. ⁸⁶Faculty of Science and Techniques, University Marien Ngouabi, Brazzaville, Republic of the Congo. 87Centre for Human Virology and Genomics, Nigerian Institute of Medical Research, Yaba, Lagos, Nigeria. 88 Nigeria Centre for Disease Control and Prevention, Abuja, Nigeria. 89 Laboratoire des Arbovirus, Fièvres Hémorragiques virales, Virus Emergents et Zoonoses, Institut Pasteur de Bangui, Bangui, Central African Republic. 90Le Laboratoire National de Biologie Clinique et de Santé Publique (LNBCSP), Bangui, Central African Republic. 91West African Centre for Cell Biology of Infectious Pathogens (WACCBIP), College of Basic and Applied Sciences University of Ghana Accra, Ghana. 92 Institute of Lassa Fever Research and Control, Irrua Specialist Teaching Hospital, Irrua, Nigeria. ⁹³PATH, Lusaka, Zambia. ⁹⁴Department of Entomology and Plant Pathology, North Carolina State University, Raleigh, NC, USA. ⁹⁵Bioinformatics Research Center, North Carolina State University, Raleigh, NC, USA. ⁹⁶School of Life Sciences and Zeeman Institute for Systems Biology and Infectious Disease Epidemiology Research (SBIDER), University of Warwick, Coventry, UK. ⁹⁷Uganda Virus Research Institute, Entebbe, Uganda. 98PathCare Vermaak, Pretoria, South Africa and Division of Virology, University of the Free State, Bloemfontein, South Africa. ⁹⁹College of Medicine and Allied Health Sciences, University of Sierra Leone, Freetown, Sierra Leone. ¹⁰⁰Botswana Harvard AIDS Institute Partnership and Botswana Harvard HIV Reference Laboratory, Gaborone, Botswana. 101 Macha Research Trust, Choma, Zambia. 102 International Livestock Research Institute (ILRI), Nairobi, Kenya. 103 INRSP, Nouakchott, Mauritania. 104 Faculté de Médecine de Nouakchott, Nouakchott, Mauritani. 105 Rwanda National Reference Laboratory, Kigali, Rwanda. ¹⁰⁶Robert Koch-Institute, Berlin, Germany. ¹⁰⁷G5 Evolutionary Genomics of RNA Viruses, Institut Pasteur, Paris, France. ¹⁰⁸Direcção Nacional da Saúde Pública, Ministério da Saúde, Luanda, Angola. 109 National Public Health Reference Laboratory National Public Health Institute of Liberia, Monrovia, Liberia.

110 Faculty of Pharmacy of Monastir, Monastir, Tunisia. ¹¹¹National Influenza Centre, Institut Pasteur d'Algérie, Algiers, Algeria. 112 Department of Virology, National Health Laboratory Service (NHLS), Charlotte Maxeke Johannesburg Academic Hospital, Johannesburg, South Africa. 113 School of Pathology, Faculty of Health Science, University of the Witwatersrand, Johannesburg, South Africa. 114 Institute of Tropical Medicine, Universitätsklinikum Tübingen, Tübingen, Germany, 115 Ministère de Santé Publique et de la Solidarité Nationale, Ndjamena, Chad. ¹¹⁶WHO Int Comoros, Moroni, Union of Comoros. ¹¹⁷World Health Organization, Africa Region, Brazzaville, Republic of the Congo. ¹¹⁸Division of Medical Virology, Faculty of Medicine and Health Sciences, Stellenbosch University, Tygerberg, Cape Town, South Africa. 119 National Health Laboratory Service (NHLS), Tygerberg,

Cape Town, South Africa. 120 UHAS COVID-19 Testing and Research Centre, University of Health and Allied Sciences, Ho, Ghana. 121 Department of Biomedical Sciences, University of Health and Allied Sciences, PMB 31, Ho, Ghana. ¹²²Ministry of Health, COVID-19 Testing Laboratory, Mbabane, Kingdom of Eswatini. 123 Satellite Molecular Laboratory, Rivers State University Teaching Hospital, Port Harcourt, Nigeria. ¹²⁴Department of Emerging Infectious Diseases, Institute of Tropical Medicine, Nagasaki University. Nagasaki, Japan. 125CHU Habib Bourguiba, Laboratory of Microbiology, Faculty of Medicine of Sfax, University of Sfax, Sfax, Tunisia. ¹²⁶Central Public Health Laboratories (CPHL), Kampala, Uganda. ¹²⁷Institut Pasteur de Côte d'Ivoire, Departement des Virus Epidemiques, Abidjan, Côte d'Ivoire. 128 Faculty of Medicine Ain Shams Research Institute (MASRI), Ain Shams University, Cairo, Egypt. 129 Doctoral School of Technical and Environmental Sciences, Department of Biology and Human Health, N'Djamena, Chad. ¹³⁰Department of Infectious Diseases Epidemiology, London School of Hygiene and Tropical Medicine, London, UK. ¹³¹Charles Nicolle Hospital, Laboratory of Microbiology, National Influenza Center, Tunis, Tunisia. 132 University of Tunis El Manar, Faculty of Medicine of Tunis, Research Laboratory LR99ES09, Tunis, Tunisia. ¹³³College of Medicine and Allied Health Science, University of Sierra Leone, Freetown, Sierra Leone. ¹³⁴Namibia Institute of Pathology, Windhoek, Namibia. ¹³⁵National Institute of Hygiene, Lomé, Togo. ¹³⁶Virology/Molecular Biology Department, Central Health Laboratory, Victoria Hospital, Ministry of Health and Wellness, Port Louis, Mauritius. ¹³⁷Department of Biostatistics and Data Science, School of Public Health and Tropical Medicine, Tulane University, New Orleans, LA, USA. ¹³⁸WHO Burundi, Gitega, Burundi. ¹³⁹Grupo de Investigação Microbiana e Imunológica, Instituto Nacional de Investigação em Saúde (National Institute for Health Research), Luanda, Angola. 140 Departamento de Bioquímica, Faculdade de Medicina, Universidade Agostinho Neto, Luanda, Angola. ¹⁴¹Vaccine and Infectious Disease Division, Fred Hutchinson Cancer Center, Seattle, WA, USA. ¹⁴²Universidade Federal de Minas Gerais, Belo Horizonte, Brazil. ¹⁴³Institute of Agricultural Sciences, Universidade Federal dos Vales do Jeguitinhonha e Mucuri, Unaí, Brazil. 144WHO South Sudan, Juba, South Sudan. 145 Faculty of Medicine, University of Burundi, Bujumbura, Burundi. ¹⁴⁶Pasteur Network, Institut Pasteur, Paris, France. ¹⁴⁷Botswana Institute for Technology Research and Innovation, Gaborone, Botswana. ¹⁴⁸Instituto Nacional de Saúde Pública, Praia, Cape Verde. ¹⁴⁹Zambia National Public Health Institute, Lusaka, Zambia. 150 Public Health Institute of Malawi, Lilongwe, Malawi. 151 National Health Laboratory, Gaborone, Botswana. ¹⁵²Laboratory of Transmissible Diseases and Biologically Active Substances (LR99ES27), Faculty of Pharmacy, University of Monastir, Monastir, Tunisia. ¹⁵³Laboratory of Microbiology, University Hospital of Monastir, Monastir, Tunisia. ¹⁵⁴Biomedical Informatics and Chemoinformatics Group, Informatics and Systems Department, National Research Centre, Cairo, Egypt. ¹⁵⁵Ministry of Health and Wellness, Gaborone, Botswana. ¹⁵⁶Eastern Technical University of Sierra Leone, Kenema, Sierra Leone. ¹⁵⁷Zoonotic Arbo and Respiratory Virus Program, Centre for Viral Zoonoses, Department of Medical Virology, University of Pretoria, Pretoria, South Africa. ¹⁵⁸Next Generation Sequencing Unit and Division of Virology, Faculty of Health Sciences, University of the Free State, Bloemfontein, South Africa. 159 National Reference Laboratory Lesotho, Maseru, Lesotho. 160 Centre for Biotechnology Research and Development, Kenya Medical Research Institute, Nairobi, Kenya. ¹⁶¹Swiss Tropical and Public Health Institute, Basel, Switzerland. ¹⁶²Laboratorio de Investigaciones de Baney, Baney, Equatorial Guinea. ¹⁶³Ifakara Health Insitute, Ifakara, Tanzania. ¹⁶⁴Department of Medical Diagnostics, Kumasi Centre for Collaborative Research in Tropical Medicine, Kwame Nkrumah University of Science and Technology, Kumasi, Ghana. ¹⁶⁵PraesensBio, Lincoln, NE, USA. ¹⁶⁶Department of Medical Laboratory Science, Niger Delta University, Bayelsa State, Nigeria. 167 Systems and Biomedical Engineering Department, Faculty of Engineering, Cairo University, Cairo, Egypt. 168 King Faisal Specialist Hospital and Research Center, Riyadh, Kingdom of Saudi Arabia. ¹⁶⁹Biological Prevention Department, Ministry of Defence, Cairo, Egypt. ¹⁷⁰Faculty of Science, Fayoum University, Fayoum, Egypt. 171 Molecular Pathology Lab, Children's Cancer Hospital, Cairo, Egypt. 172 Laboratoire Biolim FSS/Université de Lomé, Lomé, Togo. 173 Department

of Genetics and Genomics, College of Medicine and Health Sciences, United Arab Emirates University, Abu Dhabi, United Arab Emirates. 174High Institute of Biotechnology of Monastir, University of Monastir, Rue Taher Haddad 5000, Monastir, Tunisia. ¹⁷⁵Rwanda National Joint Task Force COVID-19, Rwanda Biomedical Centre, Ministry of Health, Kigali, Rwanda. 176School of Health Sciences, College of Medicine and Health Sciences, University of Rwanda, Kigali, Rwanda. ¹⁷⁷Department of Microbiology, Faculty of Medical Laboratory Sciences, Omdurman Islamic University, Sudan. 178 Instituto Nacional de Saúde (INS), Marracuene, Mozambique. ¹⁷⁹Department of Disease Control, School of Veterinary Medicine, University of Zambia, Lusaka, Zambia. 180 Internal Medicine Department, Alex Ekwueme Federal University Teaching Hospital, Abakaliki, Nigeria. ¹⁸¹Institut Pasteur de Guinée, Conarky, Guinea. ¹⁸²Virology Laboratory, Alex Ekwueme Federal University Teaching Hospital, Abakaliki, Nigeria. ¹⁸³Department of Epidemiology and Community Health, Faculty of Clinical Sciences. College of Health Sciences. University of Ilorin, Ilorin, Kwara State, Nigeria. ¹⁸⁴Department of Public Health, Ministry of Health, Ilorin, Kwara State, Nigeria. ¹⁸⁵Alex Ekwueme Federal University Teaching Hospital, Abakaliki, Nigeria. ¹⁸⁶Mayotte Hospital Center, Mayotte, France. 187The African Center of Excellence in Bioinformatics and Data-Intensive Sciences, The Infectious Diseases Institute, Kampala, Uganda. 188 Immunology and Molecular Biology, Makerere University, Kampala, Uganda. ¹⁸⁹Department of Medicine, Faculty of Clinical Sciences, College of Medicine, Ambrose Alli University, Ekpoma, Edo State, Nigeria. ¹⁹⁰Division of Virology, National Health Laboratory Service and University of the Free State,

Bloemfontein, South Africa. 191 Infectious Hazards Preparedness, World Health Organization, Eastern Mediterranean Regional Office, Cairo, Egypt. 192 Department of Microbiology and Immunology, Faculty of Pharmacy, Cairo University, Cairo, Egypt. ¹⁹³Microbiology and Immunology Research Program, Children's Cancer Hospital Egypt, Cairo, Egypt. ¹⁹⁴National Public Health Laboratory, Ministry of Public Health of Cameroon, Yaoundé, Cameroon. ¹⁹⁵Facu of Medicine and Biomedical Sciences. University of Yaoundé. Yaoundé, Cameroon. ¹⁹⁶Virology Service, Centre Pasteur of Cameroun, Yaounde, Cameroon. ¹⁹⁷Coordenadora da rede do Diagnóstico Tuberculose/HIV/COVID-19 na Instituição -Laboratório Nacional de Referência da Tuberculose em São Tomé e Príncipe, São Tomé, São Tomé and Principe. ⁸Ponto focal para Melhoria da qualidade dos Laboratórios (SLIPTA) ao nível de São Tomé e Príncipe, São Tomé, São Tomé and Principe. ¹⁹⁹National Public Health Reference Laboratory (NPHRL), Mogadishu, Somalia. ²⁰⁰Faculty of Medicine of Monastir, University of Monastir, Monastir, Tunisia. 201 University of Basel, Basel, Switzerland. 202 Clinical and Experimental Pharmacology Lab, LR16SP02, National Center of Pharmacovigilance, University of Tunis El Manar, Tunis, Tunisia. ²⁰³Harvard T.H. Chan School of Public Health, Boston, MA, USA. ²⁰⁴Centre MURAZ, Ouagadougou, Burkina Faso. ²⁰⁵National Institute of Public Health of Burkina Faso (INSP/BF), Ouagadougou, Burkina Faso. 206National Reference Center for Respiratory Viruses, Molecular Genetics of RNA Viruses, UMR 3569 CNRS, Université Paris Cité, Institut Pasteur, Paris, France. ²⁰⁷World Health Organization, Harare, Zimbabwe. ²⁰⁸Vietnamese-German Center for Medical Research, Hanoi, Vietnam. 209 Howard Hughes Medical Institute, Fred

Hutchinson Cancer Center, Seattle, WA, USA. $^{\rm 210}{\rm Sub}\text{-Saharan}$ African Network For TB/HIV Research Excellence (SANTHE), Durban, South Africa. 211 World Health Organization, WHO Lesotho, Maseru, Lesotho. 212 Med24 Medical Centre, Ruwa, Zimbabwe. ²¹³Department of Virology, Noguchi Memorial Institute for Medical Research, University of Ghana, Legon, Ghana. ²¹⁴Division of Human Genetics, Department of Pathology, University of Cape Town, Cape Town, South Africa. 215 Yemaachi Biotech, Accra, Ghana, 216 Center for Human Genetics, College of Medicine and Health Sciences, University of Rwanda, Kigali, Rwanda. ²¹⁷Laboratory of Human Genetics, GIGA Research Institute, Liège, Belgium. ²¹⁸Department of Biochemistry and Biotechnology, Pwani University, Kilifi, Kenya. 219 Department of Global Health, University of Washington, Seattle, WA, USA. *Corresponding author. Email: tulio@sun.ac.za (T.d.O.); ewilkinson@sun.ac.za (E.W.)

†These authors contributed equally to this work. ‡Africa PGI collaborators are listed in the supplementary materials.

SUPPLEMENTARY MATERIALS

science.org/doi/10.1126/science.abq5358 Africa PGI Collaborator List Figs. S1 to S16 Tables S1 to S4 Reference (77) MDAR Reproducibility Checklist

View/request a protocol for this paper from Bio-protocol.

Submitted 14 April 2022; accepted 12 September 2022 10.1126/science.abq5358



The evolving SARS-CoV-2 epidemic in Africa: Insights from rapidly expanding genomic surveillance

Houriiyah TegallyJames E. SanMatthew CottenMonika MoirBryan TegomohGerald MboowaDarren P. MartinCheryl BaxterArnold W. LambisiaAmadou DialloDaniel G. AmoakoMoussa M. DiagneAbay SisayAbdel-Rahman N. ZekriAbdou Salam GueyeAbdoul K. SangareAbdoul-Salam OuedraogoAbdourahmane SowAbdualmoniem O. MusaAbdul K. SesayAbe G. AbiasAdam I. ElzagheidAdamou LagareAdedotun-Sulaiman KemiAden Elmi AbarAdeniji A. JohnsonAdeola FowotadeAdeyemi O. OluwapelumiAdrienne A. AmuriAgnes JuruAhmed KandeilAhmed MostafaAhmed RebaiAhmed SayedAkano KazeemAladje BaldeAlan ChristoffelsAlexander J. TrotterAllan CampbellAlpha K. KeitaAmadou KoneAmal BouzidAmal SouissiAmbrose AgweyuAmel NaguibAna V. GutierrezAnatole NkeshimanaAndrew J. PageAnges YadouletonAnika VinzeAnise N. HappiAnissa ChouikhaArash IranzadehArisha MaharajArmel L. Batchi-BouyouArshad IsmailAugustina A. SylverkenAugustine GobaAvoade FemiAvotunde E. SiiuwolaBaba MarycelinBabatunde L. SalakoBamidele S. OderindeBankole BolajokoBassirou DiarraBelinda L. HerringBenjamin TsofaBernard Lekana-DoukiBernard MvulaBerthe-Marie Njanpop-LafourcadeBlessing T. MaronderaBouh Abdi KhairehBourema KouribaBright AduBrigitte PoolBronwyn McInnisCara BrookCarolyn WilliamsonCassien NduwimanaCatherine AnscombeCatherine B. PrattCathrine ScheepersChantal G. Akoua-KoffiCharles N. AgotiChastel M. MapanguyCheikh LoucoubarChika K. OnwuamahChikwe IhekweazuChristian N. MalakaChristophe PeyrefitteChukwa GraceChukwuma E. OmoruyiClotaire D. RafaïCollins M. Morang'aCyril EramehDaniel B. LuleDaniel J. BridgesDaniel Mukadi-BamulekaDanny ParkDavid A. RasmussenDavid BakerDavid J. NokesDeogratius SsemwangaDerek TshiabuilaDominic S. Y. AmuzuDominique GoedhalsDonald S. GrantDonwilliams O. OmuoyoDorcas MaruapulaDorcas W. WanjohiEbenezer Foster-NyarkoEddy K. LusamakiEdgar SimulunduEdidah M. Ong'eraEdith N. NgabanaEdward O. AbworoEdward OtienoEdwin ShumbaEdwine BarasaEl Bara AhmedElhadi A. AhmedEmmanuel LokiloEnatha MukantwariEromon PhilomenaEssia BelarbiEtienne Simon-LoriereEtilé A. AnohEusebio ManuelFabian LeendertzFahn M. TawehFares WasfiFatma AbdelmoulaFaustinos T. TakawiraFawzi DerrarFehintola V. AjogbasileFlorette TreurnichtFolarin OnikepeFrancine NtoumiFrancisca M. MuyembeFrank E. Z. RagomzingbaFred A. DratibiFred-Akintunwa IyanuGabriel K. MbunsuGaetan ThilliezGemma L. KayGeorge O. AkpedeGert U. van ZylGordon A. AwandareGrace S. KpeliGrit SchubertGugu P. MaphalalaHafaliana C. RanaivosonHannah E. OmunakweHarris OnyweraHaruka AbeHela KarrayHellen NansumbaHenda TrikiHerve Albéric Adje KadjoHesham ElgahzalyHlanai GumboHota MathieuHugo Kavunga-Membolbtihel Smetildowu B. Olawoyelfedayo M. O. Adetifalkponmwosa Odiallhem Boutiba Ben Boubakerlluoreh Ahmed Muhammadlsaac Ssewanyanalsatta Wurielyaloo S. Konstantinus Jacqueline Wemboo Afiwa Halatoko James Ayei Janaki Sonoo Jean-Claude C. Makangara Jean-Jacques M. TamfumJean-Michel HeraudJeffrey G. ShafferJennifer GiandhariJennifer MusyokiJerome NkurunzizaJessica N. UwanibeJinal N. BhimanJiro YasudaJoana MoraisJocelyn KiconcoJohn D. SandiJohn HuddlestonJohn K. OdoomJohn M. MorobeJohn O. GyapongJohn T. KayiwaJohnson C. OkolieJoicymara S. XavierJones GyamfiJoseph F. WamalaJoseph H. K. BonneyJoseph NyandwiJosie EverattJoweria NakaseeguJoyce M. NgoiJoyce NamulondoJudith U. OguzieJulia C. AndekoJulius J. LutwamaJuma J. H. MoggaJustin O'GradyKatherine J. SiddleKathleen VictoirKayode T. AdeyemiKefentse A. TumediKevin S. CarvalhoKhadija Said MohammedKoussay DellagiKunda G. MusondaKwabena O. DueduLamia Fki-BerrajahLavanya SinghLenora M. KeplerLeon BiscornetLeonardo de Oliveira MartinsLucious ChabukaLuicer OlubayoLul Deng OjokLul Lojok DengLynette I. Ochola-OyierLynn TyersMadisa MineMagalutcheemee RamuthMaha MastouriMahmoud ElHefnawiMaimouna MbanneMaitshwarelo I. MatshekaMalebogo KebabonyeMamadou DiopMambu MomohMaria da Luz Lima MendonçaMarietjie VenterMarietou F. PayeMartin FayeMartin M. NyagaMathabo MarekaMatoke-Muhia DamarisMaureen W. MburuMaximillian G. MpinaMichael OwusuMichael R. WileyMirabeau Y. TatfengMitoha Ondo'o AyekabaMohamed AbouelhodaMohamed Amine BeloufaMohamed G. SeadawyMohamed K. KhalifaMooko Marethabile MatoboMouhamed KaneMounerou SalouMphaphi B. MbulawaMulenga MwendaMushal AllamMy V. T. PhanNabil AbidNadine RujeniNadir AbuzaidNalia IsmaelNancy ElquindyNdeye Marieme TopNdongo DiaNédio MabundaNei-yuan HsiaoNelson Boricó SilochiNgiambudulu M. FranciscoNgonda SaasaNicholas BbosaNickson MurungaNicksy GumedeNicole WolterNikita SitharamNnaemeka NdodoNnennaya A. AjayiNoël TordoNokuzola MbheleNorosoa H. RazanajatovoNosamiefan IguosadoloNwando MbaOjide C. KingsleyOkogbenin SylvanusOladiji FemiOlubusuyi M. AdewumiOlumade TestimonyOlusola A. OgunsanyaOluwatosin FakayodeOnwe E. OgahOpe-Ewe OludayoOusmane FayePamela Smith-LawrencePascale OndoaPatrice CombePatricia NabisubiPatrick SemandaPaul E. OluniyiPaulo ArnaldoPeter Kojo QuashiePeter O. OkokherePhilip BejonPhilippe DussartPhillip A. BesterPlacide K. MbalaPontiano KaleebuPriscilla AbechiRabeh El-SheshenyRageema JosephRamy Karam AzizRené G. EssombaReuben Ayivor-DjanieRichard NjouomRichard O. PhillipsRichmond GormanRobert A. KingsleyRosa Maria D. E. S. A. Neto RodriguesRosemary A. AuduRosina A. A. CarrSaba GargouriSaber MasmoudiSacha BootsmaSafietou SankheSahra Isse MohamedSaibu FemiSalma MhallaSalome HoschSamar Kamal KassimSamar MethaSameh TrabelsiSara



Hassan AqwaSarah Wambui MwanqiSeydou DoumbiaSheila Makiala-MandandaSherihane AryeeteyShymaa S. AhmedSide Mohamed AhmedSiham ElhamoumiSikhulile MoyoSilvia LutucutaSimani GaseitsiweSimbirie JallohSoa Fy AndriamandimbySobajo OguntopeSolène GrayoSonia Lekana-DoukiSophie ProsolekSoumeya OuangraouaStephanie van WykStephen F. SchaffnerStephen KanyereziSteve Ahuka-MundekeSteven RudderSureshnee PillaySusan NabaddaSylvie BehillilSylvie L. BudiakiSylvie van der WerfTapfumanei MasheThabo MohaleThanh Le-VietThirumalaisamy P. VelavanTobias SchindlerTongai G. MapongaTrevor BedfordUgochukwu J. AnyanejiUgwu ChineduUpasana RamphalUwem E. GeorgeVincent EnoufVishvanath NeneVivianne GorovaWael H. RoshdyWasim Abdul KarimWilliam K. AmpofoWolfgang PreiserWonderful T. ChogaYahaya Ali AhmedYajna RamphalYaw BediakoYeshnee NaidooYvan ButeraZaydah R. de LaurentAhmed E. O. OumaAnne von GottbergGeorge GithinjiMatshidiso MoetiOyewale TomoriPardis C. SabetiAmadou A. SallSamuel O. OyolaYenew K. TebejeSofonias K. TessemaTulio de OliveiraChristian HappiRichard LessellsJohn NkengasongEduan Wilkinson, and Aaron L. Shibemba, and Abasi Ene Obong, and Abayomi Fadeyi, and Abbad Anas, and Abd Elazeez Shabaan, and Abd Monaem Adel, and Abd Moniem Ain Shoka, and Abdelhamid W., and Abdelilah Laraqui, and Abdelkader Laatiris, and Abdelkrim Meziane Bellefquih, and Abdellah Faouzi, and Abdelmoulah F., and Abdelomunim Essabbar, and Abderrahmane Bimouhen, and Abderraouf Hilali, and Abdo I., and Abdou Padane, and Abdoul Karim Sangaré, and Abdoul Karim Soumah, and Abdoulaye Djimde, and Abdoulaye Toure, and Abdoulie Kanteh, and Abdulla Bashein, and Abdullah Salama, and Abe Lojuan, and Abebe Genetu Bayih, and Abel Abera Negash, and Abel Lissom, and Abid N., and Abla A. Konou, and Abo Shama, and Abosede O., and Abouelnaga S., and Abraham Ali, and Abraham Kwabena Anang, and Abraham Tesfaye, and Adam K. Khan, and Adamu Tayachew, and Adane Mihret, and Adba Alfatih AlEmam, and Adede Hawi, and Adesegun A., and Adey Feleke Desta, and Adib Ghassan, and Adjaratou Traoré, and Adjiratou Aissatou B. A., and Adodo Sadji, and Adrian Egli, and Adriano Mendes, and Adugna Abera, and Adul Candé, and Afaf Alaoui, and Afonso Pedro, and Agbodzi B., and Ageez A.M., and Ahidjo Ayouba, and Ahlam Alarif, and Ahmed E Kaved, and Ahmed El-Taweel, and Ahmed Elsaved, and Ahmed F. Gad, and Ahmed Fakhfakh, and Ahmed Kandeil, and Ahmed M., and Ahmed Mostafa, and Ahmed O. S., and Ahmed Reggad, and Ahmed Taha, and Ahyong Vida, and Aicha Bensalem, and Aida Sivro, and Aissam Hachid, and Ajili F., and Ajogbasile F.V., and Akim A. Adegnika, and Akoele Siliadin, and Akwii Patience Natasha, and Aladje Balde, and Alan Lemtudo, and Alaoui Sanaa-amine, and Alaruusi A. M., and Alassane Ouro-Medeli, and Albert Nyunja, and Alberto Rizzo, and Alemseged Abdissa, and Alemu Tike Debela, and Alessandro Mancon, and Alessandro Marcello, and Alexander Goredema, and Alexander Greninger. and Alexis Ndjolo, and Alexis Niyomwungere, and Alfredo Mari, and Alfredo Mayor, and Ali M.A., and Ali Zumla, and Alia Ben Kahla, and Alia Grad, and Alice Kabanda, and Alie Tia, and Alimou Camara, and Alimuddin Zumla, and Alle Baba Dieng, and Almoustapha I. Maiga, and Amadou Alpha Sall, and Amadou Daou, and Amal Naguib, and Amal Souiri, and Amal Zouaki, and Amalou Ghita, and Amandine Mveang-Nzoghe, and Amariane Koné, and Amariane M. M. Koné, and Ambroise Ahouidi, and Amel Benyahia, and Amel Nagiub, and Amer K. E., and Ameyo Dorkenoo, and Amina Barkat, and Aminata Dia, and Aminata Mbaye, and Aminata Mboup, and Aminata Sileymane Thiam, and Amine Idriss Lahlou, and Amira Suliaman Wadi, and Amivi Ehlan, and Amujal Marion, and Amuri Aziza, and Amy Strydom, and Anass Abbad, and Anatole Nkeshimana, and Andargachew Mulu, and Anderrahmane Maaroufi, and Andrea E. Luquette, and Andreas Shiningavamwe, and Andres Moreira-Soto, and Andrew Azman, and Andrew J. Bennett, and Andrew Tarupiwa, and Anga Latifa, and Ange Badjo, and Angel Angelov, and Angela Brisebarre, and Angela M. Detweiler, and Angoune Ndong, and Ania Werno, and Anna Julienne Selbe NDiave, and Anna-Lena Sander, and Annair B., and Annamaria D'Aprile, and Anne van der Linden, and Annemiek van der Eijk, and Annette Erhart, and Anou M. Somboro, and Anoumou Dagnran, and Anthony Ahumibe, and Anthony Levasseur, and Antje van der Linden, and Antoine Dara, and Anu Jegede-Williams, and Aouni M., and Arjarquah A., and Arlene Uwituze, and Arlo Upton, and Armel Poda, and Arsène Somé, and Arsène Zongo, and Arsenia Massinga, and Asare K.M., and Ashaba Fred Katabazi, and Asma Ferjani, and Assane Dieng, and Astou Gaye-Gaye, and Atiga N., and Atsbeha G. Weldemariam, and Augustina Arjaquah, and Auld A., and Awa Ba-Diallo, and Awatef ElMoussi, and Awunyo Sena, and Aya Mohamed, and Ayman Farghaly, and Ayo-Ale B., and Ayoade F., and Ayola Akim Adegnika, and Ayong More, and Ayorinde Babatunde James, and Azami Nawfel, and Azaria Diergaardt, and Azuka Patrick Okwuraiwe, and Babafemi O. Taiwo, and Baboo S. B., and Bahadoor B. S., and Bahnassy A.A., and Bakary Sanyang, and Bakry U., and Bamba Fatoumata Touré, and Bamidele Iwalokun, and Banda R., and Bane S., and Bankole Johnson, and Barada Cisse, and Barbra Murwira, and Bas Oude Munnink, and Batchi-Bouyou Armel L., and Batra R., and Beatrice Dhaala, and Belayachi Lamiae, and Belkhir A.B., and Ben Ayed I., and Ben Morton, and Ben Moussa, and Ben Wulf, and Bénédicte Ndeboko, and Benhida Rachid, and Benjamiin B. Lindsey, and Benjamin H. Foulkes, and Benjamin Hounkpatin, and Benjamin Selekon, and Bensaid M., and Bernard Mpairwe, and Bernard Ssentalo Bagaya, and Bert Vanmechelen, and Bertrand Lell, and Beth Mutai, and Bethlehem Adnew, and Beuty Makamure, and Bighignoli B., and Birahim Piere Ndiaye, and Bishwo N. Adhikari, and Bitrou S.M., and Blaise MBoringong Akenji, and Bode Shobayo, and Boitumelo Zuze, and Bonifacio Manguire Nlavo, and Bouchra Belfquih, and Bouchra Boujemla, and Bouna Yatassaye,



and Bouzidi Aymane, and Brian Andika, and Bright K. Yemi, and Bronwyn Kleinhans, and Bruna Galvao, and Bubacar Delgado Pinto Embaló, and Bulelani Manene, and Butoyi Pascal, and Camille Capel, and Campbell Anu, and Carine Tchibozo, and Carla Madeira, and Carlos Cortes, and Carniel Elisabeth, and Carol Kifude, and Carolle Yanique Tayimetha, and Catherine E. Arnold, and Catherine Okoi, and Cecilia Waruhiu, and Celestin Godwe, and Celestina Obiekea, and Celine Nkenfou, and Chabuka L. B., and Chakib Nejjari, and Chambaro H., and Chanda D., and Changula K., and Charifa Drissi Touzani, and Charles Kayuki, and Charles Nyagupe, and Charles Ssuuna, and Charoute Hicham, and Chaselynn M. Watters, and Chawech H., and Cheikh Sokhna, and Chenaoui Mohamed, and Chiamaka Nwuba, and Chilufya C., and Chimaobi Chukwu, and Chinyere Anyika, and Chipimo P.J., and Chitanga S., and Chitenje M., and Chiwaula M. J., and Chouati Taha, and Chris Mansell, and Christelle Butel, and Christian A. Devaux, and Christian Drosten, and Christian Ranaivoson, and Christian Utpatel, and Christophe Malabat, and Chtourou A., and Chukwu G., and Claudia Daubenberger, and Clement G. Kakai, and Clement Masakwe, and Clotaire Donatien Rafai, and Collins Chenwi, and Collins M. Misita, and Collins Muli, and Corine H. GeurtsvanKessel, and Corinne Maufrais, and Coulibaly Mbegnan, and Crawford M., and Cristina M. Tato, and Cyrus Yiaba, and D'Amore N., and Dabiri Damilari, and Dalia Ramadan, and Damena D., and Daniel Ouso, and Daniela Pisanelli, and Danilo Licastro, and David Hammer, and David Nieuwenhuijse, and David Patrick Kateete, and David Wilkinson, and Davy Leger Mouangala, and Dawit Hailu Alemayehu, and Dawood R.M., and De Sanctis R., and Deborah N. A. Mettle, and Demba Koita, and Dennis Kenyi Lodiongo, and Dennis Laryea, and Dennis N. Wandera, and Dereje Leta, and Dersi Noureddine, and Desire Takou, and Dessalegn Abeje Tefera, and Dhwani Batra, and Dia Ndongo, and Diab A., and Diabou Diagne, and Diane A. Mabika, and Diané Bamourou, and Dianke Samaté, and Diarra B., and Didier Raoult, and Dieudonne Mutangana, and Dineo Emang Tshiamo. Gape Nyepetsi, and Diosdado Odjama Nseng Ada, and Djimabi Salah, and Donatella Cedola, and Doris Harding, and Dorothy Yeboah-Manu, and Dossou Ange, and Dowbiss Meta Djomsi, and Dragana Drinkovic, and Draper C., and Dumbuya Foday Sahr, and Ebenezer Odewale, and Eclou Sedjro, and Edang-Minko Armand, and Edgar Kigozi, and Edith Koskei, and Edith Nkwembe, and Edmilson F. de Oliveira Filho, and Edmira Maria da Costa, and Edward Kiritu, and Efrem S. Lim, and Egon A. Ozer, and Egyir B., and Ehab Abdelkader Abuelenein, and Eitel Mpoudi Ngole, and El Aliani Aissam, and El Ansari Fatima Zahra, and El Hamouchi Adil, and El Oualid Abdelmjid, and El-Shagngery H., and El-Zayat M., and Elamin Abualas, and Elannaz Hicham, and Elargoubi A., and Eleni Kidane, and Elham Rizgalla, and Elhoseny M.M., and Elhosieny F.W., and Elisabeth Carniel, and Elizabeth Nyakarungu, and Elkhateeb S.M., and Elmostafa Benaissa, and Elmostafa El Fahime, and Elouanass M., and Elsissy M.H., and Elvyre Mbongo-Nkama, and Elwaleed M. Elamin Sara A.I Latif, and Emame Edu, and Emanuele Orsini, and Emilio Skarwan, and Emily K. Stefanov, and Emmanuel Nasinghe, and Emmanuel Nepolo, and Emmanuel Ogunbayo, and Emmanuelle Munger, and Emmanuelle Permal, and Emna Gaies, and Ennibi H., and Ennibi Khalid, and Erasmus Kotey, and Erasmus Smit, and Eric A. Lelo, and Eric Adu, and Eric Delaporte, and Eric Katagirya, and Eric Kezakarayagwa, and Eric Muthanje, and Erica Luis Maria Magalhães, and Ernest Asiedu, and Esemu Livo, and Esperance Umumararungu, and Esther Chitechi, and Esther Omuseni, and Etoundi Mballa Alain, and Evelyn Bonney Quansah, and Evelyn Y. Bonney, and Ezzelarab M.H., and Faatu Cassama, and Fabien Roch Niama, and Fabio Arena, and Faggioni G., and Fahsbender E., and Faith Sigei, and Faouzi Abdellah, and Farah Jouali, and Farawyla H., and Farida Hilali, and Fatima Chgouri, and Fatima El Falaki, and Fatima Zahra El ansari, and Fatma Ebied, and Fatna Bssaibis, and Fattah Al Onifade, and Faye Ousmane, and Fayez Ahmed Khardine, and Fayez Khardine, and Fedorov A.V., and Fekadu Alemu, and Fekkak Jamal, and Fengming Sun, and Fetouma Doudou, and Fikry A.E., and Fillo S., and Fiorenza Bracchitta, and Firmin Kaboré, and L. Fki-berrajah, and Flora Donati, and Florence Fenollar, and Foday Sahr Samuel Sorie, and Folarino O., and Folorunsho Francine, and Kabatesi Francisca, and Muyembe-Mawete Francisco, and Malagon François, and Kiemdé Franklin, and Asiedu-Bekoe Franklyn Egbe, and Nkongho Frédéric, and Lemoine Frederick, and Tei-Maya, and Freitas R. H., and Fuh-Neba T., and Gaaloul I., and Gabriel Kabamba, and Gadissa Gutema, and Gaies Emna, and Galadima Gadzama, and Galal Mahmoud, and Garba Quangole, and Gargouri S., and Garry R., and Gary McAuliffe, and Gathii Kimita, and Gédéon Prince Manouana, and Gelanew Tesfaye, and George Awinda, and George B. Kyei, and George Michuki, and Georgelin Nguema Ondo, and Gerhard van Rooyen, and Gert Marais, and Getachew Abichu, and Getachew Tesfaye Beyene, and Getachew Tollera, and Getnet Hailu, and Ghamaz Hamza, and Ghazi Kayali, and Ghizlane El Amin, and Gibson Mhlanga, and Gilbert Kibet, and Gildas Hounkanrin, and Giordani F., and Gizaw Teka, and Godfrey Masete, and Goldstone R., and Gomaa C., and Gomaa Mokhtar, and Gora Lo, and Gosnell B. I., and Gottberg A., and Grace Angong Belournou, and Grace Oni, and Grace Vincent, and Gregory Wani, and Guedi Ali Barreh, and Guillermo Garcia, and Guohong Deng, and Guseva N. P., and Guy Paterne Malonga Mbembo, and Guy Stéphane Padzys, and Gwayi S., and Habiba Ben Romdhane, and Habiba Naija, and Haby Diallo, and Hadad A., and Hafez M.M., and Hafsia Ladhari, and Hagar Elshora, and Hailu Dadi, and Hajjaji Mohammed, and Hakima Kabbaj, and Hala Hafez, and Halafawy A., and Halidou Tinto, and Halimatou Diop Ndiaye, and Hamadi Assane, and Hamdani T.N., and Hamdy M.S., and Hamidah Namagembe, and Hammad M., and Hammami A., and Hamza Gharmaz, and Hana Sofia Andersson, and



Hanae Dakka, and Hanen El Jebari, and Hany K. Soliman, and Harmak Houda, and Harvey R., and Hasnae Benkirane, and Hassan Aguenaou, and Hassan Ihazmad, and Hassan R., and Hassan W., and Hatem Ahmed, and Heitzer Sogodogo, and Helena Seth-Smith, and Helisoa Razafimanjato, and Hellen Koka, and Hemlali Mouhssine, and Henry Mwebesa, and Hermes Perez, and Herve Christian Paho Tchoudjin, and Hicham Elannaz, and Hicham Oumzil, and Hilda Opoku Frempong, and Hinson Fidelia, and Hoda Ezz Elarab, and Hong Xie, and Houda Benrahma, and Housna Arrouchi, and Houssem Guedouar, and Hu Luo, and Hubert Bassène, and Huiging Si, and Ian Goodfellow, and Ibrahima Guindo, and Ibrahima Halilou, and Idil Salah Abdillahi, and Idriss-Amine Lahlou, and Idrissa Diawara, and Ifunanya Egoh, and Ige F.A., and Iknane A.A., and Ikram Bnouyahia, and Ikram Omar Osman, and Ilhem Boutiba Ben Boubaker, and Iman Abdillahi Hassan, and Iman Foda, and Imane Smyej, and Imen Kacem, and Imen Mdini, and Imen Mkada, and Inacio Mandomando, and Iñaki Comas, and Ines Mdini, and Inglês L., and Innocent Mudau, and Ireoluwa Yinka JOEL, and Irina Chestakova, and Irving Cancino, and Isaac Phiri, and Ismael Pierrick Mikelet Boussoukou, and Ismail A., and Israel Osei-Wusu, and Issaka Maman, and Ivan Barilar, and Ivy A. Asante, and Izuwayo Gerard, and Jaafar Heikel, and Jacob Souopgui, and Jacques Marx, and Jalal D., and Jalal Nourlil, and Jalil El Atar, and Jalila Ben Khelil, and Jalila Rahoui, and Jamal Fekkak, and James Mutisya, and James Ussher, and Jan Felix Exler, and Jane MaCauley, and Janet Majanja, and Jannoo N., and Jarra Manneh, and Jasmin Scharnberg, and Jasmin Schlotterbeck, and Jasper Chimedza, and Jawad Bouzid, and Jean B. Niyibigira, and Jean Claude Djontu, and Jean Maritz, and Jean-Claude Makangara Cigolo, and Jean-louis Monemou, and Jeanne d'Arc Umuringa, and Jeremy Delerce, and Jerome Ndaruhutse, and Jerome Nkurunziza, and Jesse Addo Asamoah, and Jill Sherwood, and Jing Wang, and Joan Marti-Carerras, and Joe K. Mutungi, and Joe Mutungi, and Joel Fleury Djoba Siawaya, and Joel Koivogui, and Joep de Ligt, and John Njuguna, and John Rumunu, and John Tembo, and John Waitumbi, and Johnson A. Adeniji, and Jonathan Rigby, and Jorn Hellemans, and Joseph Fokam, and Joseph L. DeRisi, and Joseph Makhema, and Joseph Mugisha, and Joseph Ojonugwa Shaibu, and Joseph Oliver-Commey, and Josephine Bwogi, and Josh Freeman, and Josiah Ayoola Isong, and Josphat Nyataya, and Joy Ayoola, and Joyce Appiah-Kubi, and Judd F. Hultquist, and Jude Gedeon, and Judith Sokei, and Julia Howard, and Julia Schneider, and Julian Campbell, and Juliet Elvy, and Jumaa A.B., and Justin Lee, and Justin Lessler, and Justin O'Grady, and Kaba Kourouma, and Kais Ghedira, and Kalantar K., and Kamela Mahlakwane, and Kamoun S., and Kangwa Mulonga, and Kapata P.C., and Kapaya F., and Kapin'a M., and Karray Hakim H., and Kasambara W., and Kasmi Yassine, and Kassahun Tesfaye, and Kathleen Subramoney, and Katyshev A.D., and Kayeyi N., and Kayla Barnes, and Kayla Delaney, and Kazorina E.V., and Kedumetse Seru, and Keita S., and Keith Durkin, and Keith R Jerome, and Keke K. René, and Kena Swanson, and Kenneth K. Maeka, and Keren Okyerebea Attiku, and Kevin Sanders, and Kevine Zang Ella, and Keyru Tuki, and Khabab A. Elhag, and Khadim Gueye, and Khaled Amer, and Khalid Ennibi Mostafa, and Kharat N., and Khumalo Z., and Kilian Stoecker, and Kim L., and Kimberly A. Bishop-Lilly, and Kimotho J., and Kitane Driss Lahlou, and Kofi Bonney, and Kokou Tegueni, and Kolawole Wasiu Wahab, and Kolomoets E.V., and Komal Jain, and Kominist Asmamaw, and Komlan Kossi, and Kondwani Jambo, and Kouadio T. Yao, and Kouriba Dürr, and Kra Ouffoué, and Krasnov Y.M., and Krishna Kumar Kandaswamy, and Kristian Andersen, and Kritsky A.A., and Kumbelembe David, and Kutlo Macheke, and Kutyrev V.V., and Kwenda S., and Kwitaka Maluzi, and Kwok Lee, and Kyle A. Long, and Kyra Grantz, and Lacy M. Simons, and Laetitia Serrano, and , and Laila Elsawy, and Laila Sbabou, and Lallepak Lamboni, and LaRinda A. Holland, and Lasata Shrestha, and Lassana Sangaré, and Latifa Anga, and Lauren Jelly, and Laurien Hoornaert, and Le Thi Kieu Linh, and Legodile Kooepile, and Leigh-Anne MC Intyre, and Léon Mutesa, and Leona Okoli, and Léopold Ouedraogo, and Lesego Kuate-lere, and Leta D., and Letaief A., and Liboro G., and Lilian Kanjau, and Lin L., and Linda Boatemaa, and Linda Houhamdi, and Lipkin W. Ian, and Lista F., and Liwewe M.M., and Lloyd Mulenga, and Logan J. Voegtly, and Loide Shipingana, and Loris Micelli, and Lorreta Kwasah, and Loubna Allam, and Louise Lefrançois, and Loukman Salma, and Lucas N. Amenga-Etego, and Ludivine Brechar, and Ludovic Mewono, and Luis A. Estrella, and Lusia Mhuulu, and Lwanga Newton, and M. Ulrich, and M.T. Mogotsi, and Maaroufi Abderrahmane, and Mad W., and Madi W.Y., and Madlen Stange, and Magdeldin S., and Maher Kharrat, and Mahlangu B., and Mahmoud Shehata, and Mahrous N., and Maida A., and Makhtar Camara, and Makori T., and Malama K., and Malena S. Bestehorn-Willmann, and Malolo I., and Mamadou Beye Cheikh Ibrahima Lo, and Mamadou Bhoye Keita, and Mamadou Saliou Bah, and Mamadou Saliou Sow, and Mamoudou Harouna Djingarey, and Mamoudou Maiga, and Manal Hamdy Elsaid, and Manal Hamdy Zahran, and Mandiou Diakite, and Manel Ben Sassi, and Mangombi Pambou, and Manickchund N., and Manoli Torres Puente, and Manrai S. S., and Mansour T., and Maowia M. Mukhtar, and Marcel Tongo, and Marchoudi Nabila, and Margaret Mills, and Maria Artesi, and Maria Pia Patrizio, and Maria Rita Gismondo, and Maria Rosaria Lipsi, and Mariam Kehinde SULAIMAN, and Mariama Kujabi, and Marie Amougou, and Marie Claire Okomo, and Marie Madeleine Chabert-Consen, and Marie-Astrid Vernet, and Marie-Pierre Hayette, and Mariem Gdoura, and Marijke Reynders, and Marion Barbet, and Marion Koopmans, and Marjan Boter, and Mark Siedner, and Markos Abebe, and Markus H. Antwerpen, and Marouane Melloul, and Martin Maidadi Foudi, and Martine Peeters, and Marvin Hsiao, and Mary DeAlmeida, and Mary Lalemi, and Mary-

Science

Ann Davies, and Masahiro K., and Masse Sambou, and Mathabo M., and Mathew D. Parker, and Mathias C. Walter, and Mathur H., and Matt Blakiston, and Matt Storey, and Matthew Bates, and Matthew Rogers, and Matthias Pauthner, and Maud Vanpeene, and Maurizio Margaglione, and Max Bloomfield, and May Abdelfattah, and May Sherif Soliman, and Mbengué Fall, and Mdlalose K., and Meei-Li Huang, and Mehta S., and Mélanie Albert, and Melchior A. Joël Aïssi, and Méline Bizard, and Merabet Mouad, and Meriem Laamarti, and Messanh Douffan, and Mhalla S., and Michael Addidle, and Michael Marks, and Michael Nagel, and Michael V. Deschenes, and Michaela Davids, and Michael Balm, and Michael Lin, and Michelle Tan, and Mihrete A., and Mikhail Olayinka BUHARI, and Milanca Agostinho Cá, and Mildred Adusei-Poku, and Milkah Mwangi, and Mina Kamel, and Miranda J., and Mireille Prince-David, and Miriam Eshun, and Misaki Wayengera, and Mitali Mishra, and Mjid Eloualid, and Mly Abdelaziz Elalaoui, and Mnguni A., and Mnyameni F., and Mogomotsi Matshaba, and Mohale T., and Mohamed Abdel-Salam Elgohary, and Mohamed Ahmed Ali, and Mohamed Ben Moussa, and Mohamed Chenaoui, and Mohamed El Sayes, and Mohamed Elhadidi, and Mohamed Gomaa Seadawy, and Mohamed Hassan Abdoelraheem, and Mohamed Hassany, and Mohamed Houmed Aboubaker, and Mohamed K.S., and Mohamed Kamal, and Mohamed Rhajaoui, and Mohamed Seadawy, and Mohamed Shamel, and Mohamed Shemis, and Mohammed K.S., and Mohammed Walid Chemao Elfihri, and Mohcine Bennani Mechita, and Mokhtar Gomaa, and Molalegne Bitew, and Momoh M., and Mona O.A. Alkarim, and Monemo Pacome, and Monilade Akinola, and Monte A., and Monuir G., and Monze M., and Mooko M., and Morales A.N., and Moreira-Soto Andres, and Moriba Povogui, and Mosepele Mosepele, and Moses Chilufya, and Moses Joloba, and Moses Luutu, and Mostafa Elouennass, and Mostfa Elhoseiny, and Mostfa Elnakib, and Mostfa Yakout, and Mouhcine Gardoul, and Mouhssine Hemlali, and Mouity Matoumba A., and Mouna Ben Sassi, and Mouna Safer, and Mouneem Essabbar, and Moustapha Mbow, and Moustapha Nzamba Maloum, and Moustapha Sakho, and Moyinoluwa Odugbemi, and Mtshali P., and Mubemba B., and Muchaneta Mugabe, and Mufinda M., and Muhammad Faisal, and Muinah Adenike Fowora, and Mukantwari Enatha, and Muleya W., and Muntaser elTayeb Ibrahim, and Mupeta F., and Murebwayire Clarisse, and Mushal Ali, and Mushal Allam, and Mustapha Mouallif, and Muuo S.N., and Mwangomba W., and Myriam Seffar, and Mzumara T. E., and N'dilimabaka N., and Nabila Soara, and Nabli A., and Nadia El Mrimar, and Nadia Rodrigues, and Nadia Sitoe, and Nafisatou Leye, and Naguib A., and Nalubamba K. S., and Nancy M. El Guindy, and Nandi Siegfried, and Nardjes Hihi, and Narjis Amar, and Naryshkina E.A., and Nathalia Endjala, and Nathalia Garus-Oas, and Nathan Kapata, and Ndack Ndiaye, and Ndahafa Frans, and Ndam N.T., and Ndéye Coumba Touré Kane, and Ndiaye Ndack, and Ndine Fainguem, and Ndodo Nnaemeka, and Ndumbu Pentikainen, and Ndwiga L., and Nedio Mabunda, and Neff N., and Negash A.A., and Nejla Stambouli, and Neto Z., and Ngonga Dikongo A.M., and Ngosa W., and Ngozi Mirabel Otuonye, and Niatou-Singa F. S., and Nicaise T. Ndam, and Nicholas Feasey, and Nicholas Mwikwabe, and Nicod J., and Nicole Vidal, and Nikki Freed, and Nischay Mishra, and Nissaf Ben Alaya, and Nitin Savaliya, and Noah Baker, and Noé Patrick Mbondoukwe, and Nokukhanya Mdlalose, and Nonso Nduka, and Noura M Abo Shama, and Nourlil Jalal, and Nsubuga Gideon, and Ntuli N., and Nuro Abilio, and Nyam Itse Yusuf, and Oby Wayoro, and Ochwoto M., and Ofonime Ebong, and Ofori-Boadu L., and Okomo Assoumou Marie Claire, and Ola Elroby, and Olabisi Olabisi, and Ojo Olajumoke, and Popoola, and Olfert L., and Olin Silander, and Olufemi Obafemi, and Olufemi Samuel Amoo, and Olukunke Oluwasemowo, and Olusola Anuoluwapo Akanbi, and Oluwakemi Laguda-Akingba, and Oluwatimilehin Adewumi, and Omar Askander, and Omar Elahmer, and Omar S., and Omilabu S., and Omnia Kutkat, and Omoare Adesuyi, and Omondi Francis Carey, and Onalethata Lesetedi, and Ongera E., and Ontlametse T. Bareng, and Onwuamah C.K., and Ope-Ewe O., and Osama Mansour, and Oscar Kanjerwa, and Oteng F., and Otmane Touzani, and Oumaima Ait Si Mohammed, and Oumy Diop, and Ouna Ouadghiri, and Ousseynou Gueye, and Owusu-Nyantakyi C., and Oyefolu A., and Oyeronke Ayansola, and P Nthiga, and Palomba S., and Panja L., and Papa Alassane Diaw, and Park D., and Pasacaline Manga, and Patel H., and Patience Motshosi, and Patoo M., and Patrick Amoth, and Patrick Descheemaeker, and Patrick Mavingui, and Patrick Tuyisenge, and Pattoo M., and Paul Dobi, and Paul Liberator, and Paulin N. Essone, and Paulina Joãozinho da Costa Jarra Manneh, and Pauline Yacine Sene, and Paulo A. Carralero R.R. Paixão, and Pavitra Roychoudhury, and Peace O. Uche, and Pei Zhou, and Penda Malhado Diallo, and Pereira A., and Petas Akogbeto, and Peter Bauer, and Peter T. Skidmore, and Petra Raimond, and Phasha-Muchemenye Mmatshepho, and Philip Ashton, and Philip C., and Philip El-Duah, and Philip M. Soglo, and Philip Wonder Phiri, and Philipp Wagner, and Philippe Colson, and Philippe Dussart, and Philippe Lavrard Meyer, and Philip Ashton, and Philonah Tushabe, and Pierre-Edouard Fournier, and Piet Maes, and Popova A. Yu, and Portia Manangazira, and Praise Adewumi, and Qi Yang, and Quaneeta Mohktar, and Quansah E.B., and Quashie P., and Quedraogo Rabia, and Magsood Rachel, and Githii Rachid, and Abi Rachid, and Benhida Rachid, and El Jaoudi Rachid, and Mentag Rahaman A., and Ahmed Rahma, and Algheriani Raiva, and Simbi Rajiha, and Abubeker Ramalia Chabi, and Nari Ramon, and Lorenzo-Redondo Ramy, and Galal, and Raouf A., and Raoul Saizonou, and Raphael Lumembe, and Ravena Mubichi, and Regina Z. Cer, and Reham Dawood, and Reham Kassab, and Rehema Liyai, and Rehn A., and Rei José Pereira, and Reina Sikkema, and Rfaki Abderrazak, and Riad Mounir Armanious, and Riadh Daghfous, and Riadh Gouider, and Richard Adegbola, and

Science

Richard Lino Loro Lako, and Richard Molenkamp, and Richard Webby, and Richmond Yeboah, and Rick S., and Rida Tagajdid, and Rine Zeh Nfor, and Rivalyn Nakoune Yandoko, and Robert Newton, and Robert Rutayisire, and Rodney S. Daniels, and Rodrigue Bikangui, and Rodrigue K. Kohoun, and Rodrigue Kamga, and Rodrigue Mintsa Nguema, and Roger Shapiro, and Rogers J., and Rogers Kamulegeya, and Rokaia Laamrti, and Roméo Aimé Laclong Lontchi, and Ronald Kiiza, and Ronald M. Galiwango, and Rosella De Nittis, and Roshdy W.H., and Rotimi Myrabelle Avome Houechenou, and Roua Ben Othman, and Rui Inndi, and Saad M.A., and Saaïd Amzazi, and Sabin Nsanzimana, and Sada Diallo, and Sadji Y.A., and Safae El Mazouri, and Safae Elkochri, and Safae Ghoulame, and Safiatou Karidioula, and Safietou Sankhe, and Sahr Gevao, and Sahr P.F., and Saibu J.O., and Saïdou Ouedraogo, and Saiid S., and Sainabou Laye Ndure, and Sakoba Keita, and Salah D., and Salah Eldin Hussein, and Salah H., and Saleh A.A., and Saleh M., and Salifou Sourakatou, and Sally Roberts, and Salma Abid, and Salma Sayed, and Salou M., and Salu O.B., and Sam Lissauer, and Sam O'neilla Oye Bingono, and Samba Ndiour, and Sameira M. Fageer, and Samia Abdou Girgis, and Samir M., and Samir O., and Samira Benkeroum, and Samira F. Ibrahim, and Samira M. Fageer, and Samira Zoa Assoumou, and Samirah Saiid, and Samoel Ashimosi Khamadi, and Samson Konongoi Limbaso, and Samuel Armoo, and Samuel Kirimunda, and Samuel Sorie, and Samwel Lifumo Symekher, and Samwel Owaka, and Sana Ferjani, and Sanaa Alaoui-Amine, and Sander Anna-Lena, and Sankhe Safietou, and Santiago Jiménez-Serrano, and Santigie Kamara, and Sara Chammam, and Sara Mahmoud, and Sarah Jefferies, and Sarah Rubin, and Sarah Stanley, and Saraswathi Sathees, and Saro Abdella, and Sarra Chamman, and Savannah Mwesigwa, and Sawa H., and Seadawy M.G., and Sean Ellis, and Sébastien Bontems, and Sefetogi Ramaologa, and Sekesai Zinyowera, and Selassie Kumordjie, and Sen Claudine Henriette Ngomtcho, and Sena Awunyo, and Serge Alain Sadeuh-Mba, and Serigne Saliou Niane, and Seth Okeyo, and Settimia Altamura, and Seyram Bless Agbenyo, and Shah Mohamed Bakhash, and Shahin Lockman, and Shahinaz.A. Bedri, and Shaimaa Soliman, and Shalaby L., and Shannon Wilson, and Sharmini Muttaiyah, and Sharon Abimbola, and Sharon Hsu, and Shcherbakova S.A., and Shebbar Osiany, and Shereen Shawky, and Sherine Helmy, and Shevtsova A.P., and Shimaa Moustafa, and Shirlee Wohl, and Shirley Johane, and Sidonie A.M. Kagnissode, and Silvanos Mukunzi Opanda, and Sim Mayaphi, and Simão Tchuda Bióté, and Simeone Dal Monego, and Simon Peter Ruhweza, and Simone Eckstein, and Sindayiheba Reuben, and Sinyange N., and Sivaramakrishna Rachakonda, and Soa Fy Andriam, and Sofia Viegas, and Sogodogo Sokhna, and Ndongo Sola, and Ajibaye Solomon, and Langat Sombo, and Fwoloshi Somda Soro Georgina, and Charlene Sonal P., and Henson Sondes, and Haddad Sonia V., and Bedié, and Sosedova E.A., and Souad Kartti, and Souissi Amira, and Souleymane Mboup, and Soundélé Maïté, and Sounkalo Dao, and Sourakatou Salifou, and Srinivas Reddy Pallerla, and Stefan Niemann, and Steffen Borrmann, and Stephen Asiimwe, and Stephen M. Eggan, and Stephen Ochola, and Stephen Wanok, and Steven J. Reynolds, and Subomi Olorunnimbe, and Sudhir Bunga, and Sujeewon C., and Sunday Babatunde, and Susan Engelbrecht, and Susan Morpeth, and Susan Taylor, and Susann Handrick, and Swaibu Gatare, and Sylvie Melingui, and Symeker S.L., and Syntyche Devatchagni, and Taha A.G., and Taha Chouati, and Taha Maatoug, and Tahar Baijou, and Takada A., and Taloa K.A., and Tamravehu Sevoum, and Tan M., and Tania Stander, and Tanja Niemann, and Tarek Aanniz, and Tarek Refaat Elnagdy Raafat Zaher, and Tatenda Takawira, and Tato C., and Tefera D.A., and Tei-Maya Fred, and Tesfaye Gelanew, and Tesfaye Rufael, and Thanh Le Viet, and Thela Tefelo, and Thérèse Kagoné, and Thibaut Armel Cherif Gnimadi, and Thiongo K., and Thomas Briese, and Thomas Van L., and Thongbotho Mphoyakgosi, and Thushan I de Silva, and Tim Roloff, and Timothy Blackmore, and Tiziana Rollo, and Tom Lutalo, and Tomade A.M. Ibrahim, and Tombolomako T. B., and Tomiwa Adepetun, and Tomkins-Tinch C., and Tony Wawina-Bokalanga, and Tope Sobajo, and Touil Nadia, and Touria Essayagh, and Toy Nwako, and Traoré, and Triki H., and Tsiry R., and Tsiry Randriambolamanantsoa, and Tumisang Madisa, and Turki M., and Ugwu C.A., and Uyi Emokpae, and Valeria Delli Carri, and Valeria Micheli, and Van Rooyen G., and Vanaerschot M., and Vanessa Magnussen, and Vanessa Mohr, and Vani Sathyendran, and Veronica Playle, and Viana R., and Vickos U., and Victor Max Corman, and Victor Mukonka, and Victor Ofula, and Vida Ahyong, and Vincent Appiah, and Vincent Bours, and Violette V. M'cormack, and Virginia Hope, and Vivi Hue-Trang Lieu, and Vololoniaina Raharinosy, and Wadegu Meshack, and Wadonda N., and Wadula J., and Wael Ali, and Waidi Sule, and Wallace D Bulimo, and Warren Cyrus Yiaba, and Waruhiu C.N., and Wasfi Fares, and Webby R., and Weldemariam A.G., and Wendy Karen Jo, and Woelfel R., and Wolh S., and Wurie I., and Xavier Crespin, and Xiaoyun Ren, and Xiuhua Wang, and Ya.M. Krasnov, and Yacine Amet Dia, and Yacine DIA Seni Ndiaye, and Yacob Mohamed Yusuf, and Yacouba Sawadogo, and Yadouleton Yahya, and Maidane Yakob Gebregziabher, and Tsegay Yao, and Layibo Yasser, and El Hady Yassine, and Sekhsokh Yassmin, and Moatasim Yawo A., and Sadji, and Yeboah C., and Youbi Mohammed, and Yousif Rabih Makki, and Youssef Akhoud, and Yuri Ushijima, and Yusuf Jimoh, and Yvette Badou, and Zablon J. Matoke D., and Zablon J.O., and Zaineb Hamzaoui, and Zakia Regragui, and Zara Wuduri, and Zein Souma, and Zeinab Ali Waberi, and Zeinab S. Imam, and Zekiba Tarnagda, and Zemmouri Faouzia, and Zhang J., and Zhenghui Li, and Zimmerman Maiga, and Zohour Kasmy, and Zong Minko O., and Zorgani A., and Zouheir Yassine, and Zoukaneirii Issa, and Zulu P., and Zuzheng Xiang



Science, 378 (6615), eabq5358. • DOI: 10.1126/science.abq5358

Surveillance across Africa

The past 2 years, during which waves of severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) variants swept the globe, have starkly highlighted health disparities across nations. Tegally *et al.* show how the coordinated efforts of talented African scientists have in a short time made great contributions to pandemic surveillance and data gathering. Their efforts and initiatives have provided early warning that has likely benefited wealthier countries more than their own. Genomic surveillance identified the emergence of the highly transmissible Beta and Omicron variants and now the appearance of Omicron sublineages in Africa. However, it is imperative that technology transfer for diagnostics and vaccines, as well the logistic wherewithal to produce and deploy them, match the data-gathering effort. —CA

View the article online

https://www.science.org/doi/10.1126/science.abq5358

Permissions

https://www.science.org/help/reprints-and-permissions