

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

Hourijyah Tegally^{1,2}†, James E. San^{1,2}†, Matthew Cotten^{3,4}, Bryan Tegomoh^{5,6}, Gerald Mboowa⁷, Darren P. Martin^{8,9}, Cheryl Baxter^{1,10}, Monika Moir¹, Arnold Lambisia¹¹, Amadou Diallo¹², Daniel G. Amoako^{13,14}, Moussa M. Diagne¹², Abay Sisay^{15,16}, Abdel-Rahman N. Zekri¹⁷, Abdelhamid Barakat¹⁸, Abdou Salam Gueye¹⁹, Abdoul K. Sangare²⁰, Abdoul-Salam Ouedraogo²¹, Abdourahmane SOW²², Abdualmoniem O. Musa^{23,24,25}, Abdul K. Sesay²⁶, Adamou LAGARE²⁷, Adedotun-Sulaiman Kemi²⁸, Aden Elmi Abar²⁹, Adeniji A. Johnson³⁰, Adeola Fowotade^{31,32}, Adewumi M. Olubusuyi^{30,33}, Adeyemi O. Oluwapelumi^{34,35}, Adrienne A. Amuri^{36,37}, Agnes Juru³⁸, Ahmad Mabrouk Ramadan³⁹, Ahmed Kandeil⁴⁰, Ahmed Mostafa⁴⁰, Ahmed Rebai⁴¹, Ahmed Sayed⁴², Akano Kazeem⁴³, Aladje Balde^{44,45}, Alan Christoffels⁷, Alexander J. Trotter⁴⁶, Allan Campbell⁴⁷, Alpha Kabinet KEITA^{48,49}, Amadou Kone⁵⁰, Amal Bouzid⁵¹, Amal Souissi⁴¹, Ambrose Agweyu¹¹, Ana V. Gutierrez⁴⁶, Andrew J. Page⁴⁶, Anges Yadouleton⁵², Anika Vinze⁵³, Anise N. Happi⁴³, Anissa Chouikha⁵⁴, Arash Iranzadeh^{8,9}, Arisha Maharaj¹, Armel Landry Batchi-Bouyou⁵⁵, Arshad Ismail¹³, Augustina Sylverken⁵⁶, Augustine Goba^{57,58}, Ayoade Femi^{43,59}, Ayotunde Elijah Sijuwola⁴³, Azeddine Ibrahim⁶⁰, Baba Marycelin^{61,62}, Babatunde Lawal Salako^{28,33}, Bamidele S. Oderinde⁶¹, Bankole Bolajoko⁴³, Beatrice Dhaala³, Belinda L. Herring¹⁹, Benjamin Tsofa¹¹, Bernard Mvula⁶³, Berthe-Marie Njanpop-Lafourcade¹⁹, Blessing T. Marondera⁶⁴, Bouh Abdi KHAIREH^{65,66}, Bourema Kouriba²¹, Bright Adu⁶⁷, Brigitte Pool⁶⁸, Bronwyn McInnis¹⁷, Cara Brook^{69,70}, Carolyn Williamson^{9,10,71}, Catherine Anscombe^{72,73}, Catherine B. Pratt⁷⁴, Cathrine Scheepers^{13,75}, Chantal G. Akoua-Koffi^{76,77}, Charles N. Agoti^{11,78}, Cheikh Loucoubar¹², Chika Kingsley Onwuamah⁷⁹, Chikwe Ihekweazu⁸⁰, Christian Noël MALAKA⁸¹, Christophe Peyrefitte¹², Chukwuma Ewean Omoruyi^{31,32}, Clotaire Donatien Rafai⁸², Collins M. Morang'a⁸³, D. James Nokes^{11,84}, Daniel Bugembe Lule³, Daniel J. Bridges⁸⁵, Daniel Mukadi-Bamuleka³⁶, Danny Park⁵³, David Baker⁴⁶, Deelan Doolabh⁹, Deogratius Ssemwanga^{3,86}, Derek Tshiabuila², Diarra Bassirou¹², Dominic S.Y. Amuzu⁸³, Dominique Goedhals⁸⁷, Donald S. Grant^{57,58,88}, Donwilliams O. Omuoyo¹¹, Dorcas Maruapula⁸⁹, Dorcas Waruguru Wanjohi⁷, Ebenezer Foster-Nyarko⁴⁶, Eddy K. Lusamaki^{36,37,49}, Edgar Simulundu⁹⁰, Edidah M. Ong'era¹¹, Edith N. Ngabana^{36,37}, Edward O. Abworo⁹¹, Edward Otieno¹¹, Edwin Shumba⁶⁴, Edwine Barasa¹¹, EL BARA AHMED^{92,93}, Elizabeth Kampira⁹⁴, Elmostafa EL FAHIME⁹⁵, Emmanuel Lokilo³⁶, Enatha Mukantwari⁹⁶, Erameh Cyril⁹⁷, Eromon Philomena⁴³, Essia Belarbi⁹⁸, Etienne Simon-Loriere⁹⁹, Etilé A. Anoh⁷⁶, Fabian Leendertz⁹⁸, Fahn M. Taweh¹⁰⁰, Fares Wasfi⁵⁴, Fatma Abdelmoula^{41,101}, Faustinos T. Takawira³⁸, Fawzi Derrar¹⁰², Fehintola V Ajogbasile⁴³, Florette Treurnicht¹⁰³, Folarin Onikepe^{43,59}, Francine Ntoumi^{55,104}, Francisca M. Muyembe^{36,37}, FRANCISCO NGIAMBUDULU¹⁰⁵, Frank Edgard ZONGO Ragomzingba¹⁰⁶, Fred Athanasius, DRATIBI^{107,108}, Fred-Akintunwa Iyanu⁴³, Gabriel K. Mbunu³⁷, Gaetan Thilliez⁴⁶, Gemma L. Kay⁴⁶, George O. Akpede⁹⁷, George Uwem E.¹⁰⁹, Gert van Zyl^{110,111}, Gordon A. Awandare⁸³, Grit Schubert⁹⁸, Gugu P. Maphalala¹¹², Hafaliana C. Ranaivoson⁷⁰, Hajar Lemriss¹¹³, Hannah E Omunakwe¹¹⁴, Harris Onywera⁷, Haruka Abe¹¹⁵, HELE KARRY¹¹⁶, Hellen Nansumba¹¹⁷, Henda Triki⁵⁴, Herve Albéric ADJE KADJO¹¹⁸, Hesham Elgahzaly¹¹⁹, Hlanai Gumbo³⁸, HOTA mathieu¹²⁰, Hugo Kavunga-Membo³⁶, Ibtihel Smeti⁴¹, Idowu B. Olawoye⁴³, Ifedayo Adetifa³¹, Ikponmwosa Odia⁹⁷, Ilhem Boutiba-Ben Boubaker^{121,122}, Isaac Ssewanyana¹¹⁷, Isatta Wurie¹²³, Iyaloo S Konstantinus¹²⁴, Jacqueline Wemboo Afiwa

Halatoko¹²⁵, James Ayei¹²⁶, Janaki Sonoo¹²⁷, Jean Bernard LEKANA-DOUKI¹²⁸, Jean-Claude C. Makangara^{36,37}, Jean-Jacques M. Tamfum^{36,37}, Jean-Michel Heraud^{12,70}, Jeffrey G. Shaffer¹²⁹, Jennifer Giandhari², Jennifer Musyoki¹¹, Jessica N. Uwanibe⁴³, Jinal N. Bhiman^{13,130}, Jiro Yasuda¹¹⁵, Joana Morais^{105,131}, Joana Q. Mends^{132,133}, Jocelyn Kiconco⁸⁶, John Demby Sandi^{57,58}, John Huddleston¹³⁴, John Kofi Odoom⁶⁷, John M. Morobe¹¹, John O. Gyapong^{132,135}, John T. Kayiwa³, Johnson C. Okolie⁴³, Joicymara Santos Xavier^{1,136,137}, Jones Gyamfi¹³², Joseph Humphrey Kofi Bonney⁶⁷, Joseph Nyandwi^{138,139}, Josie Everett¹³, Jouali Farah¹⁴⁰, Joweria Nakaseegu⁸⁶, Joyce M. Ngoi⁸³, Joyce Namulondo⁸⁶, Judith U. Oguzie⁴³, Julia C. Andeko¹²⁸, Julius J. Lutwama³, Justin O'Grady⁴⁶, Katherine J Siddle⁵³, Kathleen Victoir¹⁴¹, Kayode T. Adeyemi⁴³, Kefentse A. Tumedi^{135,142}, Kevin Sanders Carvalho¹⁴³, Khadija Said Mohammed¹¹, Kunda G. Musonda¹⁴⁴, Kwabena O. Duedu¹³², Lahcen Belyamani⁶⁰, Lamia Fki-Berrajah¹¹⁶, Lavanya Singh², Leon Biscornet⁶⁸, Leonardo de Oliveira Martins⁴⁶, Lucious Chabuka¹⁴⁵, Luicer Olubayo⁸, Lul Lojok Deng¹²⁶, Lynette Isabella Ochola-Oyier¹¹, Madisa Mine¹⁴⁶, Magalutcheemee Ramuth¹²⁷, Maha Mastouri^{122,147}, Mahmoud ElHefnawi¹⁴⁸, Maimouna Mbanne¹², Maitshwarelo I. Matsheka¹⁴², Malebogo Kebabonye¹⁴⁹, Mamadou Diop¹², Mambu Momoh^{57,58,150}, Maria da Luz Lima Mendonça¹⁴³, Marietjie Venter¹⁵¹, Marietou F Paye⁵³, Martin Faye¹², Martin M. Nyaga¹⁵², Mathabo Mareka¹⁵³, Matoke-Muhia Damaris¹⁵⁴, Maureen W. Mburu¹¹, Maximillian Mpina^{155,156,157}, MFOUTOU MAPANGUY Claujens Chastel^{55,158}, Michael Owusu¹⁵⁹, Michael R. Wiley^{74,160}, Mirabeau Youtchou Tatfeng¹⁶¹, Mitoha Ondo'o Ayekaba¹⁵⁶, Mohamed Abouelhoda^{162,163}, Mohamed Amine Beloufa¹⁰², Mohamed G Seadawy¹⁶⁴, Mohamed K. Khalifa¹⁶⁵, Mohammed Koussai DELLAGI¹⁴¹, Mooko Marethabile Matobo¹⁵³, Mouhamed Kane¹², Mouna Ouadghiri⁶⁰, Mounerou Salou¹⁶⁶, Mphaphi B. Mbulawa¹⁴⁹, Mudashiru Femi Saibu⁴³, Mulenga Mwenda⁸⁵, Muluken Kaba⁹⁴, My V.T. Phan³, Nabil Abid^{122,167}, Nadia Touil¹⁶⁸, Nadine Rujeni^{169,170}, Nalia Ismael¹⁷¹, Ndeye Marieme Top¹², Ndongo Dia¹², Nédio Mabunda¹⁷¹, Nei-yuan Hsiao^{9,71}, Nelson Boricó Silochi¹⁵⁶, Ngonda Saasa¹⁷², Nicholas Bbosa³, Nickson Murunga¹¹, Nicksy Gumede¹⁹, Nicole Wolter^{13,130}, Nikita Sitharam¹, Nnaemeka Ndodo⁸⁰, Nnennaya A. Ajayi¹⁷³, Noël Tordo¹⁷⁴, Nokuzola Mbhele⁹, Norosoa H Razanajatovo⁷⁰, Nosamiefan Iguosadolo⁴³, Nwando MBA⁸⁰, Ojide C. Kingsley¹⁷⁵, Okogbenin Sylvanus⁹⁷, Okokhere Peter⁹⁷, Oladiji Femi¹⁷⁶, Olumade Testimony⁴³, Olusola Akinola Ogunsanya⁴³, Oluwatosin Fakayode¹⁷⁷, Onwe E. Ogah¹⁷⁸, Ousmane Faye¹², Pamela Smith-Lawrence¹⁴⁹, Pascale Ondo⁶⁴, Patrice Combe¹⁷⁹, Patricia Nabisubi^{180,181}, Patrick Semanda¹¹⁷, Paul E. Oluniyi⁴³, Paulo Arnaldo¹⁷¹, Peter Kojo Quashie⁸³, Philip Bejon¹¹, Philippe Dussart⁷⁰, Phillip A. Bester¹⁸², Placide K. Mbala^{36,37}, Pontiano Kaleebu^{3,86}, Priscilla Abechi⁴³, Rabeh El-Shesheny^{40,183}, Rageema Joseph⁹, Ramy Karam Aziz^{184,185}, René Ghislain Essomba^{186,187}, Reuben Ayivor-Djanie¹³², Richard Njouom¹⁸⁸, Richard O. Phillips¹⁸⁹, Richmond Gorman¹⁸⁹, Robert A. Kingsley⁴⁶, Rosemary Audu²⁸, Rosina A.A. Carr¹³², Saâd El Kabbaj¹⁹⁰, Saba Gargouri¹¹⁶, Saber Masmoudi⁴¹, Safietou Sankhe¹², Sahra Isse Mohamed¹⁹¹, Salma MHALLA¹⁹², Salome Hosch^{155,193}, Samar Kamal Kassim¹¹⁹, Samar Metha⁵³, Sameh Trabelsi¹⁹⁴, Sanaâ Lemriss¹⁹⁵, Sara Hassan Agwa¹¹⁹, Sarah Wambui Mwangi⁷, Seydou Doumbia⁵⁰, Sheila Makiala-Mandanda^{36,37}, Sherihane Aryeetey¹⁸⁹, Shymaa S. Ahmed¹⁹⁶, SIDI MOHAMED AHMED⁹², Siham Elhamoumi⁵³, Sikhulile Moyo^{89,197}, Silvia Lutucuta¹⁰⁵, Simani Gaseitsiwe^{89,197}, Simbirie Jalloh^{57,58}, Soafy Andriamandimby⁷⁰, Sobajo Oguntope⁴³, Solène Grayo¹⁷⁴, Sonia Lekana-Douki¹²⁸, Sophie Prosolek⁴⁶, Soumeya Ouangraoua^{198,199}, Stephanie van Wyk¹, Stephen F. Schaffner⁵³, Stephen Kanyerezi^{180,181}, Steve AHUKA-MUNDEKE^{36,37}, Steven Rudder⁴⁶,

Sureshnee Pillay², Susan Nabadda¹¹⁷, Sylvie Behillii²⁰⁰, Sylvie L. Budiaki¹⁵³, Sylvie van der Werf²⁰⁰, Tapfumanei Mashe^{38,201}, Tarik Aanniz⁶⁰, Thabo Mohale¹³, Thanh Le-Viet⁴⁶, Thirumalaisamy P. Velavan^{104,202}, Tobias Schindler^{155,156,193}, Tongai Maponga¹¹⁰, Trevor Bedford^{134,203}, Ugochukwu J. Anyaneji², Ugwu Chinedu⁴³, Upasana Ramphal^{2,10,204}, Vincent Enouf²⁰⁰, Vishvanath Nene⁹¹, Vivianne Gorova^{205,206}, Wael H. Roshdy¹⁹⁶, Wasim Abdul Karim¹, William K. Ampofo²⁰⁷, Wolfgang Preiser^{110,111}, Wonderful T. Choga^{89,208}, Yahaya ALI AHMED¹⁹, Yajna Ramphal¹, Yaw Bediako^{83,209}, Yeshnee Naidoo², Yvan Butera^{169,210,211}, Zaydah R. de Laurent¹¹, Ahmed E.O. Ouma⁷, Anne von Gottberg^{13,130}, George Githinji^{11,212}, Matshidiso Moeti¹⁹, Oyewale Tomori⁴³, Pardis C. Sabeti⁵³, Amadou A. Sall¹², Samuel O. Oyola⁹¹, Yenew K. Tebeje⁷, Sofonias K. Tessema⁷, Tulio de Oliveira^{1,2,10,213*}, Christian Happi⁴³, 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 & LSHTM Uganda Research Unit; Entebbe, Uganda.

⁴MRC-University of Glasgow Centre for Virus Research; Glasgow, United Kingdom.

⁵The Biotechnology Centre of the University of Yaoundé I; Yaoundé, Cameroon

⁶CDC Foundation; Atlanta, Georgia, Nebraska Department of Health and Human Services; Lincoln, Nebraska

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.

¹⁰Centre for the AIDS Programme of Research in South Africa (Cape Town); Durban, South Africa.

¹¹KEMRI-Wellcome Trust Research Programme, Kilifi Kenya

¹²Virology Department, Institut Pasteur de Dakar; Dakar, Senegal.

¹³National Institute for Communicable Diseases (NICD) of the National Health Laboratory Service (NHL); Johannesburg, South Africa.

¹⁴School of Health Sciences, College of Health Sciences, University of KwaZulu-Natal, KwaZulu-Natal, South Africa

¹⁵Department of Medical Laboratory Sciences, College of Health Sciences, Addis Ababa University, Addis Ababa, Ethiopia

¹⁶Department 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, 11796, Egypt

¹⁸Genomic Sequencing Center, Institut Pasteur du Maroc, Casablanca, Morocco

¹⁹World Health Organization, Africa Region; Brazzaville Congo

²⁰Centre d'Infectiologie Charles Mérieux-Mali (CICM-Mali); Bamako, Mali.

²¹Bacteriology and Virology Department Souro Sanou University Hospital, Bobo-Dioulasso, Burkina Faso.

²²West African Health Organisation

²³Faculty of Medicine and Health Sciences. Kassala University, Kassala city, Sudan

²⁴Department of Microbiology, Faculty of Medical Laboratory Sciences, University of Gezira, Sudan

²⁵General Administration of Laboratories and Blood Banks, Ministry of Health, Kassala State, Sudan)

²⁶MRC Unit The Gambia at LSHTM; Fajara, Gambia.

²⁷Center for Medical and Sanitary Research (CERMES)

²⁸The Nigerian Institute of Medical Research; Yaba, Lagos, Nigeria

²⁹Laboratoire de la Caisse Nationale de Sécurité Sociale, Djibouti, Republic of Djibouti

³⁰Department of Virology, College of Medicine, University of Ibadan; Ibadan, Nigeria

³¹Medical Microbiology and Parasitology Department, College of Medicine, University of Ibadan; Ibadan, Nigeria

³²Biorepository Clinical Virology Laboratory, College of Medicine, University of Ibadan Nigeria

³³Infectious Disease Institute, College of Medicine, University of Ibadan, Ibadan, Nigeria

³⁴Department of Medical Microbiology and Parasitology. Faculty of Basic Clinical Sciences. College of Health Sciences. University of Ilorin; Ilorin, Kwara State, Nigeria.

³⁵The Pirbright Institute, Woking, United Kingdom.

³⁶Pathogen Sequencing Lab, Institut National de Recherche Biomédicale (INRB); Kinshasa, Democratic Republic of the Congo.

³⁷Université de Kinshasa (UNIKIN); Kinshasa, the Democratic Republic of the Congo.

³⁸National Microbiology Reference Laboratory; Harare, Zimbabwe.

³⁹Libyan Center for Biotechnology Research, Tripoli 30313, Libya.

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 dept.,CCHE57357

⁴³African Centre of Excellence for Genomics of Infectious Diseases (ACEGID), Redeemer's University; Ede, Osun State, Nigeria.

⁴⁴Laboratório de Biologia Molecular Jean Piaget

⁴⁵University Jean Piaget in Guinea-Bissau

⁴⁶Quadram Institute Bioscience, Norwich, United Kingdom.

⁴⁷Central Public Health Laboratories, Sierra Leone

⁴⁸Centre de Recherche et de Formation en Infectiologie de Guinée (CERFIG), Université de Conakry; Conakry, Guinea

⁴⁹TransVIHMI, Institut de Recherche pour le Développement, Institut National de la Santé et de la Recherche Médicale (INSERM), Montpellier University, 34090 Montpellier, France.

⁵⁰University Clinical Research Center (UCRC), University of Sciences, Techniques and Technology of Bamako; Bamako, Mali.

⁵¹Sharjah Institute for Medical Research, College of Medicine, University of Sharjah, Sharjah, United Arab Emirates

- ⁵²Laboratoire des Fièvres Hémorragiques Virales du Benin
- ⁵³Broad Institute of Harvard and MIT; Cambridge, Massachusetts, United States of America.
- ⁵⁴Laboratory of Clinical Virology, Institut Pasteur de Tunis, Tunis, Tunisia
- ⁵⁵Fondation Congolaise pour la Recherche Médicale
- ⁵⁶Kwame Nkrumah University of Science and Technology, Department of Theoretical and Applied Biology, Kumasi, Ghana
- ⁵⁷Viral Haemorrhagic Fever Laboratory, Kenema Government Hospital, Kenema Sierra Leone
- ⁵⁸Ministry of Health and Sanitation, Freetown, Sierra Leone. Eastern Province, Sierra Leone
- ⁵⁹Department of Biological Sciences, Faculty of Natural Sciences, Redeemer's University, Ede, Osun State, Nigeria
- ⁶⁰Medical Biotechnology Laboratory, Rabat Medical and Pharmacy School, Mohammed The Vth University; Rabat, Morocco.
- ⁶¹Department of Immunology, University of Maiduguri Teaching Hospital, P.M.B. 1414; Maiduguri, Nigeria
- ⁶²Department of Medical Laboratory Science, College of Medical Sciences, University of Maiduguri, P.M.B. 1069, Maiduguri, Borno State
- ⁶³National HIV Reference Laboratory, Community Health Sciences Unit, Ministry of Health, Malawi
- ⁶⁴African Society for Laboratory Medicine; Addis Ababa, Ethiopia
- ⁶⁵National Medical and Molecular Biology Laboratory, Ministry of Health, Djibouti, Republic of Djibouti
- ⁶⁶Africa CDC, Rapid Responder, Team Djibouti
- ⁶⁷Noguchi Memorial Institute for Medical Research, University of Ghana, Legon, Ghana
- ⁶⁸Seychelles Public Health Laboratory, Public Health Authority, Ministry of Health Seychelles
- ⁶⁹Department of Ecology and Evolution; University of Chicago; Chicago, Illinois, United States of America.
- ⁷⁰Virology Unit, Institut Pasteur de Madagascar; Antananarivo, Madagascar.
- ⁷¹National Health Laboratory Service (N HLS); Cape Town, South Africa.
- ⁷²Malawi-Liverpool-Wellcome Trust Clinical Research Programme
- ⁷³Liverpool school of tropical medicine, UK
- ⁷⁴University of Nebraska Medical Center (UNMC); Omaha, Nebraska, United States of America.
- ⁷⁵Antibody Immunity Research Unit, School of Pathology, University of the Witwatersrand; Johannesburg, South Africa
- ⁷⁶CHU de Bouaké, Laboratoire / Unité de Diagnostic des Virus des Fièvres Hémorragiques et Virus Émergents; Bouaké, Côte d'Ivoire.
- ⁷⁷UNIVERSITE ALASSANE OUATTARA
- ⁷⁸School of Public Health, Pwani University; Kilifi, Kenya.
- ⁷⁹Centre for Human Virology and Genomics, Nigerian Institute of Medical Research, Yaba, Lagos, Nigeria.
- ⁸⁰Nigeria Centre for Disease Control; Abuja, Nigeria.
- ⁸¹Laboratoire des Arboviruses, Fièvres Hémorragiques virales, Virus Emergents et Zoonoses, Institut Pasteur de Bangui, Central African Republic
- ⁸²Le Laboratoire National de Biologie Clinique et de Santé Publique (LNBCSP), Bangui, Central African Republic

- ⁸³West African Centre for Cell Biology of Infectious Pathogens (WACCBIP), College of Basic and Applied Sciences, University of Ghana; Accra, Ghana.
- ⁸⁴School of Life Sciences and Zeeman Institute for Systems Biology and Infectious Disease Epidemiology Research (SBIDER), University of Warwick; Coventry, United Kingdom.
- ⁸⁵PATH, Lusaka, Zambia.
- ⁸⁶Uganda Virus Research Institute; Entebbe, Uganda.
- ⁸⁷PathCare Vermaak, Pretoria, South Africa and Division of Virology, University of the Free State, Bloemfontein, South Africa
- ⁸⁸College of Medicine and Allied Health Sciences, University of Sierra Leone
- ⁸⁹Botswana Harvard AIDS Institute Partnership & Botswana Harvard HIV Reference Laboratory; Gaborone, Botswana.
- ⁹⁰Macha Research Trust, Choma, Zambia
- ⁹¹International Livestock Research Institute (ILRI), Nairobi, Kenya
- ⁹²INRSP, Nouakchott, Mauritania
- ⁹³Faculté DE MÉDECINE DE Nouakchott
- ⁹⁴Malawi CDC
- ⁹⁵Functional genomic Platform / National Centre for Scientific and Technical Research (CNRST); Rabat, Morocco.
- ⁹⁶Rwanda National Reference Laboratory; Kigali, Rwanda.
- ⁹⁷Institute of Lassa Fever Research and Control, Irrua Specialist Teaching Hospital, Irrua, Nigeria
- ⁹⁸Robert Koch-Institute; Berlin, Germany.
- ⁹⁹G5 Evolutionary Genomics of RNA viruses, Institut Pasteur; Paris, France.
- ¹⁰⁰National Public Health Reference Laboratory-National Public Health Institute of Liberia
- ¹⁰¹Faculty of pharmacy of Monastir; Monastir, Tunisia.
- ¹⁰²National Influenza Centre, Institut Pasteur d'Algérie, Algiers; Algeria
- ¹⁰³Department of Virology, Charlotte Maxeke Johannesburg Academic Hospital, Johannesburg, South Africa
- ¹⁰⁴Institute of Tropical Medicine, Universitätsklinikum Tübingen, Germany.
- ¹⁰⁵Instituto Nacional de Investigação em Saúde (National Institute for Health Research), Luanda, 3635, Angola.
- ¹⁰⁶Ministère de Santé Publique et de la Solidarité Nationale, Chad
- ¹⁰⁷WHO Int Comoros
- ¹⁰⁸World Health Organization, Africa Region, Brazzaville Congo
- ¹⁰⁹Department of Natural Sciences, Redeemer's University, Ede, Osun State, Nigeria.
- ¹¹⁰Division of Medical Virology, Faculty of Medicine and Health Sciences, Stellenbosch University; Tygerberg, Cape Town, South Africa.
- ¹¹¹National Health Laboratory Service (N HLS); Tygerberg, Cape Town, South Africa.
- ¹¹²Institution and Department: Ministry Of Health, COVID-19 Testing Laboratory; Mbabane, Kingdom of Eswatini.
- ¹¹³Laboratory of Health Sciences and Technology, High Institute of Health Sciences, Hassan 1st University of Settat; Settat, Morocco
- ¹¹⁴Rivers State University Teaching Hospital, Port Harcourt, Nigeria

- ¹¹⁵Department of Emerging Infectious Diseases, Institute of Tropical Medicine, Nagasaki University; Nagasaki, Japan.
- ¹¹⁶CHU Habib Bourguiba, Laboratory of Microbiology, Faculty of Medicine of Sfax, University of Sfax; Sfax, Tunisia
- ¹¹⁷Central Public Health Laboratories (CPHL); Kampala, Uganda.
- ¹¹⁸Institut Pasteur de Cote d'Ivoire, Departement des Virus Epidemiques
- ¹¹⁹Faculty of Medicine Ain Shams Research institute (MASRI), Ain Shams University; Cairo, Egypt.
- ¹²⁰Doctoral School of Technical and Environmental Sciences, Department of Biology and Human Health, Chad
- ¹²¹Charles Nicolle Hospital, Laboratory of Microbiology, National Influenza Center, 1006, Tunis, Tunisia
- ¹²²Laboratory of Transmissible Diseases and Biologically Active Substances (LR99ES27), Faculty of Pharmacy, University of Monastir. Tunisia
- ¹²³College of Medicine and Allied Health Science, University of Sierra Leone
- ¹²⁴Namibia Institute of Pathology; Windhoek, Namibia
- ¹²⁵National Institute of Hygiene, Lomé, Togo
- ¹²⁶National Public Health Laboratory, Ministry of Health, Juba, Republic of South Sudan
- ¹²⁷Virology/Molecular Biology Department, Central Health Laboratory, Victoria Hospital, Ministry of Health and Wellness, Mauritius
- ¹²⁸Centre Interdisciplinaires de Recherches Medicales de Franceville (CIRMF); Franceville, Gabon.
- ¹²⁹Department of Biostatistics and Data Science, School of Public Health and Tropical Medicine, Tulane University; New Orleans, Louisiana, United States of America.
- ¹³⁰School of Pathology, Faculty of Health Science, University of the Witwatersrand, Johannesburg, South Africa
- ¹³¹Departamento de Bioquímica, Faculdade de Medicina, Universidade Agostinho Neto
- ¹³²UHAS COVID-19 Testing and Research Centre, University of Health and Allied Sciences; Ho, Ghana.
- ¹³³University of Edinburgh, Scotland
- ¹³⁴Vaccine and Infectious Disease Division, Fred Hutchinson Cancer Research Center, Seattle, WA, United States
- ¹³⁵Department of Biomedical Sciences, University of Health and Allied Sciences, PMB 31, Ho. Ghana
- ¹³⁶Universidade Federal de Minas Gerais, Belo Horizonte, Brazil
- ¹³⁷Institute of Agricultural Sciences, Universidade Federal dos Vales do Jequitinhonha e Mucuri, Unaí, Brazil
- ¹³⁸National Institute of Public Health, Burundi
- ¹³⁹Faculty of medicine, University of Burundi
- ¹⁴⁰Anoual Laboratory; Casablanca, Morocco.
- ¹⁴¹Institut Pasteur, 25-26 rue du Docteur Roux, 75015 Paris
- ¹⁴²Botswana Institute for Technology Research and Innovation; Gaborone, Botswana.
- ¹⁴³Instituto Nacional de Saúde Pública, Cape Verde
- ¹⁴⁴Zambia National Public Health Institute, Lusaka, Zambia.

- ¹⁴⁵Public Health Institute of Malawi
- ¹⁴⁶National Health laboratory; Gaborone, Botswana.
- ¹⁴⁷Laboratory of Microbiology, University Hospital of Monastir; Monastir, Tunisia.
- ¹⁴⁸Biomedical Informatics and Chemoinformatics Group, Informatics and Systems Department, National Research Centre, Cairo, Egypt
- ¹⁴⁹Ministry of health and wellness; Gaborone, Botswana.
- ¹⁵⁰Eastern Technical University of Sierra Leone, Kenema Sierra Leone
- ¹⁵¹Zoonotic Arbo and Respiratory Virus Program, Centre for Viral Zoonoses, Department of Medical Virology, University of Pretoria, Pretoria, South Africa
- ¹⁵²Next Generation Sequencing Unit and Division of Virology, Faculty of Health Sciences, University of the Free State, Bloemfontein 9300, South Africa
- ¹⁵³National Reference Laboratory Lesotho; Maseru, Lesotho
- ¹⁵⁴Centre for Biotechnology Research and Development, Kenya Medical Research Institute, Nairobi, Kenya
- ¹⁵⁵Swiss Tropical and Public Health Institute; Basel, Switzerland
- ¹⁵⁶Laboratorio de Investigaciones de Baney; Baney, Equatorial Guinea.
- ¹⁵⁷Department of Medical Diagnostics, Kwame Nkrumah University of Science and Technology, Kumasi, Ghana
- ¹⁵⁸Université Marien Ngouabi, République du Congo
- ¹⁵⁹Department of Medical Diagnostics, Kumasi Centre for Collaborative Research in Tropical Medicine, Kwame Nkrumah University of Science and Technology; Kumasi, Ghana.
- ¹⁶⁰PraesensBio, Lincoln, Nebraska, USA
- ¹⁶¹Department of Medical Laboratory Science, Niger Delta University; Bayelsa State, Nigeria
- ¹⁶²Systems and Biomedical Engineering Department, Faculty of Engineering, Cairo University, Cairo 12613, Egypt.
- ¹⁶³King Faisal Specialist Hospital and Research Center, Riyadh, Kingdom of Saudi Arabia.
- ¹⁶⁴Biological Prevention Department, Egyptian Army, Faculty of Science, Fayoum University, Fayoum; Cairo, Egypt
- ¹⁶⁵Molecular Pathology Lab Childeren Cancer Hospital 57357 Egypt
- ¹⁶⁶Laboratoire Biolim FSS/Université de Lomé -Togo
- ¹⁶⁷High Institute of Biotechnology of Monastir; University of Monastir; Rue Taher Haddad 5000, Monastir, Tunisia.
- ¹⁶⁸Genomic Center for Human Pathologies (GENOPATH), Faculty of Medicine and Pharmacy, University Mohammed V; Rabat, Morocco.
- ¹⁶⁹Rwanda National Joint Task Force COVID-19, Rwanda Biomedical Centre, Ministry of Health; Kigali, Rwanda
- ¹⁷⁰School of Health Sciences, College of Medicine and Health Sciences, University of Rwanda; Kigali, Rwanda
- ¹⁷¹Instituto Nacional de Saude (INS); Marracuene, Mozambique.
- ¹⁷²University of Zambia, School of Veterinary Medicine, Department of Disease Control; Lusaka, Zambia.
- ¹⁷³Internal Medicine Department, Alex Ekwueme Federal University Teaching Hospital; Abakaliki, Nigeria.
- ¹⁷⁴Institut Pasteur de Guinée

- ¹⁷⁵Virology Laboratory, Alex Ekwueme Federal University Teaching Hospital; Abakaliki, Nigeria.
- ¹⁷⁶Department of Epidemiology and Community Health, Faculty of Clinical Sciences. College of Health Sciences. University of Ilorin; Ilorin, Kwara State, Nigeria.
- ¹⁷⁷Department of Public Health. Ministry of Health; Ilorin, Kwara State, Nigeria.
- ¹⁷⁸Alex Ekwueme Federal University Teaching Hospital; Abakaliki, Nigeria.
- ¹⁷⁹Mayotte Hospital Center; Mayotte, France.
- ¹⁸⁰The African Center of Excellence in Bioinformatics and Data-Intensive Sciences, The Infectious Diseases Institute, Kampala, Uganda
- ¹⁸¹Immunology and Molecular Biology, Makerere University, Kampala, Uganda
- ¹⁸²Division of Virology, National Health Laboratory Service and University of the Free State; Bloemfontein, South Africa.
- ¹⁸³Infectious Hazards Preparedness, World Health Organization, Eastern Mediterranean Regional Office, Cairo, Egypt.
- ¹⁸⁴Department of Microbiology and Immunology, Faculty of Pharmacy, Cairo University, 11562 Cairo, Egypt.
- ¹⁸⁵Microbiology and Immunology Research Program, Children's Cancer Hospital Egypt 57357, 11617 Cairo, Egypt
- ¹⁸⁶National Public Health Laboratory, Ministry of Public Health of Cameroon
- ¹⁸⁷Faculty of Medicine and Biomedical Sciences, University of Yaoundé
- ¹⁸⁸Virology Service, Centre Pasteur of Cameroun; Yaounde, Cameroun.
- ¹⁸⁹Kumasi Centre for Collaborative Research in Tropical Medicine, Kwame Nkrumah University of Science and Technology; Kumasi, Ghana
- ¹⁹⁰Laboratoire de Recherche et d'Analyses Médicales de la Gendarmerie Royale; Rabat, Morocco.
- ¹⁹¹National Public Health Reference Laboratory (NPHRL), Somalia
- ¹⁹²Faculty of medicine of Monastir, University of Monastir, Tunisia
- ¹⁹³University of Basel
- ¹⁹⁴Clinical and Experimental Pharmacology Lab, LR16SP02, National Center of Pharmacovigilance, University of Tunis El Manar; Tunis, Tunisia.
- ¹⁹⁵Laboratoire de Recherche et d'Analyses Médicales de la Gendarmerie Royale; Rabat, Morocco.
- ¹⁹⁶Central Public Health Laboratories (CPHL); Cairo, Egypt.
- ¹⁹⁷Harvard T.H. Chan School of Public Health; Boston, Massachusetts, United States of America.
- ¹⁹⁸Centre MURAZ; Ouagadougou, Burkina Faso
- ¹⁹⁹National Institute of Public Health of Burkina Faso (INSP/BF); Ouagadougou, Burkina Faso
- ²⁰⁰National Reference Center for Respiratory Viruses, Molecular Genetics of RNA Viruses, UMR 3569 CNRS, Université Paris Cité, Institut Pasteur; Paris, France
- ²⁰¹World Health Organization; Harare, Zimbabwe
- ²⁰²Vietnamese-German Center for Medical Research, Hanoi, Vietnam
- ²⁰³Howard Hughes Medical Institute, Seattle, Washington, United States
- ²⁰⁴Sub-Saharan African Network For TB/HIV Research Excellence (SANTHE); Durban, South Africa
- ²⁰⁵World Health Organization, WHO Lesotho; Maseru, Lesotho

²⁰⁶Med24 Medical Centre; Ruwa, Zimbabwe.

²⁰⁷Department of Virology, Noguchi Memorial Institute for Medical Research, University of Ghana, Legon, Ghana

²⁰⁸Division of Human Genetics, Department of Pathology, University of Cape Town; Cape Town, South Africa.

²⁰⁹Yemaachi Biotech, Accra, Ghana

²¹⁰Center for Human Genetics, College of Medicine and Health Sciences, University of Rwanda; Kigali, Rwanda

²¹¹Laboratory of Human Genetics, GIGA Research Institute; Liège, Belgium

²¹²Department of Biochemistry and Biotechnology, Pwani University, Kilifi, Kenya

²¹³Department of Global Health, University of Washington; Seattle, USA.

† These authors contributed equally

*Corresponding author: Prof Tullio de Oliveira and Dr. Eduan Wilkinson, email:

tulio@sun.ac.za; ewilkinson@sun.ac.za

Abstract: Investment in Africa over the past year with regards to SARS-CoV-2 genotyping has led to a massive increase in the number of sequences, exceeding 100,000 genomes generated to track the pandemic on the continent. Our results show an increase in the number of African countries able to sequence within their own borders, coupled with a decrease in sequencing turnaround time. Findings from this genomic surveillance underscores the heterogeneous nature of the pandemic but we observe repeated dissemination of SARS-CoV-2 variants within the continent. Sustained investment for genomic surveillance in Africa is needed as the virus continues to evolve, particularly in the low vaccination landscape. These investments are very crucial for preparedness and response for future pathogen outbreaks.

One-Sentence Summary: Expanding Africa SARS-CoV-2 sequencing capacity in a fast evolving pandemic.

Introduction

What originally started as a small cluster of pneumonia cases in Wuhan, China over two years ago (1), quickly turned into a global pandemic. Coronavirus Disease 2019 (COVID-19) is the clinical manifestation of severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) infection; by March 2022 there had been over 437 million reported cases and over 5.9 million reported deaths (2). Though 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 confer some viral adaptive advantages that increase their odds of 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, two of which were first detected on the African continent (Beta and Omicron) (3, 4, 12), while two more (Alpha and Delta) (12, 13) have spread extensively on the continent in successive waves. The remaining VOC, Gamma (14), which originated in Brazil has had a limited influence in Africa with only four recorded sequenced cases.

For genomic surveillance to be useful for public health responses, sampling 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 be able 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 blindspots were identified with regards to genomic surveillance on the continent. Since then, a lot of investment has been poured into building capacity for genomic surveillance in Africa coordinated mostly by the Africa Centers for Disease Control (Africa CDC) and the WHO AFRO, but also provided by several national and international partners, resulting in an additional 90,000 sequences shared over the past year. This makes the sequencing effort for SARS-CoV-2 a phenomenal milestone making it the fastest and most concentrated sequencing efforts for any human pathogen in the history of the continent and globally. In comparison, only 12,000 whole genome influenza sequences (16) and only ~3,100 whole genome HIV sequences (17) have been shared publicly even though HIV has plagued the continent for decades.

Here we describe how the first 100,000 SARS-CoV-2 genomes from Africa have helped with understanding the epidemic on the continent, how this genomic surveillance in Africa 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.

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 heterogeneous across the continent, but most countries have experienced multiple waves of infection (18–29), with significant local and regional diversity in the first and to a lesser extent the second waves, followed by successive sweeps of the continent with Delta and Omicron VOCs (**Figure 1A**). In all regions of the continent, different lineages and VOIs evolved and co-circulated with VOCs and in some cases, contributed considerably to epidemic waves.

In North Africa (**Figure 1B, Supp Figure S1A**), B.1 lineages and Alpha dominated the pandemic in the first and second wave 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 sub-lineage dominated the epidemic in Egypt (~40% of infections) before July 2021 when it was 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 (**Figure 1C, Supp Figure S1C**), we see a similar pandemic profile with B.1 dominating the first wave, but 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, due to its mutational profile and predicted capacity for immune escape (31). However, the C.1.2 lineage did not cause many infections in the region as it was circulating at a time when Delta was dominant. In West Africa (**Figure 1D, Supp Figure 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 Omicron VOCs in successive waves. In Central Africa (**Figure 1E, Supp Figure 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 (**Figure 1F, Supp Figure 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

Finally, we directly compared the official recorded cases in Africa with the ongoing SARS-CoV-2 genomic surveillance (GISAID date of access 2022-02-15) for a crude estimation of variants' contribution to cases, with the obvious caveats that testing and reporting practices vary widely across the continent and genomic surveillance volumes have varied throughout the pandemic. We observe that Delta was responsible for an epidemic wave between May and October 2021 (**Figure 1A**) and had the greatest impact on the continent with almost 38.5% of overall infections in Africa possibly attributed to it. The second most impactful variant was Beta, responsible for an epidemic wave at the end of 2020 and beginning of 2021 (**Figure 1A**), with 15.7% 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 significance in Africa, accounting for just 4.7% of infections.

At the time of writing, the Omicron wave was still unfolding globally and in Africa and its full impact may turn out to be much bigger. However, due to increased population immunity, from SARS-CoV-2 infection and vaccination, the impact of Omicron on mortality has been less than with the other VOCs, as can be observed by the relatively low death rates in South Africa during the Omicron wave (34). The findings from mapping epidemiological numbers onto genomic surveillance data are reliable given that the sampling of genomes across Africa appears to scale proportionally to the size and timing of epidemic waves (**Supp Figure S2**).

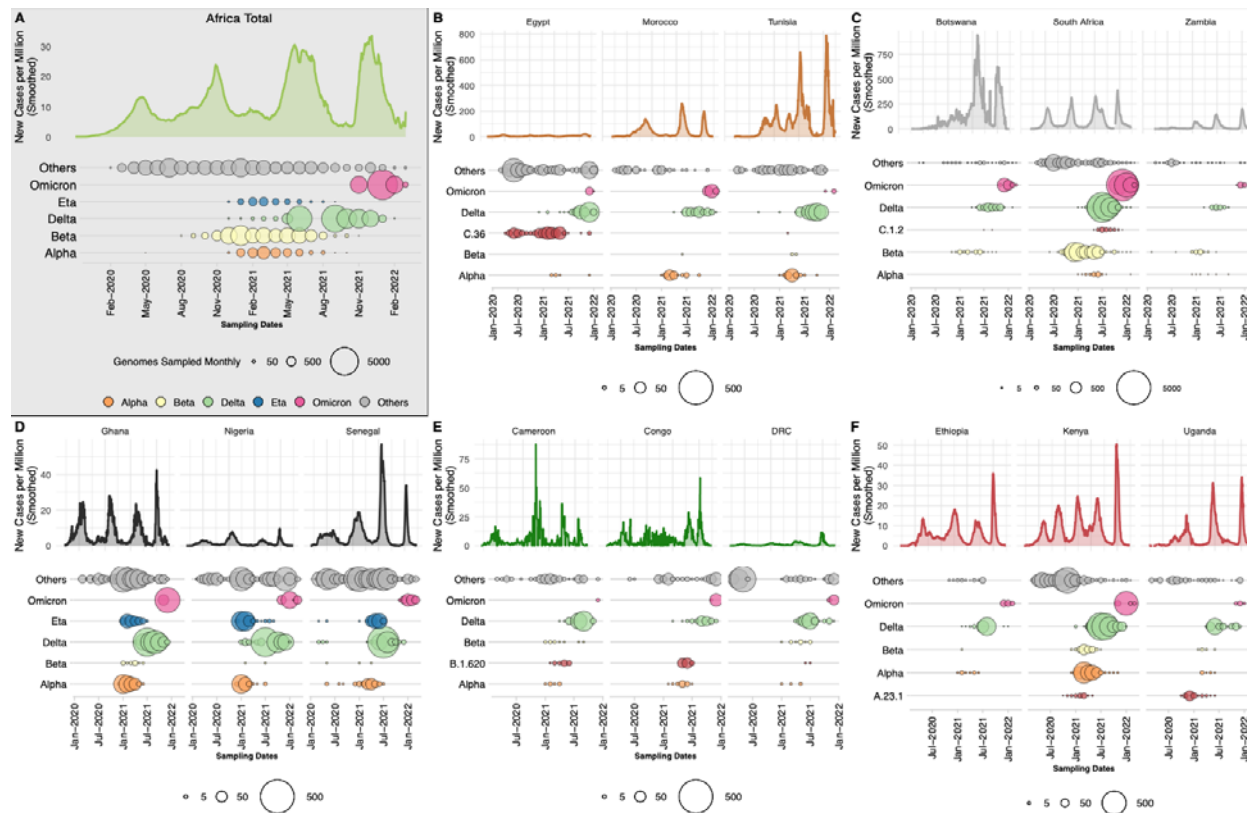


Figure 1: Epidemiological progression of the COVID-19 pandemic on the African continent. A) Total new case counts per million inhabitants in Africa along with the distribution of VOCs, the Eta VOI and other lineages through time (size of circles proportional to the number of genomes sampled per month for each category). (B-F) Breakdown of new cases per million 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). Note that the y-axis scales differ between regional sub-plots.

Regional (**Supplementary Table S1**) and country (**Supplementary Table S2**) specific Nextstrain builds for Africa also clearly support the changing nature of the pandemic over time with B.1-like viruses circulating during the first wave, and were replaced by either the Alpha or

Beta variant in the second wave, the Delta variant in the third wave and Omicron in the fourth wave.

Phylogenetic insights into the rise and spread of Variants of Concern in Africa

During the first wave of infections in 2020 in Africa, as was the case globally, the majority of corresponding genomes were classified as PANGO B.1 (n=2 456) or B.1.1 viruses (n=1 329). Towards the end of 2020, more distinct viral lineages started to appear. The most important of which that impacted the African continent are: B.1.525 (n=797), B.1.1.318 (n=398) (35), B.1.1.418 (n=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, 36), 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) (**Supp Fig S3A,B**). From our discrete state phylogeographic inference we infer at least 2,151 (95% CI: 2,132 - 2,170) viral introductions of non-VOC lineages into African countries over the course of the pandemic. While there was an initial dominance of introductions from Europe, 62% of the overall viral introductions (n=1 333; 95% CI: 1,319 - 1,347) were attributed to another African country (**Supp Fig S3C,D**). Of the 38% foreign viral introductions (n=818; 95% CI: 805 - 831) the bulk were attributed to the United Kingdom UK (~255), US (~179), and EU countries (~176). Conversely, 930 (95% CI: 919 - 941) viral exports from Africa to countries outside the continent were observed with the bulk attributed to Europe (~432), North America (~213) and Asia (~195).

As with the unfolding of events globally, VOCs became increasingly important to the epidemic in Africa towards the end of 2020. The Alpha, Beta, Delta and Omicron variants demonstrate many similarities as well as differences in the way they spread on the continent. For all these VOCs, we observe large regional monophyletic transmission clusters in each of their phylogenetic reconstructions in Africa (**Supp Fig S4**). This suggests an important extent of continental dissemination within Africa. While the Alpha and Beta variants were epidemiologically important in distinct regions of the continent, the Delta and Omicron variants sequentially dominated the majority of infections on the entire continent shortly after their emergence (**Figure 2A**).

The Alpha variant was first identified in December 2020 in the UK and has since spread globally. In Africa, Alpha has been detected in 43 countries with evidence of community transmission, based on phylogenetic clustering, in many countries including Ghana, Nigeria, Kenya, Gabon and Angola. Even in southern Africa where the Beta variant was dominating the pandemic landscape at the time, Alpha still managed to cause significant community transmission (**Supp Fig S4**). Discrete state maximum likelihood reconstruction inferred at least 155 introductions (95% CI: 152 - 159) into Africa with the bulk of imports attributed to the UK (>97%) (**Figure 2B**). However, >66% of imports into any particular African country were actually attributed to another African nation, suggesting substantial dispersal of the Alpha variant within the continent, mostly originating from West African countries (~71% of viral transfers within Africa) (**Figure 2C**). This is in spite of initial importations of the Alpha variant from Europe into all regions of the continent (**Supp Fig S5B**).

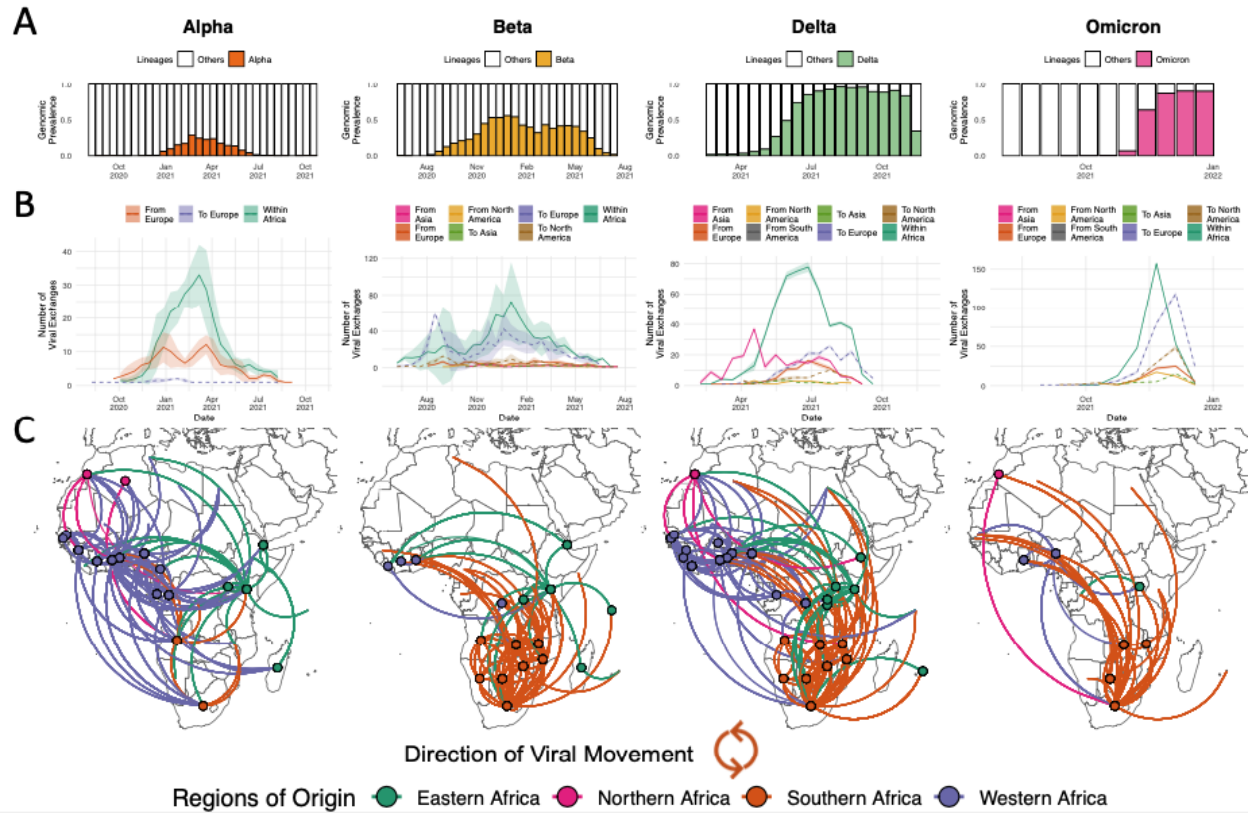


Figure 2: Inferred viral dissemination patterns of VOCs within Africa. A) Genomic prevalence of VOCs Alpha, Beta, Delta and Omicron in Africa over time. B) Inferred viral exchange patterns to, from and within the Africa continent for the four VOCs. Introductions and viral transitions within Africa are shown in solid lines and exports from Africa are shown in dotted lines and these are coloured by continent. The shaded areas around the lines represent uncertainty of this analysis from ten replicates (+/- s.d.). Note that the y-axis scales differ between subplots. C) Dissemination patterns of the VOCs within Africa, from inferred ancestral state reconstructions, annotated and coloured 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 an anti-clockwise direction.

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 in South Africa (**Supp Fig S4**). 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 substantially reduced the Beta variant's initial growth, limiting its spread to largely Southern Africa, and to a lesser extent the East Africa regions. Beta spread to at least 114 countries globally, including 37 countries and territories in Africa. Of the 900 (95% CI: 895 - 905) inferred introductions of the Beta variant into African countries, only 108 (95% CI: 105 - 111) were attributed to sources outside the continent (**Supp Fig 5C**), while nearly half of introductions were attributed to South Africa (51.3%) (**Figure 2C**). This is in line with

expectations as the variant originated in South Africa and the dominance of Alpha in most other parts of the world, which may have limited the international spread of the Beta variant (**Figure 2B**). Beyond southern Africa, most of the introductions back into the African continent were attributed to France and other EU countries into the French overseas territories, Mayotte and Reunion, and other Francophone African countries.

The fourth VOC was Delta (13), which rose to prominence in April 2021 in India, where it fueled an explosive second wave. Since its emergence, Delta was detected in >170 countries, including 37 African countries and territories (**Supp Fig S4**). Our analysis infers at least 399 (95% CI: 393 - 405) introductions of the Delta variant into Africa, with the bulk attributed to India (~52%), mainland Europe (~25%), the UK (~14%), and the US (~6%). However, as with the Alpha variant, while all regions of the continent saw a large number of introductions from Asia (**Supp Fig S5D**), the overall majority of Delta introductions into any African country were in fact from another African country (mean 67%; 95% CI: 65 - 69%) (**Figure 1B**), suggesting evidence of rapid spread within and between regions. We infer that viral dissemination of Delta within Africa was not restricted to or dominated by any particular region unlike Alpha and Beta, but rather spread across the entire continent (**Figure 1C**). Following introductions from Asia in the middle of 2021, Delta rapidly replaced the other circulating variants. For example, in southern African countries, the Delta variant rapidly displaced Beta and by June-2021 was circulating at very high (>90%) frequencies (37).

The latest VOC, Omicron, was identified and characterized in November 2021, in southern Africa (3). At the time of writing, the variant has been detected in >150 countries including 36 African countries and one overseas territory (**Supp Fig S4**). Our discrete ancestral state reconstruction infers at least 397 (95% CI: 391 - 403) introductions of the Omicron variant into various African countries, of which only 121 (95% CI: 115 - 127) were attributed to sources outside of the continent (**Figure 1B**). Similar to Beta, viral dissemination of Omicron within the continent mostly originated from southern Africa (**Figure 1C**). Outside introductions of the Omicron back into the African continent were dominated by the UK (~47%), USA (~33%), Australia and New Zealand (~9%). These reintroductions of the Omicron variant back into Africa were primarily directed to countries outside of the southern African region (e.g. Nigeria, Ghana or Kenya) where the variant was first identified (**Supp Fig 5D**). As expected, we also infer a considerable proportion of viral exportation events of Omicron from Africa to Europe (**Figure 1B**).

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 (**Figure 3A**). 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 to other sequencing technologies and better adapted to modest laboratory infrastructures, illustrates one component of how this rapid scale-up of local sequencing was achieved (**Supp Fig S6**). 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, there were 52/55 countries in Africa with SARS-CoV-2 genomes deposited in GISAID, however, there were still 16 countries with no reported local sequencing capacity (**Figure 3A**) and undoubtedly many with limited capacity for the demands during pandemic waves.

To tackle this, regional sequencing hubs were established aimed at maximizing 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 (**Figure 3B**). Sequencing is further supplemented by a number of countries utilizing 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 blindspots (**Figure 3C**). Countries in sub-Saharan Africa, particularly in Southern and East Africa, most benefited from the regional sequencing networks, while 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 frequency of sampling. African countries were positioned across a range of different combinations of overall proportion and frequency of genomic sampling (**Figure 3D**). While the ultimate goal would be to optimize both of these parameters, a lower proportion of sampling can also be useful if frequency of sampling is maintained as high as possible. For instance, South Africa and Nigeria, who have both sequenced ~1% of cases overall, can be considered to have successful genomic surveillance programs on the basis 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 that sequencing turn-around time in Africa improved (**Figure 3E**), decreasing from a mean of 171 days between October to December 2020 to a mean of 32 days over the same period a year later, although this does come with several caveats. First, we measure sequencing turn-around time in the most accessible manner, which is by comparing the date of sampling of a specimen to the date its sequence is deposited in GISAID. Overall, 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 country-to-country variation in these trends (**Supp Figure S7**), delays in data sharing, and potential retrospective sequencing, particularly by countries joining 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 last four months) may still be ongoing. The shortening duration between sampling and genomic data sharing is

nevertheless a positive takeaway, given that this data also feeds 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 (**Supp Figure S8**). As expected, we found that the route taken to sequencing matters to the speed of data generation. Local sequencing has significantly faster sequencing turn-around times of the three frameworks we investigated (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) (**Figure 3F**). These findings support the investments in local genomic surveillance, to generate timely data for local and regional decision making.

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 the variant originated in September 2020. This three-month lag in detection means that a new variant, like Beta, has ample time to spread over a large geographic region prior to 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 or HPD: 30 September - 20 October 2021) and the variant was described on 23rd November 2021 (3). Thus, Omicron was detected within ~5 weeks from origin compared to the Beta variant (~16 weeks) and the Alpha variant, detected in the UK (~10 weeks). More importantly, the time between sequence deposition to the WHO declaring the new variant a VOC was substantially shortened to 72 hours for the Omicron variant.

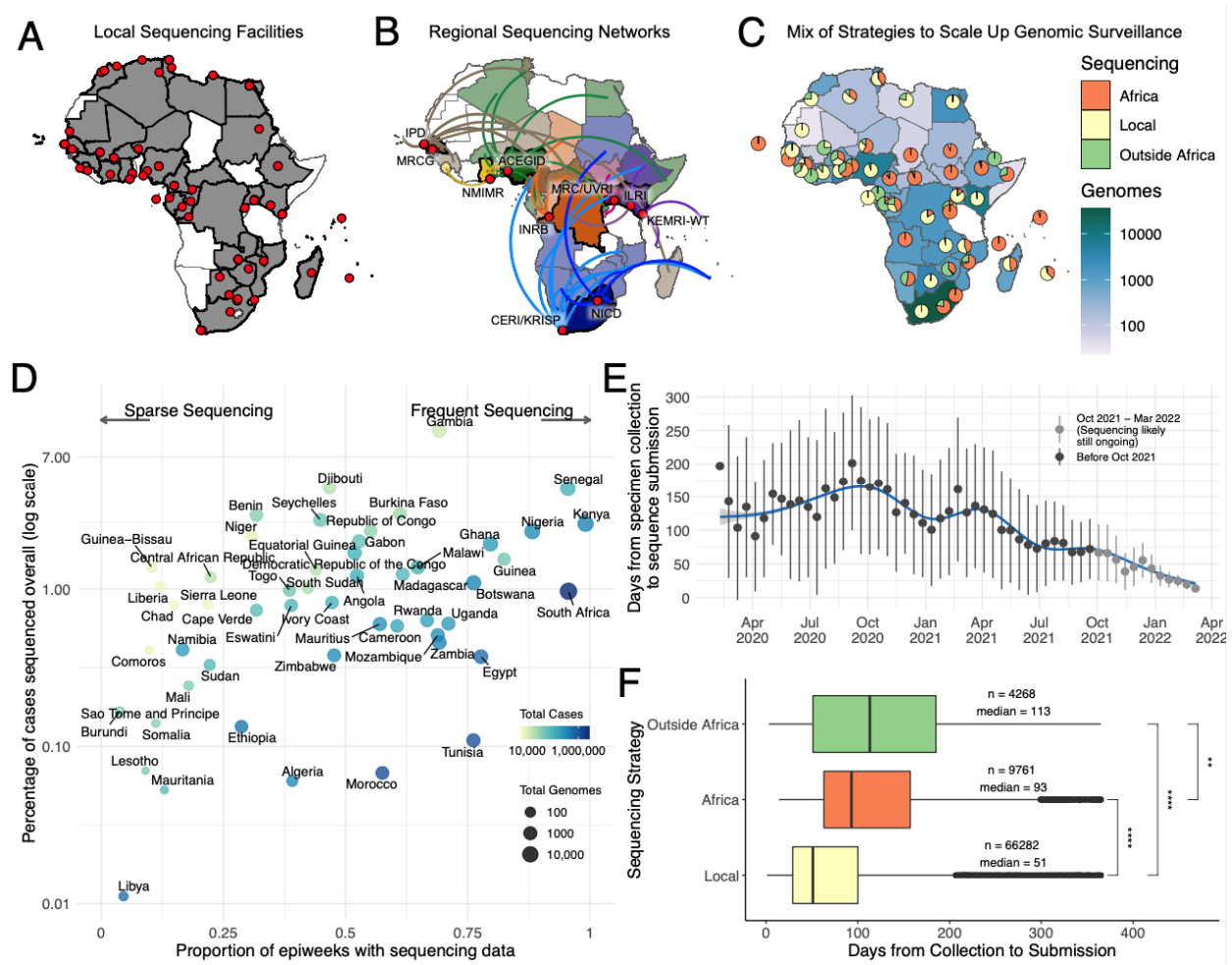


Figure 3: Sequencing strategies and outputs in Africa. A) Geographical representation of all countries (shaded in gray) and institutions (red dots) in Africa with their own on-site sequencing facilities. B) Key regional sequencing hubs and networks in Africa showing countries (shaded in bright colors) and institutions (red dots) that have sequenced for other countries (shaded in corresponding light colors and linking curves) on the continent. CERI: Centre for Epidemic Response and Innovation; KRISP: KwaZulu-Natal Research Innovation and Sequencing Platform; NICD: National Institute for Communicable Diseases; KEMRI-WT: Kenya Medical Research Institute - Wellcome Trust; ILRI: International Livestock Research Institute; MRC/UVRI: Medical Research Council/Uganda Virus Research Institute; INRB: Institut National de Recherche Biomédicale; ACEGID: African Centre of Excellence for Genomics of Infectious Diseases; NMIMR: Noguchi Memorial Institute for Medical Research; MRCG: Medical Research Council Unit - The Gambia; IPD: Institut Pasteur de Dakar 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) Progression of

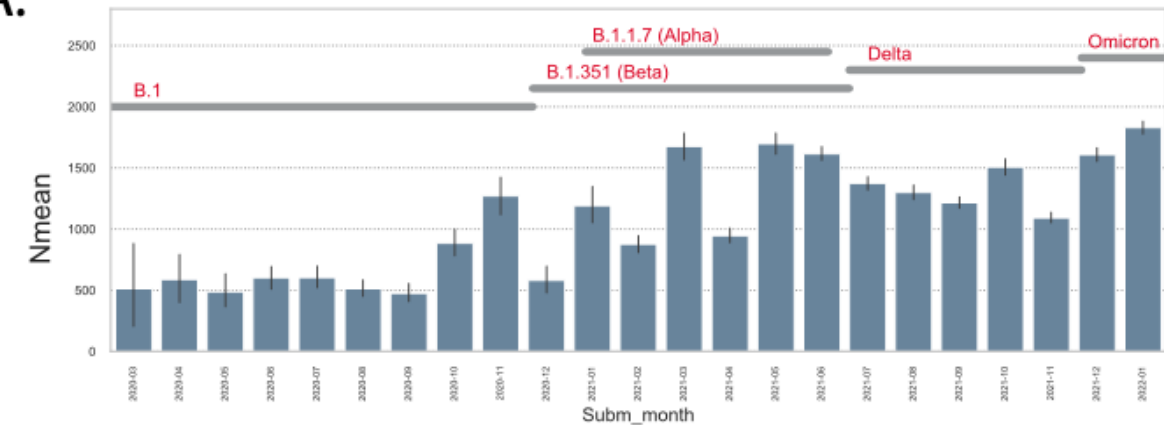
sequencing turnaround time (calculated at the difference between the date of sampling and date of sequencing submission of each sample) for the whole continent over the course of the pandemic. The dot and error bars for each group denote the mean and range for two s.d., respectively F) Comparison of sequencing turn-around times (lag times from sample collection to submission) for the three strategies of sequencing in Africa, showing a significant difference in the means (p -value<0.0001). The box and whisker plot denote the lower quartile, median and upper quartile (box), the minimum maximum values (whisker), and outliers (black dots)

Optimizing and developing sequencing protocols in Africa

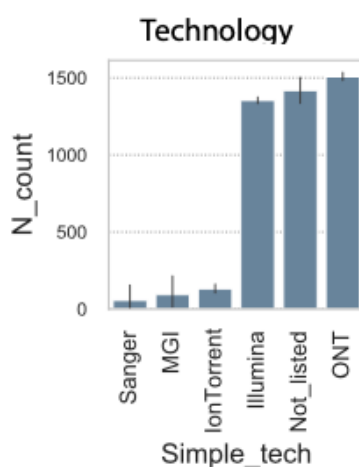
Yet, the rapidity of SARS-CoV-2 evolution has complicated sequencing efforts. Common methods of RNA sequencing include reverse transcription followed by double stranded DNA amplification using sequence-specific primer sets (38). Ongoing SARS-CoV-2 evolution has necessitated the continual evaluation and updating of these primer sets to ensure their sustained utility during genomic surveillance efforts.

In addition to increasing the volume of genomic surveillance on the continent, it is critical to also maintain sequencing quality. For the sequencing of fast-evolving pathogens like SARS-CoV-2, viral evolution is an important factor to consider in the process of designing primers. Here, we examined the current set of genomes to determine aspects of the sequencing that might be improved upon in the future. Many of the primer sets used were designed using viral sequences at the start of the pandemic and may require updating to keep pace with evolution. Indeed, the ARTIC primer sets are currently in version 4.1 (39). The Entebbe primer set was designed mid-2020 way into the first year of the epidemic and used an algorithm and design that accommodates evolution (40). The effects of viral evolution on sequencing patterns can be seen with low median N values for the first 12 months of the epidemic with an increase from October 2020 (**Figure 4A**). Additional challenges appear (indicated by increasing median N values) as the virus further evolved to adapt to humans and avoid immune responses (**Figure 4A**). Examining the role of sequencing technology, it appears that the two major technologies used (Illumina and ONT) have similar gap profiles while Ion Torrent, MGI and Sanger show reduced N content (**Figure 4B**). Likely factors for this pattern are the primers used in sequencings, with primer choice playing a key role in the presence of gaps (**Figure 4C**). The position of gaps shows an enrichment in the genome regions after position 19 000 with frequent gaps disrupting the spike coding region (**Figure 4D**).

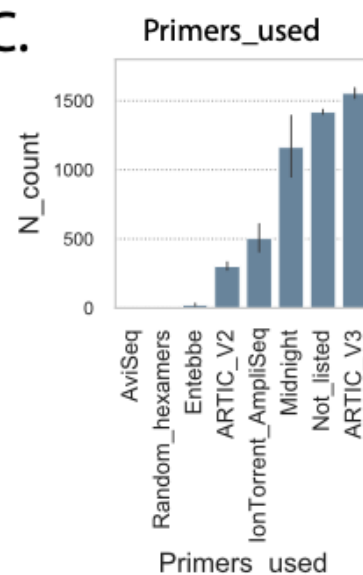
A. Mean number Ns in genomes, by Month, Technology, Primers



B.



C.



D.

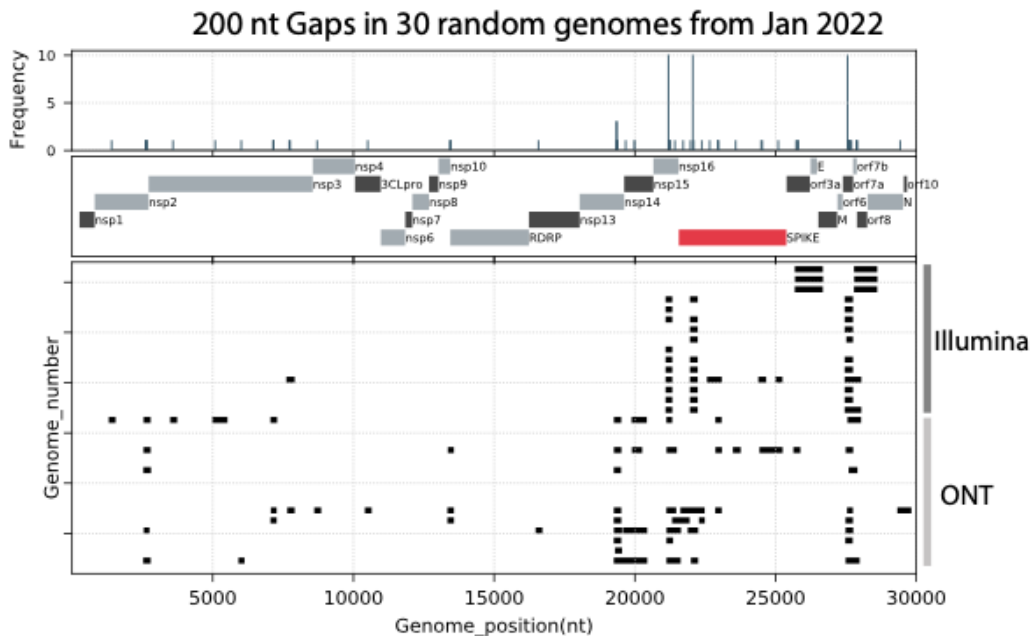


Figure 4: Genome gap analysis. A) Shows the mean N count per genome by month of submission to GISAID. The dates for the detection of important SARS-CoV-2 lineages are indicated at the top of the figure. B) Illustrates the mean N count per genome stratified by sequencing technology. C) Shows the mean N count per genome stratified by the sequencing primers sets used. For panels A to C, error bars indicate 95% confidence intervals. D) Shows the genome locations of 200nt gaps (strings of 200N) in 30 random African genomes from January 2022 (15 generated using Illumina methods, 15 with ONT). The upper histogram displays frequency of 200 base pair gaps (by start nt of gap), the middle graph shows SARS-CoV-2 genome features, bottom graph shows the positions of 200 nt gaps (black bars) with each vertical line a single genome.

Conclusions

The investment in genomic surveillance in Africa has started bearing fruit. It should be intensified as the virus continues to evolve, requiring the adaptation of methodologies locally to keep pace with the emergence of variants. The continued development of sequencing protocols in Africa is of crucial importance (40–42) given the number of variants and lineages that emerged in, and were introduced to, the continent. In Northern Africa, the SARS-CoV-2 pandemic was caused by waves of infections that were similar to those seen in Europe (first wave = B.1 descendants, second wave = Alpha, third wave = Delta and firth wave = 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 sometimes mirroring East Africa and other times southern Africa. In West Africa Eta made a significant contribution to both a second wave (together with alpha) and a third wave (together with Delta).

In spite of the recent successful expansion of genomics surveillance in Africa, additional work remains necessary. Even with the Africa CDC - Africa PGI's investment, there are still 16 countries with no sequencing capacity within their own borders. These countries' only option is to send samples to continental sequencing hubs or to centers outside of the continent, which increases the turn-around 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 which could bring substantial economic harm. Such hesitancy has obvious potential ramifications for the future of genomic surveillance on the continent.

Sampling remains heterogeneous across the continent with high relative sampling frequencies in more developed countries such as South Africa, while consistent temporally and geographically representative sampling remains elusive in most of the countries on the continent. This heterogeneous sampling creates challenges for accurate epidemic reconstruction through phylogenetic means, in particular when inferring viral dispersal patterns as uneven sampling can bias results. Unfortunately, uneven sampling will continue and can only be addressed through sensitivity analyses such as the downsampling strategies we previously

performed (15). There also exists the limitation that our genomic epidemiology understanding is influenced by the low level of testing in most if not all countries in Africa. Yet, this limitation may not be so problematic as even when countries have much higher coverage of cases/sequences (Denmark, Australia, UK), the findings may not differ substantially especially when surveillance is consistent and regular like we demonstrate happened in several African countries.

Africa needs to continue expanding genomic sequencing technologies on the continent. This holds true not just for SARS-CoV-2 but for other emerging or re-emerging pathogens on the continent. For example, WHO announced in February 2022 the re-emergence of wild polio in Africa, while sporadic influenza H1N1 and Ebola outbreaks continue to emerge on the continent. The Africa CDC has estimated that over 200 pathogen outbreaks are reported across the continent every year. Beyond the current pandemic, continued investment in these areas could serve the public health of the continent well into the 21st century.

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Author contribution

Conceptualization: HT, CB, SKT, TdO, RL, EW

Methodology: HT, TdO, RL, EW, JES, MC, BT, GM, DPM, AL, GG

Genomic Data Generation: HT, TdO, RL, EW, JES, MC, BT, GM, DPM, AL, GG, SKT, AD, DGA, MMD, AS, ANZ, AB, ASG, AKS, AO, AS, AOM, AKS, AL, AK, AEA, AAJ, AF, AMO, AOO, AAA, AJ, AMR, AK, AM, AR, AS, AK, AB, AC, AJT, AC, AKK, AK, AB, AS, AA, AVG, AJP, AY, AV, ANH, AC, AI, AM, ALB, AI, AS, AG, AF, AES, AI, BM, BLS, BSO, BB, BD, BLH, BT, BM, BN, BTM, BAK, BK, BA, BP, BM, CB, CW, CA, CBP, CS, CGA, CNA, CL, CKO, CI, CNM, CP, CEO, CDR, CMM, DJN, DBL, DJB, DM, DP, DB, DD, DS, DT, DB, DSA, DG, DSG, DOO, DM, DWW, EF, EKL, ES, EMO, ENN, EOA, EO, ES, EB, EBA, EEF, EL, EM, EC, EP, EB, ES, EAA, FL, FMT, FW, FA, FTT, FD, FVA, FR, FO, FN, FMM, FN, FEZR, FAD, FI, GKM, GT, GLK, GOA, GUE, GvZ, GAA, GS, GPM, HCR, HL, HEO, HO, HA, HK, HN, HT, HAAK, HE, HG, Hm, HK, IS, IBO, IA, IO, IBB, IS, IW, ISK, JWAH, JA, JS, JBL, JCM, JMT, JH, JGS, JG, JM, JNU, JNB, JY, JM, JQM, JK, JDS, JH, JKO, JMM, JOG, JTK, JCO, JSX, JG, JHKB, JN, JE, JF, JN, JMN, JN, JUO, JCA, JJL, JO, KJS, KV, KTA, KAT, KSC, KSM, KGM, KOD, LB, LF, LS, LB, LdOM, LC, LO, LLD, LIO, MM, MR, MM, ME, MM, MIM, MK, MD, MM, MdLLM, MV, MFP, MF, MMN, MM, MD, MWM, MM, MMCC, MO, MRW, MYT, MOA, MA, MAB, MGS, MKK, MKD, MM, MMM, MK, MO, MS, MBM, MFS, MM, MVP, NA, NT, NR, NI, NMT, ND, NM, NH, NBS, NS, NB, NM, NG, NW, NS, NN, NAA, NT, NM, NHR, NI, NM, OCK, OS, OP, OF, OT, OAO, OF, OEO, OF, PS, PO, PC, PN, PS, PEO, PA, PKQ, PB, PD, PAB, PKM, PK, PA, RE, RJ, RKA, RGE, RA, RN, ROP, RG, RAK, RA, RAC, SEK, SG, SM, SS, SIM, SM, SH, SKK, SM, ST, SL, SHA, SWM, SD, SM, SA, SSA, SMA, SE, SM, SL, SG, SJ, SA, SO, SG, SL, SP, SO, SvW, SFS, SK, SA, SR, SP, SN, SB, SLB, SvdW, TM, TA, TM, TL, TPV, TS, TM, TB, UJA, UC, UR, VE, VN, VG, WHR,

WAK, WKA, WP, WTC, YAA, YR, YB, YN, YB, ZRdL, AEO, AvG, MM, OT, PCS, AAS, SOO, YKT, CH, JN

Data Analysis: HT, TdO, RL, EW, JES, MC, BT, GM, DPM, AL, GG, MM, SvW

Funding acquisition: TdO, GG, SKT, AEO, AvG, MM, OT, AAS, SOO, YKT, CH

Project administration: TdO, GG, SKT, AEO, AvG, MM, OT, AAS, SOO, YKT, CH, RL, EW, PCS, JN, GM, AD, DGA, MMD, AC, DWW, HO, SWM

Supervision: TdO, GG, SKT, AEO, AvG, MM, OT, AAS, SOO, YKT, CH, RL, EW, PCS, JN

Writing – original draft: TdO, SKT, RL, EW, GM, HT, JES, MC, DPM, CB

Writing – review & editing: TdO, SKT, RL, EW, GM, HT, JES, MC, DPM, CB, GG, AEO, AvG, MM, OT, AAS, SOO, YKT, CH, PCS, JN, AD, DGA, MMD, AC, DWW, HO, SWM, BT, AL, AS, ANZ, AB, ASG, AKS, AO, AS, AOM, AKS, AL, AK, AEA, AAJ, AF, AMO, AOO, AAA, AJ, AMR, AK, AM, AR, AS, AK, AB, AJT, AC, AKK, AK, AB, AS, AA, AVG, AJP, AY, AV, ANH, AC, AI, AM, ALB, AI, AS, AG, AF, AES, AI, BM, BLS, BSO, BB, BD, BLH, BT, BM, BN, BTM, BAK, BK, BA, BP, BM, CB, CW, CA, CBP, CS, CGA, CNA, CL, CKO, CI, CNM, CP, CEO, CDR, CMM, DJN, DBL, DJB, DM, DP, DB, DD, DS, DT, DB, DSA, DG, DSG, DOO, DM, EF, EKL, ES, EMO, ENN, EOA, EO, ES, EB, EBA, EEF, EL, EM, EC, EP, EB, ES, EAA, FL, FMT, FW, FA, FTT, FD, FVA, FR, FO, FN, FMM, FN, FEZR, FAD, FI, GKM, GT, GLK, GOA, GUE, GvZ, GAA, GS, GPM, HCR, HL, HEO, HA, HK, HN, HT, HAAK, HE, HG, Hm, HK, IS, IBO, IA, IO, IBB, IS, IW, ISK, JWAH, JA, JS, JBL, JCM, JMT, JH, JGS, JG, JM, JNU, JNB, JY, JM, JQM, JK, JDS, JH, JKO, JMM, JOG, JTK, JCO, JSX, JG, JHKB, JN, JE, JF, JN, JMN, JN, JUO, JCA, JJL, JO, KJS, KV, KTA, KAT, KSC, KSM, KGM, KOD, LB, LF, LS, LB, LdOM, LC, LO, LLD, LIO, MM, MR, MM, ME, MM, MIM, MK, MD, MM, MdLLM, MV, MFP, MF, MMN, MM, MD, MWM, MM, MMCC, MO, MRW, MYT, MOA, MA, MAB, MGS, MKK, MKD, MM, MMM, MK, MO, MS, MBM, MFS, MM, MVP, NA, NT, NR, NI, NMT, ND, NM, NH, NBS, NS, NB, NM, NG, NW, NS, NN, NAA, NT, NM, NHR, NI, NM, OCK, OS, OP, OF, OT, OAO, OF, OEO, OF, PS, PO, PC, PN, PS, PEO, PA, PKQ, PB, PD, PAB, PKM, PK, PA, RE, RJ, RKA, RGE, RA, RN, ROP, RG, RAK, RA, RAC, SEK, SG, SM, SS, SIM, SM, SH, SKK, SM, ST, SL, SHA, SD, SM, SA, SSA, SMA, SE, SM, SL, SG, SJ, SA, SO, SG, SL, SP, SO, SvW, SFS, SK, SA, SR, SP, SN, SB, SLB, SvdW, TM, TA, TM, TL, TPV, TS, TM, TB, UJA, UC, UR, VE, VN, VG, WHR, WAK, WKA, WP, WTC, YAA, YR, YB, YN, YB, ZRdL

Competing interests: With the exception of Pardis Sabeti 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, we the authors have no conflicts of interest to declare.

Data and Code 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. A full list of the African sequences as well as global references are presented and acknowledged in Supplementary Table S3 and in

our data repository, which will be made public at final publication. The github repo also contains all of the alignments, raw and time scaled ML tree topologies, data analysis and visualization scripts used here which will allow for the independent reproduction of results.

Figure legends

Supplementary Materials

Methods and Methods

Ethics statement

This project relied on sequence data and associated metadata 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 (IRB) references or material transfer agreements (MTAs) for countries are listed below.

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 March 22, 2020), Kenya - samples were collected under the Ministry of Health protocols as part of the national COVID-19 public health response. 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 were 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. 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 were approved by the Uganda Virus Research Institute - Research and Ethics Committee UVRI-REC Federalwide Assurance [FWA] 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 15 February 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 **Figure 3** was ‘new cases per million (smoothed)’. To calculate the genomic sampling proportion and frequency for each

country for **Figure 1**, the total number of recorded cases at 15 February 2022 was considered, as well as the total length of time for which each country has recorded cases of SARS-CoV-2.

Genomic metadata was downloaded for all African entries on GISAID for the same time period (date of access: 15 February 2022). From this, information extracted from all entries for this study included: date of sampling, country of sampling, viral lineage and clade, originating laboratory, sequencing laboratory, 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 elapsed from specimen collection to sequence submission to GISAID. Sequencing technology information for all African entries was also downloaded from GISAID on 15 February 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 28 January 2022 yielding 74 785 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 74 785 entries (~19%). The total N per genome were counted, results from which were then used for genome quality analysis and visualization. Gap locations in the genomes were mapped and visualized compared to the original Wuhan strain (43).

Phylogenetic investigation

All African sequences on the GISAID sequence database (16) were downloaded on the 28th of October 2021 (n=48 145). Prior to any phylogenetic inference we performed some quality assessment on the sequences to exclude incomplete or problematic sequences. Briefly, all African sequences were passed through the NextClade analysis pipeline (44) in order to identify and exclude: (i) sequences missing >10% of the SARS-CoV-2 genome (n=6 495), (ii) sequences that deviate by >70 nucleotides from the Wuhan reference strain (n=12), (iii) sequences with >10 ambiguous bases (n=165), (iv) clustered mutations (n=99), and (v) sequences flagged as problematic by NextClade (n=544). Additionally, 492 sequences were removed due to a lack of adequate metadata (i.e. exact sampling date). This produced a final African dataset of 37,787 high quality genotypes for analysis. Due to the sheer size of the dataset we opted to perform independent phylogenetic inferences on the main VOCs (Alpha n=2 365; Beta n=10 809; Delta n=13 911) that have spread on the African continent, as well as a separate inference for all non-VOC SARS-CoV-2 sequences (n=13 252). Due to the discovery of Omicron in November 2021 an additional Omicron dataset was also constructed following the exact quality control measures (n=3 371; 659 did not pass QC; date of access 10th January 2022).

In order 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. Due to the oversampling of some variants or lineages we performed a random downsampling while retaining the oldest two known variants from each country. In total, 9 335 Alpha, 9 077 Beta, 9 999 Delta, 3 941 Omicron and 16 231 non-VOCs global references were respectively aligned with their African counterparts independently with NextAlign (44). Each of the alignments were then used to infer maximum likelihood (ML) tree topologies in FastTree v 2.0 (45) using the General Time Reversible (GTR) model of nucleotide substitution and a total of 100 bootstrap replicates (46). The resulting ML tree topologies were first inspected in TempEst (47) to identify any sequences that deviate more than 0.0001 from the residual mean. Following the removal of potential outliers in R with the ape package (48), the resulting ML-trees were then transformed into time calibrated phylogenies in TreeTime (49) by applying a rate of 8x10⁻⁴ substitution per site per year (50) in order to transform the branches into units of calendar time. Time calibrated trees were then visualized along with associated metadata in R using ggtree (51) and other packages.

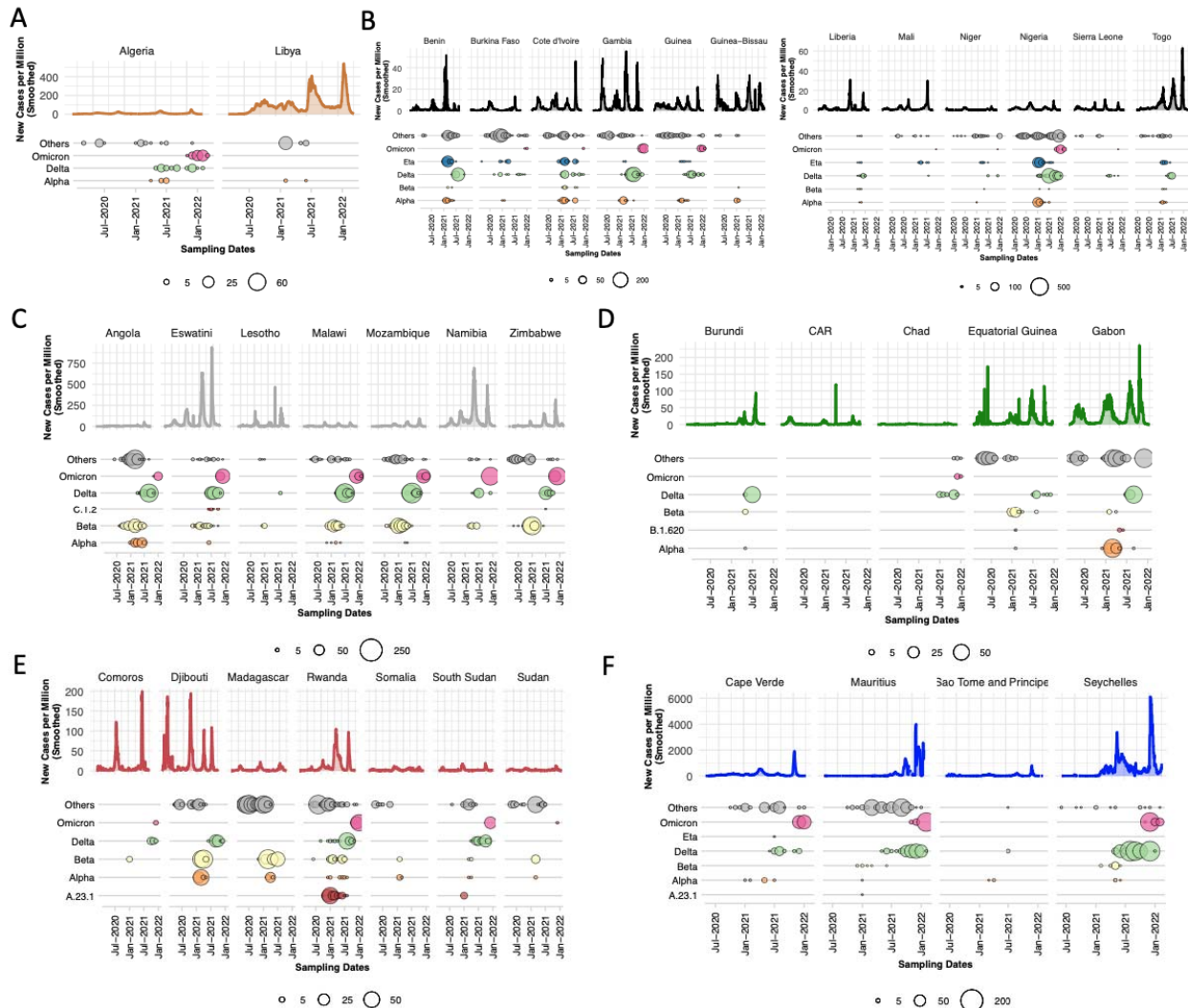
We performed a basic viral dispersal analysis for each of the VOCs (excluding Gamma), as well as for the non-VOC dataset. Briefly, a migration model was fitted to each of the time calibrated tree topologies in TreeTime, mapping the country location of sampled sequences to the external tips of the trees. The *mugration* model of TreeTime also infer 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 serve as the estimated import or export event. To infer some confidence around these estimates, we performed ten replicates for each of the dataset by random selection from the 100 bootstrap trees. 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). In a previous analysis (15) we performed a sensitivity analysis to address some of these issues and found no substantial variations in estimates.

All data analytics were performed using custom python and R scripts and results visualized using the ggplot libraries (52).

Regional and country specific Nextstrain builds

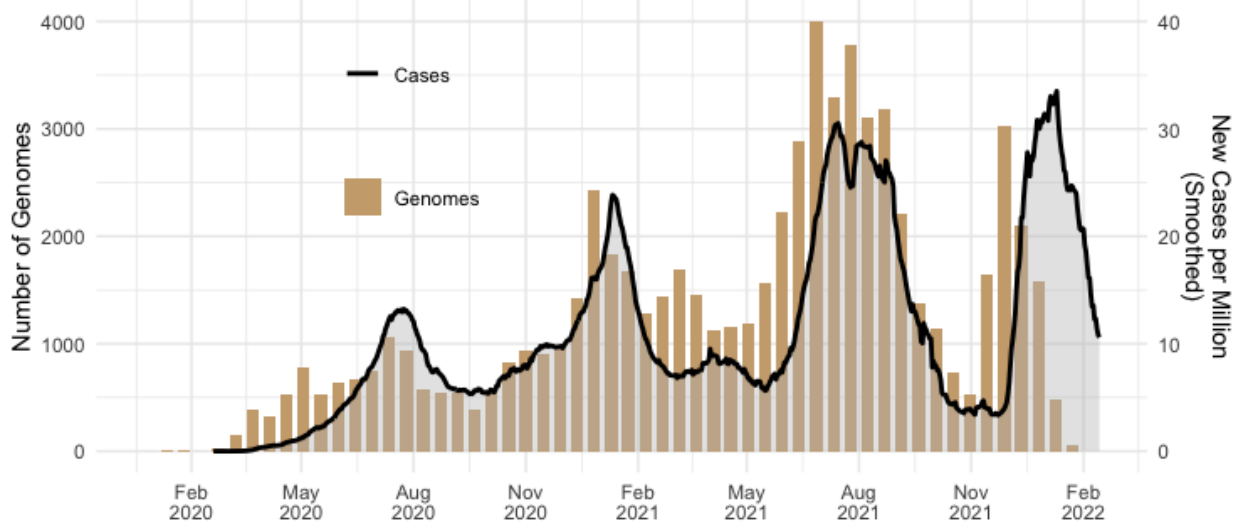
The Nextstrain pipeline (<https://github.com/nextstrain/ncov>) was used to generate the regional and country-specific builds (53). Sequencing data from the rest of the world were included for phylogenetic context based on genomic proximity and even sampling over time. For computational feasibility, ease of interpretation and to balance heterogeneous sampling efforts between countries, genomes were subsampled across time and geography resulting in final builds of ~6000 genomes for each build.

Supplementary Figures

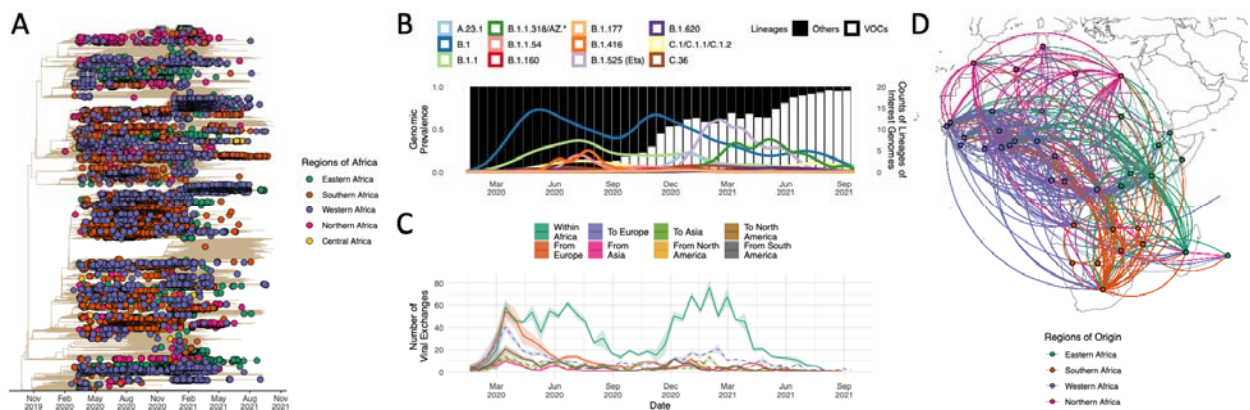


Supplementary Figure S1: Epidemiological progression of the COVID-19 pandemic in all African countries overlaid with the distribution of VOCs, the Eta VOI and other lineages through time (size of circles proportional to the number of genomes sampled per month for each category). The graphs show a breakdown of new cases per million and monthly sampling of VOCs, regional variant or lineage of interest and other lineages for all African countries not shown in Figure 1, grouped by region: A) North Africa, B) West Africa, C) Southern Africa, D) Central Africa, E) East Africa, F) Cape Verde, Mauritius, Sao Tome and Principe and Seychelles, from the beginning of the pandemic to February 2022.

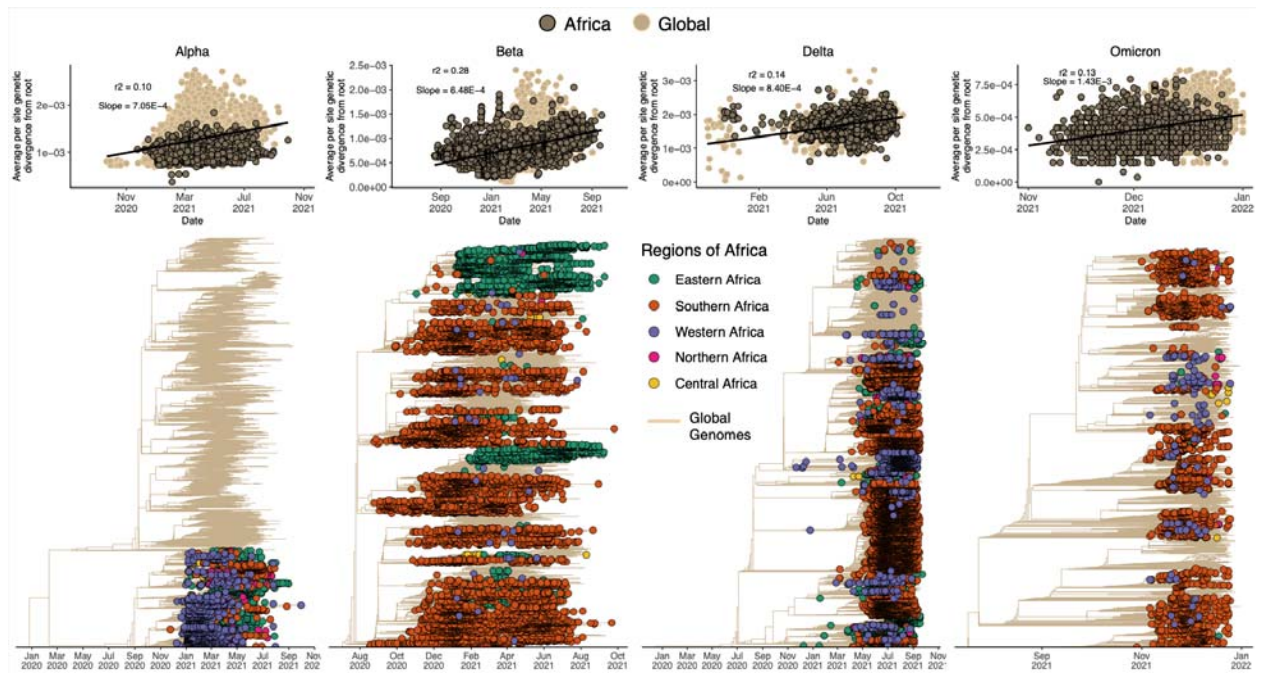
Overall Production of Sequences vs Epidemic in Africa



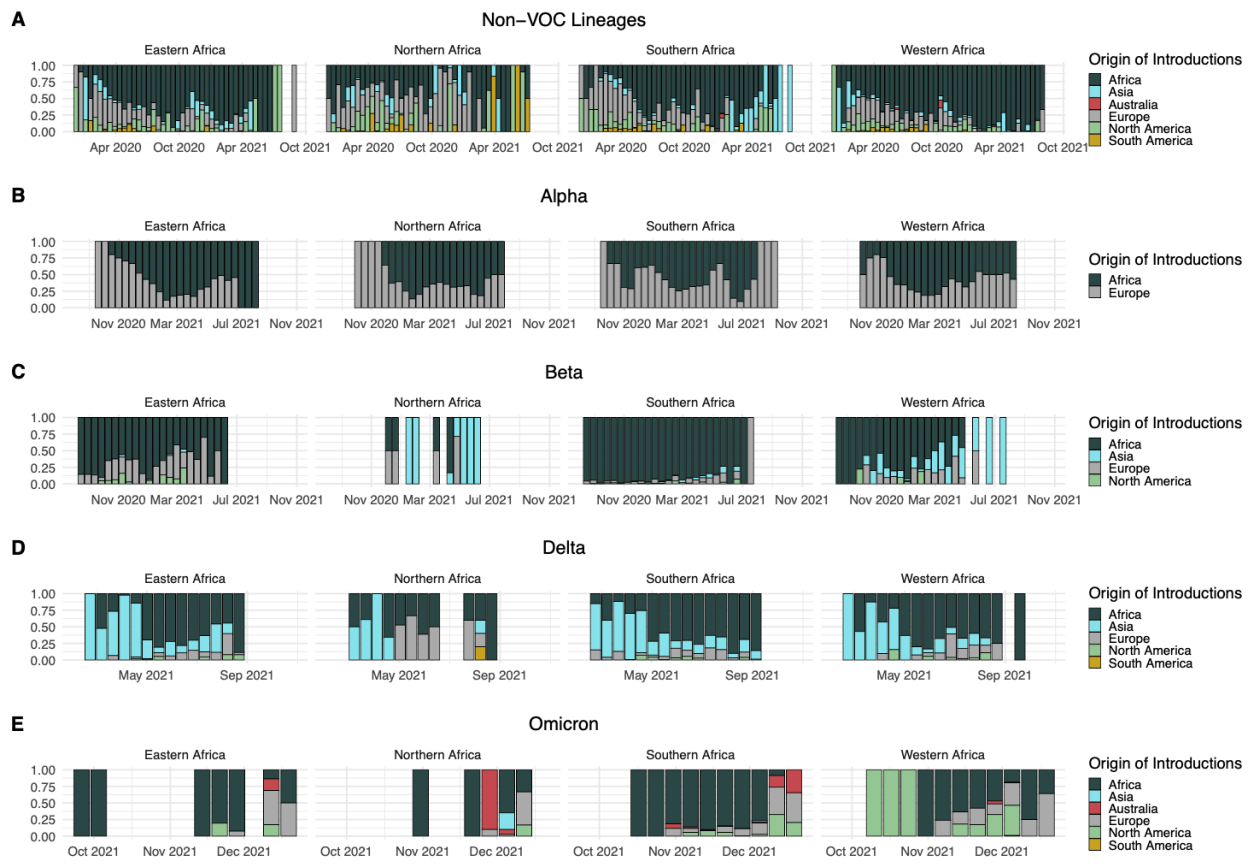
Supplementary Figure S2: Corresponding progression of genomic sequence production and epidemic size in Africa.



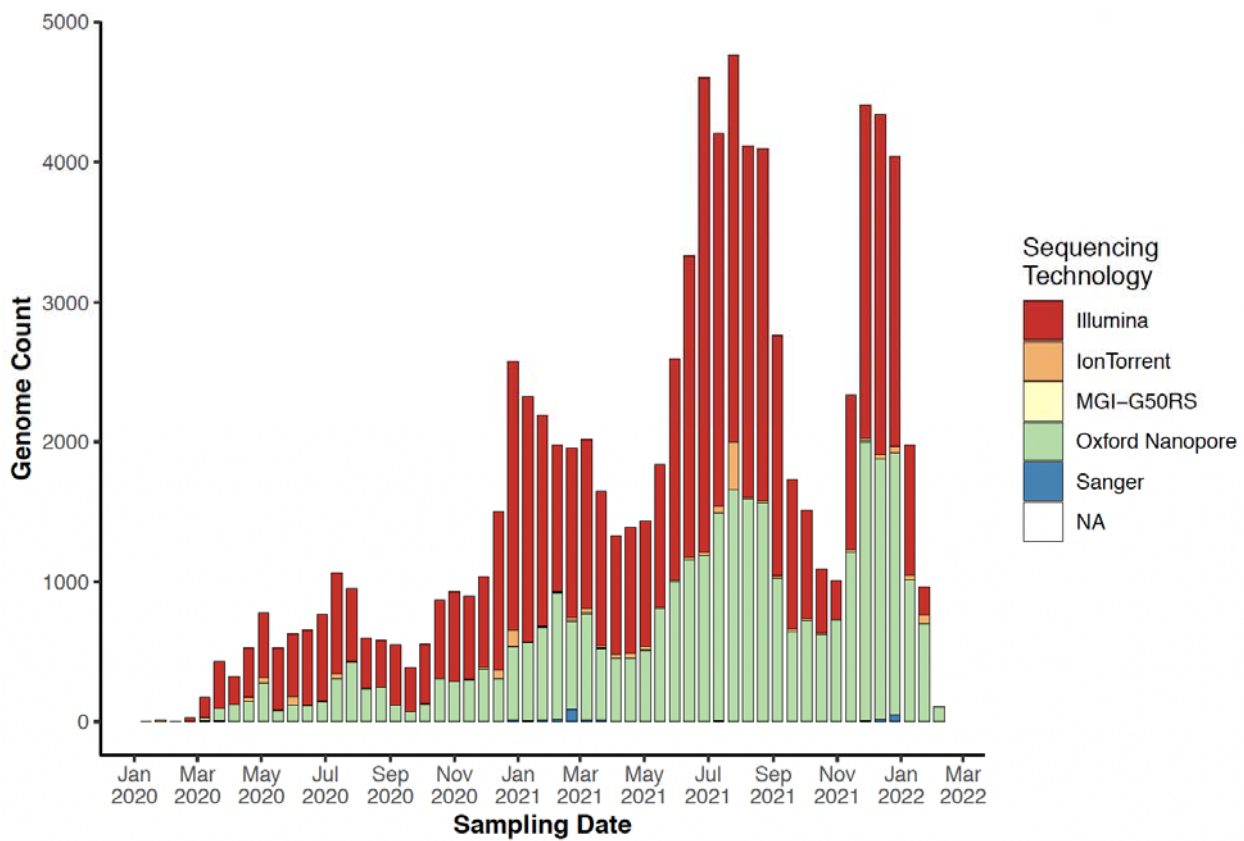
Supplementary Figure S3: Phylogenetic inference of non-VOC lineages in Africa. A) Maximum-Likelihood timetrees of non-VOC genomes in Africa from the beginning of the pandemic till October 2021 against a global reference with African genomes denoted by tippoint circles coloured by regions of Africa. B) Genomic prevalence of non-VOC (black) vs VOC (white) lineages in Africa overlaid by frequency progressions of some lineages of interest in Africa. C) *Inferred viral dissemination patterns of non-VOC lineages to, from and within the Africa continent from the beginning of the pandemic to October 2021. Introductions and viral transitions within Africa are shown in solid lines and exports from Africa are shown in dotted lines and these are coloured by continent. The shaded areas around the lines represent uncertainty of this analysis from ten replicates.* D) *Dissemination patterns of the non-VOC lineages within Africa, from inferred ancestral state reconstructions, annotated and coloured 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 an anti-clockwise direction.*



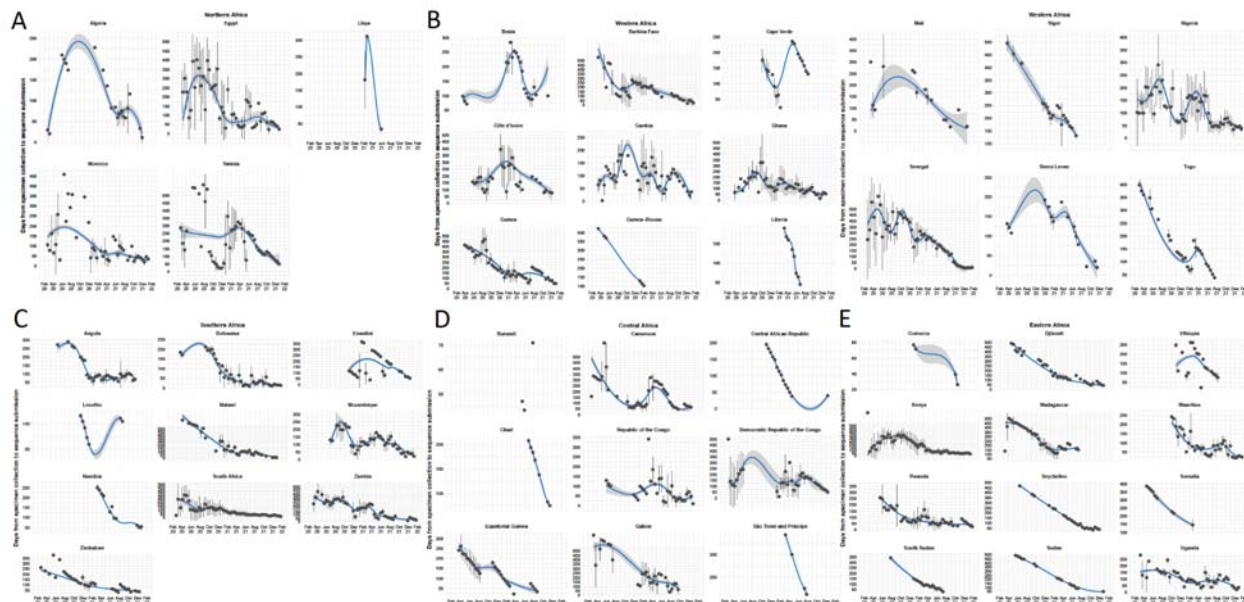
Supplementary Figure S4: Phylogenetic inference of VOCs in Africa. Top - Molecular clock evolution. Bottom - VOC-specific Maximum-Likelihood timetrees with African genomes denoted by tippoints coloured by regions of Africa.



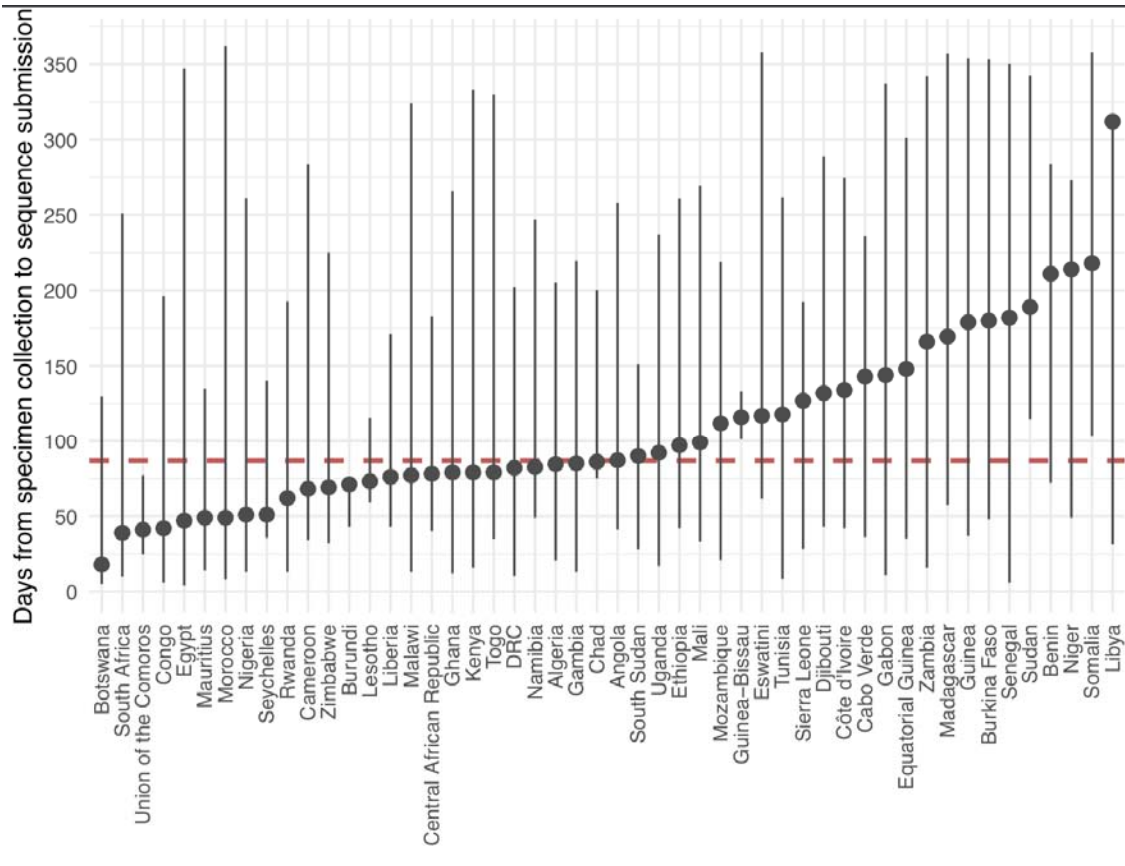
Supplementary Figure S5: Patterns of viral importations into different regions of Africa showing the proportions of introductions into Eastern, Northern, Southern and Western Africa attributed to specified origins for A) non-VOC lineages, B) Alpha VOC, C) Beta VOC, D) Delta VOC, and E) Omicron VOC during the relevant time periods.



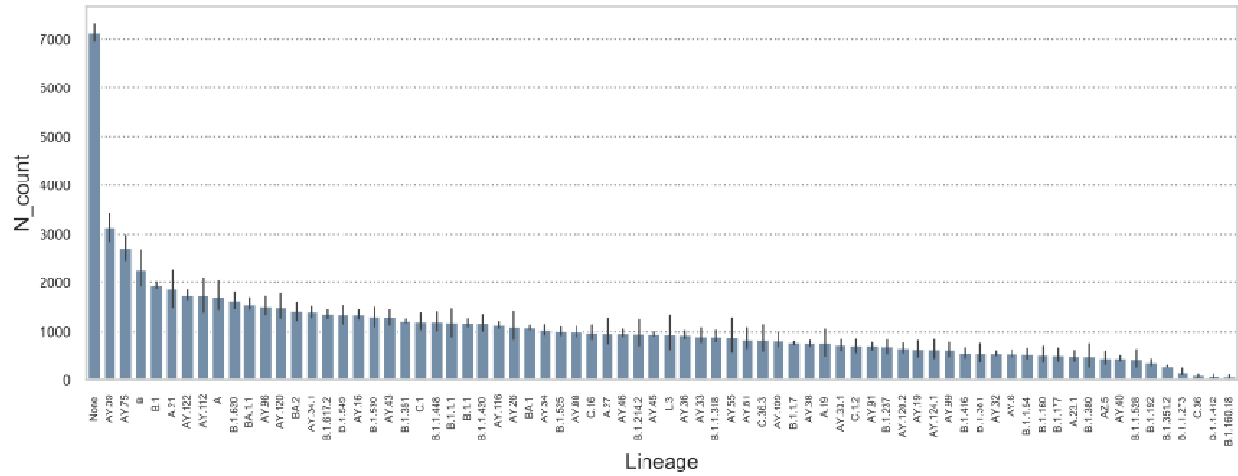
Supplementary Figure S6: The progression of sequencing technologies used for SARS-CoV-2 sequencing in Africa



Supplementary Figure S7: Sequencing turnaround time progression per country in each region of Africa in A) North Africa, B) West Africa, C) Southern Africa, D) Central Africa, and E) East Africa



Supplementary Figure S8: Sequencing turnaround time average by country. The circles indicate the mean number of days between specimen collection and sequence submission to GISAID and the bars indicate the distribution days.



Supplementary Figure S9. Gapped genomes by lineage. The mean N data were stratified by SARS-CoV-2 lineages to investigate if there were lineage-specific patterns in the genome gaps, an indirect measure of primer mismatch. For lineages present in at least 100 times in the Africa 75K genome data, there was a modest difference in N content across the lineages. Lineages that returned no classification with Pangolin ("None") showed the highest N

content, which is not surprising as the high N content was probably the basis for the failed classification. The more recent lineages Delta (e.g. AY.39, AY.75) and Omicron (BA.1.1, BA.2) also showed higher N content. There were however a number sequences classified into more basal lineages (e.g. B, B.1) and it is possible that the basal lineage accumulated genome sequences that, because of their N content, failed to return a more specific classification.

Supplementary Table S1: NextStrain builds of the five major geographical regions in Africa. Each build focuses on a specific region within Africa and includes sequences from outside the region and the continent to place the regional sequences into context of the global pandemic. All the data used in the builds are publicly available on GISAID and are maintained and updated by the Africa CDC in collaboration with the NextStrain team on a weekly basis.

Region	Regional Builds
Central African Region	https://nextstrain.org/groups/africa-cdc/ncov/central-africa
Eastern African Region	https://nextstrain.org/groups/africa-cdc/ncov/eastern-africa
Northern African Region	https://nextstrain.org/groups/africa-cdc/ncov/northern-africa
Southern African Region	https://nextstrain.org/groups/africa-cdc/ncov/southern-africa
Western African Region	https://nextstrain.org/groups/africa-cdc/ncov/western-africa

Supplementary Table S2: Individual country specific NextStrain builds from the African continent. Each build focuses on a specific country within Africa and includes sequences from the rest of the world in order to place the country's sequences into context of the global pandemic. All the data used in the builds are publicly available on GISAID and are maintained and updated by the Africa CDC in collaboration with the NextStrain team on a weekly basis.

Region	Country	Build
Central African Region	Burundi	https://nextstrain.org/groups/africa-cdc/ncov/burundi
	Cameroon	https://nextstrain.org/groups/africa-cdc/ncov/cameroon
	Central African Republic	https://nextstrain.org/groups/africa-cdc/ncov/central-african-republic
	Chad	https://nextstrain.org/groups/africa-cdc/ncov/chad
	Republic of Congo	https://nextstrain.org/groups/africa-cdc/ncov/republic-of-the-congo

	Democratic Republic of Congo	https://nextstrain.org/groups/africa-cdc/ncov/democratic-republic-of-the-congo
	Equatorial Guinea	https://nextstrain.org/groups/africa-cdc/ncov/equatorial-guinea
	Gabon	https://nextstrain.org/groups/africa-cdc/ncov/gabon
Eastern African Region	Comoros	https://nextstrain.org/groups/africa-cdc/ncov/comoros
	Djibouti	https://nextstrain.org/groups/africa-cdc/ncov/djibouti
	Ethiopia	https://nextstrain.org/groups/africa-cdc/ncov/ethiopia
	Kenya	https://nextstrain.org/groups/africa-cdc/ncov/kenya
	Madagascar	https://nextstrain.org/groups/africa-cdc/ncov/madagascar
	Mauritius	https://nextstrain.org/groups/africa-cdc/ncov/mauritius
	Rwanda	https://nextstrain.org/groups/africa-cdc/ncov/rwanda
	Seychelles	https://nextstrain.org/groups/africa-cdc/ncov/seychelles
	Somalia	https://nextstrain.org/groups/africa-cdc/ncov/somalia
	South Sudan	https://nextstrain.org/groups/africa-cdc/ncov/south-sudan
Northern African Region	Sudan	https://nextstrain.org/groups/africa-cdc/ncov/sudan
	Uganda	https://nextstrain.org/groups/africa-cdc/ncov/uganda
	Algeria	https://nextstrain.org/groups/africa-cdc/ncov/algeria
	Egypt	https://nextstrain.org/groups/africa-cdc/ncov/egypt
	Libya	https://nextstrain.org/groups/africa-cdc/ncov/libya
	Morocco	https://nextstrain.org/groups/africa-cdc/ncov/morocco

	Tunisia	https://nextstrain.org/groups/africa-cdc/ncov/tunisia
Southern African Region	Angola	https://nextstrain.org/groups/africa-cdc/ncov/angola
	Botswana	https://nextstrain.org/groups/africa-cdc/ncov/botswana
	Eswatini	https://nextstrain.org/groups/africa-cdc/ncov/eswatini
	Lesotho	https://nextstrain.org/groups/africa-cdc/ncov/lesotho
	Malawi	https://nextstrain.org/groups/africa-cdc/ncov/malawi
	Mozambique	https://nextstrain.org/groups/africa-cdc/ncov/mozambique
	Namibia	https://nextstrain.org/groups/africa-cdc/ncov/namibia
	South Africa	https://nextstrain.org/groups/africa-cdc/ncov/south-africa
	Zambia	https://nextstrain.org/groups/africa-cdc/ncov/zambia
	Zimbabwe	https://nextstrain.org/groups/africa-cdc/ncov/zimbabwe
Western African Region	Benin	https://nextstrain.org/groups/africa-cdc/ncov/benin
	Burkina Faso	https://nextstrain.org/groups/africa-cdc/ncov/burkina-faso
	Cabo Verde	https://nextstrain.org/groups/africa-cdc/ncov/cabo-verde
	Côte d'Ivoire	https://nextstrain.org/groups/africa-cdc/ncov/cote-divoire
	Gambia	https://nextstrain.org/groups/africa-cdc/ncov/gambia
	Ghana	https://nextstrain.org/groups/africa-cdc/ncov/ghana
	Guinea	https://nextstrain.org/groups/africa-cdc/ncov/guinea
	Guinea-Bissau	https://nextstrain.org/groups/africa-cdc/ncov/guinea-bissau

Liberia	https://nextstrain.org/groups/africa-cdc/ncov/liberia
Mali	https://nextstrain.org/groups/africa-cdc/ncov/mali
Niger	https://nextstrain.org/groups/africa-cdc/ncov/niger
Nigeria	https://nextstrain.org/groups/africa-cdc/ncov/nigeria
Senegal	https://nextstrain.org/groups/africa-cdc/ncov/senegal
Sierra Leone	https://nextstrain.org/groups/africa-cdc/ncov/sierra-leone
Togo	https://nextstrain.org/groups/africa-cdc/ncov/togo

Supplementary Author List

Aaron L. Shibemba, Abasi Ene Obong, Abayomi FADEYI, Abbad Anas, Abd Elazeez Shabaan, Abd Monaem Adel, Abd Moniem Ain Shoka, Abdelhamid W., Abdelilah Laraqui, Abdelkader Laatiris, Abdelkrim Meziane Bellefquih, Abdellah Faouzi, Abdelmoula F., Abdelomunim Essabbar, Abderrahmane Bimouhen, Abderraouf Hilali, Abdo I., Abdou Padane, Abdoul Karim Sangaré, Abdoul Karim Soumah, Abdoulaye Djimde, Abdoulaye Toure, Abdoulie Kanteh, Abdulla Bashein, Abdullah Salama, Abe G. Abias, Abebe Genetu Bayih, Abel Abera Negash, Abel Lissom, Abid N., Abla A. Konou, Abo Shama, Abosede O., Abouelnaga S., Abraham Ali, Abraham Kwabena Anang, Abraham Tesfaye, Adam K. Khan, Adamu Tayachew, Adane Mihret, Adba Alfatih AlEmam, Adede Hawi, Adesegun A., Adey Feleke Desta, Adib Ghassan, Adjaratou Traoré, Adjiratou Aissatou B. A., Adodo Sadji, Adrian Egli, Adriano Mendes, Adugna Abera, Adul Candé, Afaf Alaoui, Afonso Pedro, Agbodzi B., Ageez A.M., Ahidjo Ayouba, Ahmed E Kayed, Ahmed El-Taweel, Ahmed Elsayed, Ahmed F. Gad, Ahmed Fakhfakh, Ahmed Kandeil, Ahmed M., Ahmed Mostafa, Ahmed O. S., Ahmed Reggad, Ahmed Taha, Ahyong Vida, Aicha Bensalem, Aida Sivro, Aissam Hachid, Ajili F., Ajogbasile F.V., Akim A. Adegnika, Akoele Siliadin, Akwii Patience Natasha, Aladje Balde, Alan Lemtudo, Alaoui Sanaa-amine, Alaruusi A. M., Alassane Ouro-Medeli, Albert Nyunja, Alberto Rizzo, Alemsegged Abdissa, Alemu Tike Debela, Alessandro Mancon, Alessandro Marcello, Alexander Goredema, Alexander Greninger, Alexis Ndjolo, Alexis Niyomwungere, Alfredo Mari, Alfredo Mayor, Ali M.A., Ali Zumla, Alia Ben Kahla, Alia Grad, Alice Kabanda, Alie Tia, Alimou Camara, Alimuddin Zumla, Alle Baba Dieng, Almoustapha I. Maiga, Amadou Alpha Sall, Amadou Daou, Amal Naguib, Amal Souiri, Amal Zouaki, Amalou Ghita, Amandine Mveang-Nzoghe, Amariane Koné, Amariane M. M. Koné, Ambroise Ahoudi, Amel Benyahia, Amel Nagiub, Amer K. E., Ameyo Dorkenoo, Amina Barkat, Aminata Dia, Aminata Mbaye, Aminata Mboup, Aminata Sileymane Thiam, Amine Idriss Lahlou, Amira Suliaman Wadi, Amivi Ehlan, Amujal Marion, Amuri Aziza, Amy Strydom, Anass Abbad, Anatole Nkeshimana, Andargachew Mulu, Anderrahmane Maaroufi, Andrea E. Luquette,

Andreas Shiningavamwe, Andres Moreira-Soto, Andrew Azman, Andrew J. Bennett, Andrew Tarupiwa, Anga Latifa, Ange Badjo, Angel Angelov, Angela Brisebarre, Angela M. Detweiler, Angoune Ndong, Anja Werno, Anna Julienne Selbe NDiaye, Anna-Lena Sander, Annajr B., Annamaria D'Aprile, Anne van der Linden, Annemiek van der Eijk, Annette Erhart, Anou M. Somboro, Anoumou Dagnran, Anthony Ahumibe, Anthony Levasseur, Antje van der Linden, Antoine Dara, Anu Jegede-Williams, Aouni M., Arjarquah A., Arlene Uwituze, Arlo Upton, Armel Poda, Arsène Somé, Arsène Zongo, Arsenia Massinga, Asare K.M., Ashaba Fred Katabazi, Asma Ferjani, Assane Dieng, Astou Gaye-Gaye, Atiga N., Atsbeha G. Weldemariam, Augustina Arjaquah, Auld A., Awa Ba-Diallo, Awatef ElMoussi, Awunyo Sena, Aya Mohamed, Ayman Farghaly, Ayo-Ale B., Ayoade F., Ayola Akim Adegnika, Ayong More, Ayorinde Babatunde James, Azami Nawfel, Azaria Diergaardt, Azuka Patrick Okwuraiwe, Babafemi O. Taiwo, Baboo S. B., Bahadoor B. S., Bahnassy A.A., Bakary Sanyang, Bakry U., Bamba Fatoumata Touré, Bamidele Iwalokun, Banda R, Bane S., Bankole Johnson, Barada Cisse, Barbra Murwira, Bas Oude Munnink, Batchi-Bouyou Armel L., Batra R., Belayachi Lamiae, Belkhir A.B., Ben Ayed I., Ben Morton, Ben Moussa, Ben Wulf, Bénédicte Ndeboko, Benhida Rachid, Benjamiin B. Lindsey, Benjamin H. Foulkes, Benjamin Hounkpatin, Benjamin Selekon, Bensaid M., Bernard Ssentalo Bagaya, Bert Vanmechelen, Bertrand Lell, Beth Mutai, Bethlehem Adnew, Beuty Makamure, Bighignoli B., Birahim Piere Ndiaye, Bishwo N. Adhikari, Bitrou S.M., Blaise MBoringong Akenji, Bode Shobayo, Boitumelo Zuze, Bonifacio Manguire Nlavo, Bouchra Belfquih, Bouchra Boujemla, Bouna Yatassaye, Bouzidi Aymane, Brian Andika, Bright K. Yemi, Bronwyn Kleinhans, Bruna Galvao, Bubacar Delgado Pinto Embaló, Bulelani Manene, Camille Capel, Campbell Anu, Carine Tchibozo, Carla Madeira, Carlos Cortes, Carniel Elisabeth, Carol Kifude, Carolle Yanique Tayimetha, Catherine E. Arnold, Catherine Okoi, Cecilia Waruhiu, Celestin Godwe, Celestina Obiekea, Celine Nkenfou, Chabuka L. B., Chakib Nejjari, Chambaro H., Chanda D., Changula K., Charifa Drissi Touzani, Charles Kayuki, Charles Nyagupe, Charoute Hicham, Chaselynn M. Watters, Chawech H., Cheikh Sokhna, Chenaoui Mohamed, Chiamaka Nwuba, Chilufya C., Chimaobi Chukwu, Chinyere Anyika, Chipimo P.J., Chitanga S., Chitenje M., Chiwaula M. J., Chouati Taha, Chris Mansell, Christelle Butel, Christian A. Devaux., Christian Drosten, Christian Ranaivoson, Christian Utpatel, Christophe Malabat, Chtourou A., Chukwu G., Claudia Daubenberger, Clement G. Kakai, Clement Masakwe, Clotaire Donatien Rafai, Collins Chenwi, Collins M. Misita, Collins Muli, Corine H. GeurtsvanKessel, Corinne Maufrais, Coulibaly Mbegnan, Crawford M., Cristina M. Tato, Cyrus Yiaba, D'Amore N., Dabiri Damilar, Dalia Ramadan, Damena D., Daniel Ouso, Daniela Pisanelli, Danilo Licastro, David Hammer, David Nieuwenhuijse, David Patrick Kateete, David Wilkinson, Davy Leger Mouangala, Dawit Hailu Alemayehu, Dawood R.M., De Sanctis R., Deborah N. A. Mettle, Demba Koita, Dennis Kenyi Lodiongo, Dennis Laryea, Department of Virology, College of Medicine, University of Ibadan, Dereje Leta, Dersi Noureddine, Desire Takou, Dessalegn Abeje Tefera, Dhwani Batra, Dia Ndongo, Diab A., Diabou Diagne, Diane A. Mabika, Diané Bamourou, Dianke Samaté, Diarra B., Didier Raoult, Dieudonne Mutangana, Dineo Emang Tshiamo. Gape Nyepetsi, Diosdado Odjama Nseng Ada, Djimabi Salah, Donatella Cedola, Doris Harding, Dorothy Yeboah-Manu, Dossou Ange, Dowbiss Meta Djomsi, Dragana Drinkovic, Draper C., Dumbuya Foday Sahr, Ebenezer Odewale, Eclou Sedjro, Edang-Minko Armand, Edgar Kigozi, Edith Koskei, Edith Nkwembe, Edmilson F. de Oliveira Filho, Edmira Maria da Costa, Edward Kiritu, Efrem S. Lim, Egon A. Ozer, Egyir B., Ehab Abdelkader

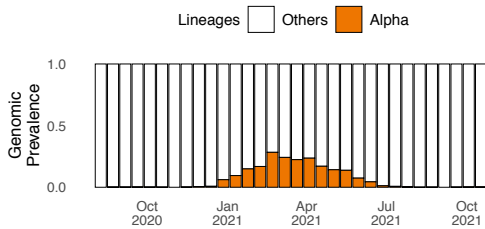
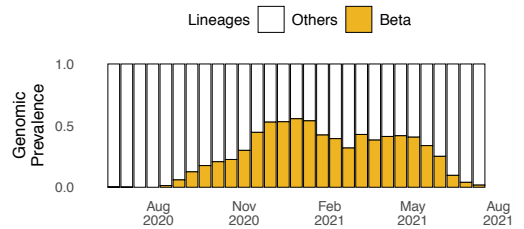
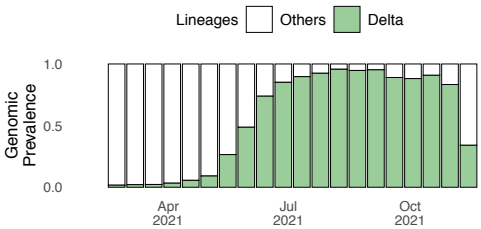
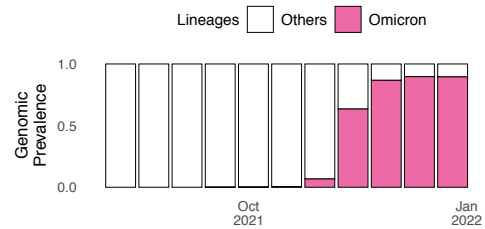
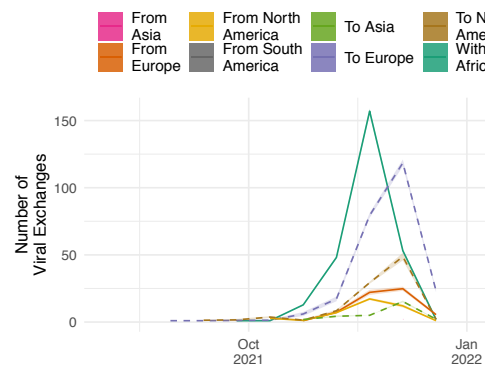
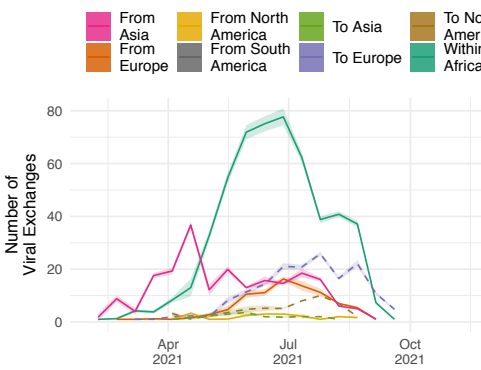
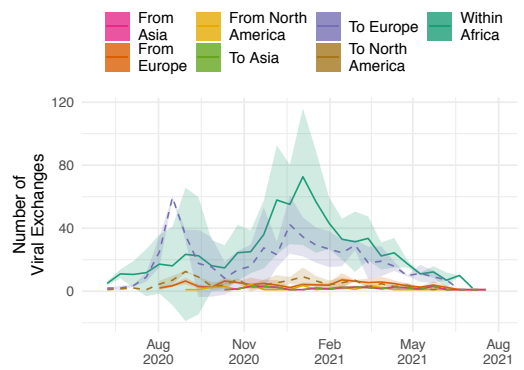
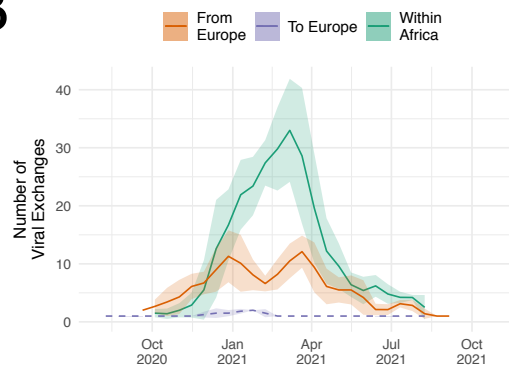
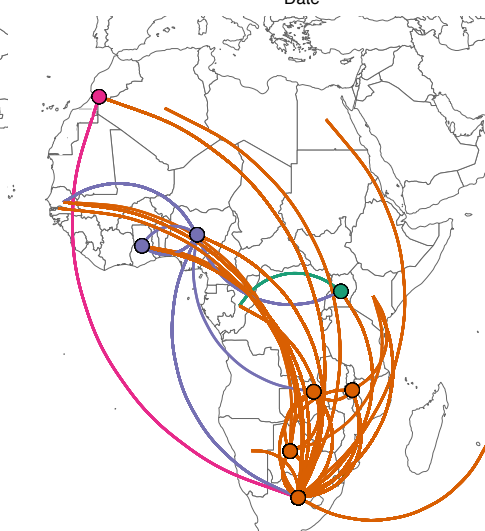
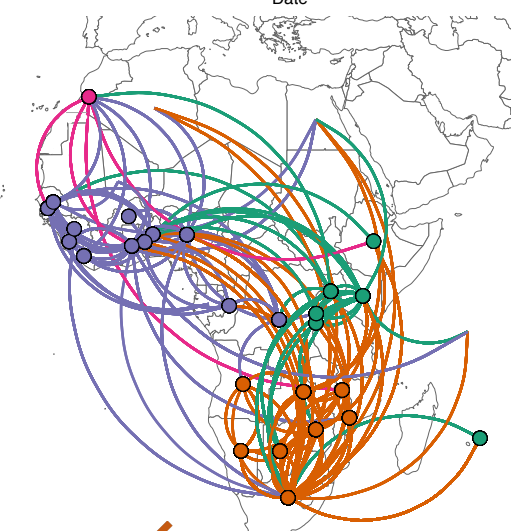
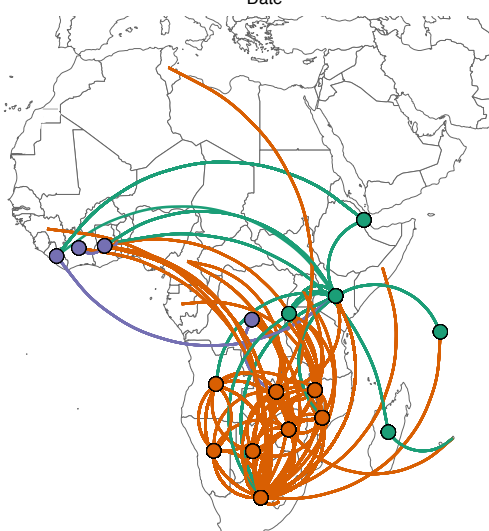
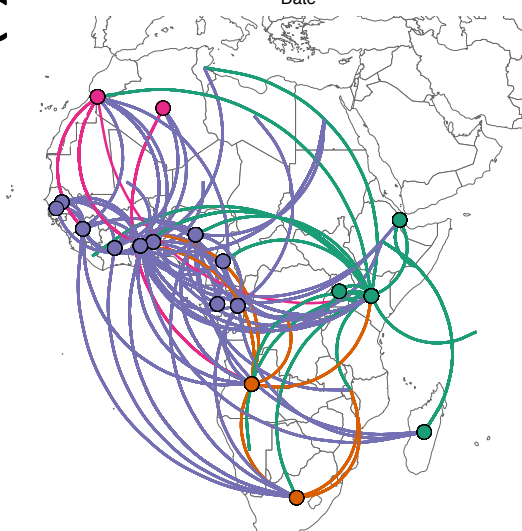
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Maud Vanpeene, Maurizio Margaglione, Max Bloomfield, May Abdelfattah, May Sherif Soliman, Mbengué Fall, Mdlalose K., Meei-Li Huang, Mehta S., Mélanie Albert, Melchior A. Joël Aïssi, Méline Bizard, Merabet Mouad, Meriem Laamarti, Messanh Douffan, Mhalla S., Michael Addidle, Michael Marks, Michael Nagel, Michael V. Deschenes, Micheala Davids, Michelle Balm, Michelle Lin, Michelle Tan, Mihrete A., Mikhail Olayinka BUHARI, Milanca Agostinho Cá, Mildred Adusei-Poku, Milkah Mwangi, Mina Kamel, Miranda J., Mireille Prince-David, Miriam Eshun, Misaki Wayengera, Mitali Mishra, Mjid Eloualid, Mly Abdelaziz Elalaoui, Mnguni A., Mnyameni F., Mogomotsi Matshaba, Mohale T., Mohamed Abdel-Salam Elgohary, Mohamed Ahmed Ali, Mohamed Ben Moussa, Mohamed Chenaoui, Mohamed El Sayes, Mohamed Elhadidi, Mohamed Gomaa Seadawy, Mohamed Hassan Abdoelraheem, Mohamed Hassany, Mohamed Houmed Aboubaker, Mohamed K.S., Mohamed Kamal, Mohamed Rhajaoui, Mohamed Seadawy, Mohamed Shamel, Mohamed Shemis, Mohammed K.S., Mohammed Walid Chemao Elfihri, Mohcine Bennani Mechita, Mokhtar Gomaa, Molalegne Bitew, Momoh M., Mona O.A. Alkarim, Monemo Pacome, Monilade Akinola, Monte A., Monuir G., Monze M., Mooko M., Morales A.N., Moreira-Soto Andres, Moriba Povogui, Mosepele Mosepele, Moses Chilufya, Moses Joloba, Moses Luutu, Mostafa Elouennass, Mostfa Elhoseiny, Mostfa Elnakib, Mostfa Yakout, Mouhcine Gardoul, Mouhssine Hemlali, Mouity Matoumba A., Mouna Ben Sassi, Mouna Safer, Mouneem Essabbar, Moustapha Mbow, Moustapha Nzamba Maloum, Moustapha Sakho, Moyinoluwa Odugbemi, Mtshali P., Mubemba B., Muchaneta Mugabe, Mufinda M., Muhammad Faisal, Muinah Adenike Fowora, Mukantwari Enatha, Muleya W., Muntaser elTayeb Ibrahim, Mupeta F., Murebwayire Clarisse, Mushal Ali, Mushal Allam, Mustapha Mouallif, Muuo S.N., Mwangomba W., Myriam Seffar, Mzumara T. E., N'dilimabaka N., Nabila Soara, Nabli A., Nadia El Mrimar, Nadia Rodrigues, Nadia Sitoe, Nafisatou Leye, Naguib A., Nalubamba K. S., Nancy M. El Guindy, Nandi Siegfried, Nardjes Hihi, Narjis Amar, Naryshkina E.A., Nathalia Endjala, Nathalia Garus-Oas, Nathan Kapata, Ndack Ndiaye, Ndahafa Frans, Ndam N.T., Ndéye Coumba Touré Kane, Ndiaye Ndack, Ndine Fainguem, Ndodo Nnaemeka, Ndumbu Pentikainen, Ndwiga L., Nedio Mabunda, Neff N., Negash A.A., Nejla Stambouli, Neto Z., Ngonga Dikongo A.M., Ngosa W., Ngozi Mirabel Otuonye, Niatou-Singa F. S., Nicaise T. Ndam, Nicholas Feasey, Nicholas Mwikwabe, Nicod J., Nicole Vidal, Nikki Freed, Nischay Mishra, Nissaf Ben Alaya, Nitin Savaliya, Noah Baker, Noé Patrick Mbondoukwe, Nokukhanya Mdlalose, Nonso Nduka, Noura M Abo Shama, Nourlil Jalal, Nsubuga Gideon, Ntuli N., Nuro Abilio, Nyam Itse Yusuf, Oby Wayoro, Ochwoto M., Ofonime Ebong, Ofori-Boadu L., Okomo Assoumou Marie Claire, Ola Elroby, Olabisi, Olabisi Ojo, Olajumoke Popoola, Olfert L., Olin Silander, Olufemi Obafemi, Olufemi Samuel Amoo, Olukunke Oluwasemowo, Olusola Anuoluwapo Akanbi, Oluwakemi Laguda-Akingba, Oluwatimilehin Adewumi, Omar Askander, Omar Elahmer-Abdulla Bashein- Rahma Algheriani-Ahram Alarif, Omar S., Omilabu S., Omnia Kutkat, Omoare Adesuyi, Omondi Francis Carey, Onalethata Lesetedi, Ongera E., Ontlametse T. Bareng, Onwuamah C.K., Ope-Ewe O., Osama Mansour., Oscar Kanjerwa, Oteng F., Otmane Touzani, Oumaima Ait Si Mohammed, Oumy Diop, Ouna Ouadghiri, Ousseynou Gueye, Owusu-Nyantakyi C., Oyefolu A., Oyeronke Ayansola, P Nthiga, Palomba S., Panja L., Papa Alassane Diaw, Park D., Pasacaline Manga, Patel H., Patience Motshosi, Pattoo M., Patrick Amoth, Patrick Descheemaeker, Patrick Mavingui, Patrick Tuyisenge, Pattoo M., Paul Dobi, Paul Liberator, Paulin N. Essone, Paulina Joãozinho da Costa Jarra Manneh, Pauline Yacine Sene, Paulo A Carralero RR Paixão J. P., Pavitra Roychoudhury, Peace O. Uche, Pei

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A**Alpha****Beta****Delta****Omicron****B****C****Direction of Viral Movement**

Regions of Origin

Eastern Africa

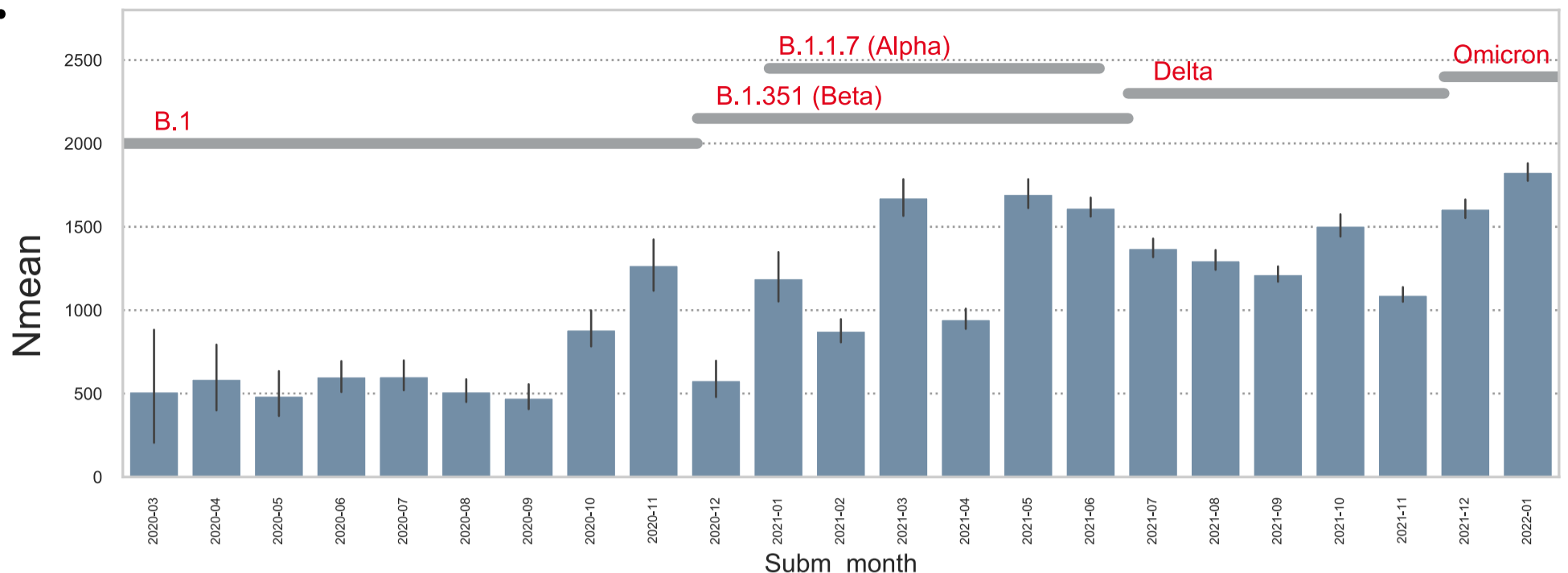
Northern Africa

Southern Africa

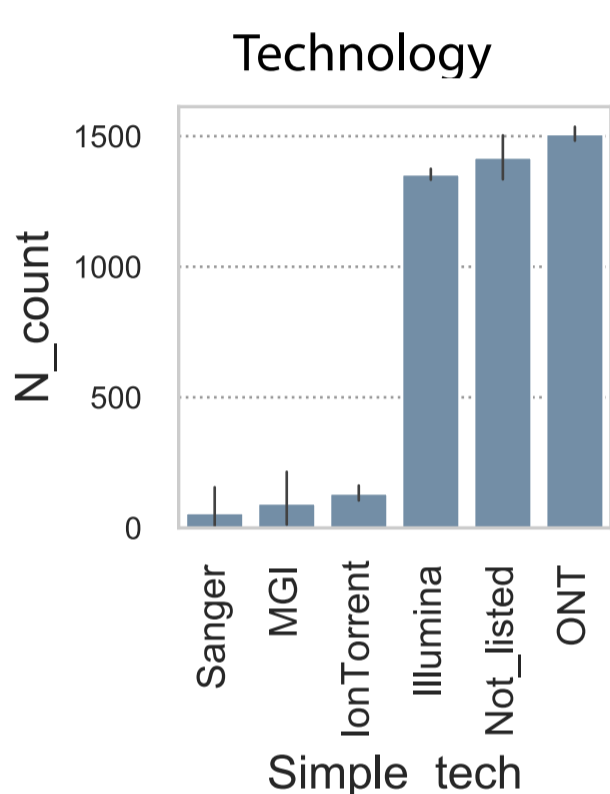
Western Africa

Mean number Ns in genomes, by Month, Technology, Primers

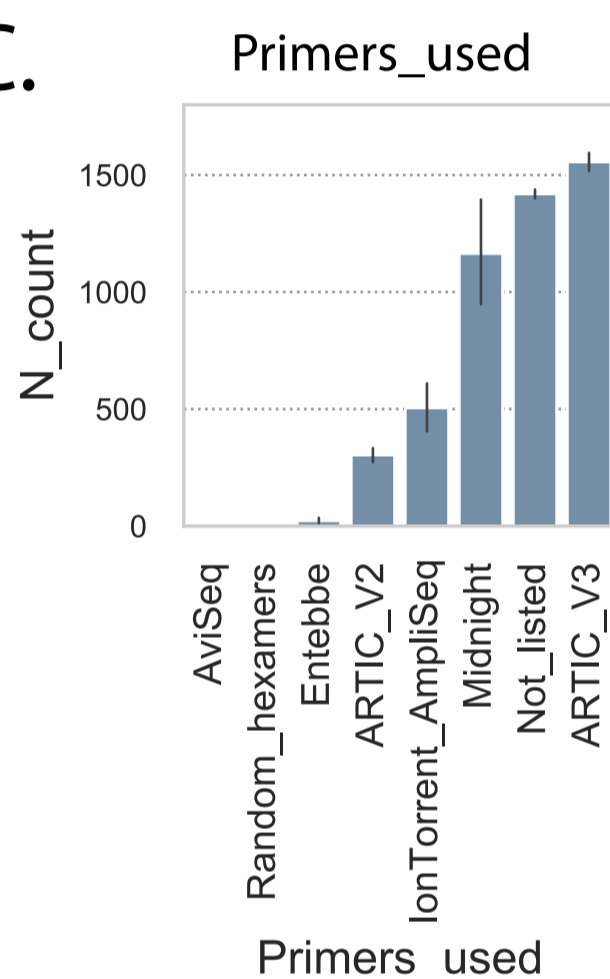
A.



B.



C.



D.

200 nt Gaps in 30 random genomes from Jan 2022

