2 Origin of imported SARS-CoV-2 strains in The

3 Gambia identified from whole genome sequences.

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24 Abstract

25 Severe Acute Respiratory Syndrome Coronavirus 2 (SARS-CoV-2) is a positive-sense 26 single stranded RNA virus with high human transmissibility. This study generated 27 Whole Genome data to determine the origin and pattern of transmission of SARS-28 CoV-2 from the first six cases tested in The Gambia. Total RNA from SARS-CoV-2 29 was extracted from inactivated nasopharyngeal-oropharyngeal swabs of six cases and converted to cDNA following the ARTIC COVID-19 sequencing protocol. Libraries 30 31 were constructed with the NEBNext ultra II DNA library prep kit for Illumina and Oxford 32 Nanopore Ligation sequencing kit and sequenced on Illumina MiSeg and Nanopore 33 GridION, respectively. Sequencing reads were mapped to the Wuhan reference 34 genome and compared to eleven other SARS-CoV-2 strains of Asian, European and 35 American origins. A phylogenetic tree was constructed with the consensus genomes 36 for local and non-African strains. Three of the Gambian strains had a European origin 37 (UK and Spain), two strains were of Asian origin (Japan). In The Gambia, Nanopore 38 and Illumina sequencers were successfully used to identify the sources of SARS-CoV-39 2 infection in COVID-19 cases.

40 Keywords: Coronavirus, SARS-CoV-2, nasopharyngeal-oropharyngeal, RNA,

41 genome, Nanopore, Illumina, sequencing, ARTIC protocol.

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43 Introduction

The emerging and re-emerging of pathogens such as severe acute respiratory syndrome-coronavirus 2 (SARS-CoV-2) pose a grave threat to human health.¹ The SARS-CoV-2 disease, first detected in Wuhan, China, in December 2019 has become a global pandemic,² and is causing an unprecedented burden on the health care 48 systems and economies globally.^{3–6} Worldwide, the number of cases has been 49 increasing exponentially,⁶ especially in Europe and America, with significant but variable case-fatality rates between continents. By April 28th, 2020, there were more 50 51 than 3.1 million SARS-CoV-2 confirmed cases and more than 200,000 deaths.7 Nevertheless, SARS-CoV-2 confirmed cases in sub-Saharan Africa are currently 52 53 relatively low, possibly due to much lower international air traffic than in other continents and thus a low number of imported cases.⁸ By the 28th April 2020, The 54 55 Gambia, a tourism hotspot, had reported a total of ten SARS-CoV-2 cases, including 56 one death. While the travel history of index cases may suggest the origin of infection, 57 phylogenetic analysis of the strains isolated from these cases and contacts will provide 58 a precise link between local transmission and other global populations.

59 The first SARS-CoV-2 case was reported to be an acquired zoonotic infection,^{9,10} followed by efficient and rapid human-to-human transmission from Wuhan, China, to 60 other Asian countries and then other continents.^{11–13} The single stranded positive 61 62 sense RNA genome of the SARS-CoV-2 is closely related to the Middle East 63 Respiratory Syndrome-Coronavirus (MERS-CoV) and the Severe Acute Respiratory Syndrome Coronavirus (SARS-CoV).⁹⁻¹⁴ These pathogens pose significant risk to 64 global health and modern-day life, hence the need for effective strategies to detect the 65 66 sources of infections, outbreaks and transmission patterns in different geographical settings. 67

The phylogenetic analyses of global SARS-CoV-2 sequences provide insight into the relatedness of strains from different areas and suggest the transmission of four superclades,¹⁵ geographically clustering into viral isolates from Asia (China), US (two super clades) and Europe. The objective of this analysis was to provide genome data on six

cases of SARS-CoV-2 in The Gambia, determine the source of these strains, baseline for subsequent local transmission, and contribute genomic diversity data towards local and global vaccine design. The Oxford Nanopore GridION and Illumina MiSeq platforms were utilized to sequence the viral genomes from four confirmed SARS-CoV-2, one inconclusive and one negative case by rRT-PCR. We also analysed the genomes of samples classified as indeterminate and negative by RT-PCR (COVID-19 detection assay) from two different cases respectively.

79 Methods

80 Sample acquisition

Nasopharyngeal-Oropharyngeal (NP-OP) swabs (451) from SARS-CoV-2 suspected cases and their contacts were transported to the Medical Research Council Unit The Gambia at London School of Hygiene and Tropical Medicine (MRCG at LSHTM). Of the 451 samples screened by rRT-PCR, ten were confirmed as SARS-CoV-2 cases and 5 as indeterminate cases (positive for the screening gene and negative for the SARS-CoV-2 confirmatory gene)

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For WGS, four SARS-CoV-2 confirmed cases, one indeterminate case and one negative case were processed Table 1. In one of the confirmed cases, different isolates from samples collected up to 10 days apart were sequenced. Of the 6 cases sequenced, 4 were male; 2 females, there was one death, two recoveries and two active cases.

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Table 1: Sample information for COVID-19 sequenced cases from The Gambia
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Case ID	Travelled from	Date Reported	Current Status	Number of	Time points	Library prep type			Sequencing	
				samples submitted		Depletion	ARTIC amplicon (NEB)	ARTIC amplicon (ONT -LSK109)	Illumina (MiSeq)	Nanopore (GridION)
A	London	16/03/20	Recovered	4	Days 0,4,7,10	2	4	4	4	4
В	Bangladesh	19/03/20	Dead	1	Day 0	0	1	1	1	1
С	France	20/03/20	Recovered	1	Day 0,	0	1	1	1	1
D	France	26/03/20	Active	2	Day 0,11	0	1	2	1	2
E	Netherland	23/03/20	Active	2	Day 0,14	0	1	2	1	2
F	Italy	13/03/20	Recovered	1	Day 0	0	1	1	1	1
Total 11			11		2	9	11	9	11	
Cases A-D = Confirmed RT-PCR COVID-19 cases Case E = Indeterminate by RT-PCR Case F = RT-PCR COVID-19 negative Cases A and D travelled to The Gambia in the same flight				ight						
Cases C and D both travelled from France										

101 RNA extraction

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- 103 Total RNA was purified from eleven samples Table 1 using the QiaAmp viral RNA mini 104 kit (Qiagen – 52906) following viral inactivation at the MRCG at LSHTM containment 105 level 3 facility. The purified RNA samples were quantified using Qubit RNA reagent kit on a Qubit fluorometer 3.0 (concentration range 3-7 ng/µl) (Invitrogen). RNA integrity 106 107 (RINe) was checked on the Agilent TapeStation 108 4200 yielding a RINe range of 2.1–5. 109 110 **Ribosomal RNA depletion, cDNA synthesis and Multiplex** 111
- 112 **PCR**

113 Two of the samples (day 0 and 4) from Case A were depleted using the RiboMinus 114 transcription isolation kit from ThermoFisher and purified using RNA purification beads 115 from Beckman Coulter. The purified rRNA-depleted samples were converted to cDNA 116 as per the NEBNext ultra II RNA library prep kit for Illumina (NEB, E7770L). Total RNA 117 from the rest of the samples was converted to cDNA according to the ARTIC amplicon sequencing protocol for SARS-CoV-2.^{artic} ARTIC protocol primer ^{artic} schemes for 118 119 SARS-CoV-2 (Version 2) were used for the multiplex PCR. Two primer pools at 10 µM 120 containing 98 primers each were used for the PCR amplification. The samples were subjected to 35 cycles of PCR. The purified products were visualised and quantified. 121 122

123 Illumina and Nanopore library Preparation and Sequencing
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125 Illumina and Nanopore library preparation

The purified cDNA from the depletion and PCR products from the ARTIC protocol were 126 normalised to 100 ng with EB buffer (10 mM Tris-HCI) to a final volume of 25 µl for 127 128 Illumina library preparation using the NEBNext ultra II DNA library prep kit for Illumina 129 (New England Biolabs, UK; E7645). Following 7 cycles of PCR enrichment, the 130 libraries were purified and quantified using the high sensitivity dsDNA Qubit kit and 131 sized using D1000 ScreenTape on the Agilent TapeStation 4200 (amplicon size range 132 519-572 bp). Each sample was normalised to 10 nM before pooling. The pool was run 133 at a final concentration of 10 pM on an Illumina MiSeg instrument using MiSeg V3 134 reagent kit. The pool was denatured with sodium hydroxide according to Illumina 135 recommendation and spiked with 5% PhiX (PhiX control v3 Illumina Catalogue FC-136 110-3001) before loading (Fig 1).

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138 Nanopore sequencing library preparation was performed according to the 139 manufacturer's instructions for the Ligation Sequencing Kit (SQK-LSK109, Oxford 140 Nanopore Technologies). Briefly, the cDNA samples were amplified using the ARTIC protocol and purified with 1X AMPure XP beads. Individual samples were then 141 142 subjected to end repair and adapter ligation following SQK-LSK109 protocol. 20 ng of each library was loaded on the Oxford Nanopore GridION on individual R9.4.1 flow 143 144 cells and sequencing data monitored on the fly using Rampart (v1.1.0). Fig 1 145 summarised the steps involved in sequencing using for both Nanopore and Illumina 146 platform.

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Fig 1. Summary of the Library preparation steps for Illumina and Oxford
Nanopore Sequencing Technology platforms. Library preparation took ~ 8 hours
for the Nanopore workflow and ~10 hours for the Illumina workflow.

153 **Quality control and read mapping for Illumina and Nanopore**

154 platform

Although a minimum read depth of 30X for the SARS-CoV-2 genome was targeted, more than 100X coverage was generated on both platforms. FASTQ files were subjected to various quality control checks and analysed following standard analysis pipelines (SARS-CoV-2 novel Coronavirus bioinformatics protocol; SAMTOOLS).

For Nanopore data, sequencing reads were quality checked using MinIONQC,¹⁶ and only reads with a minimum Q score of 7 were included in our subsequent analysis. Quality checked reads were run through what's in My Pot (WIMP) pipeline on the Oxford Nanopore EPI2ME platform to verify the number of reads characterised as SARS-CoV-2.

We used SARS-CoV-2 novel Coronavirus bioinformatics protocol developed by Nick Loman *et. al.* to analyse the Nanopore data.¹⁷ Firstly, we used "ARTIC Guppylex" to remove chimeric reads from each sample with the following parameters (--min-length 400 –max-length 700). Filtered reads were then mapped to Wuhan-Hu-1 reference genomes (accession number MN908947.3) and Single nucleotide polymorphism (SNPs) were generated based on the reference and a consensus genome for each sample was generated using SAMTOOLS (v1.9). To further validate our results, weran all genomes through the coronavirus typing tool (v1.13).

172 Similarly, 250 bp paired end Illumina reads were guality checked using FASTQC (v0.11.5).¹⁸ Reads with only a minimum Phred score of 30 were included in our 173 174 downstream analysis. One sample which was SARS-CoV-2 negative by reverse-175 transcription real-time polymerase chain reaction (rRT-PCR) was characterised as Bat 176 coronavirus using kraken and thus excluded from the analysis. Filtered reads were 177 then mapped to the Wuhan-Hu-1 reference genomes (accession number MN908947.3) using BWA-mem (0.7.17-r1188). BCFtools Mpileup (v1.8) was used in 178 179 creating a variant file. Finally, BCFtools consensus was used in generating the FASTA 180 consensus sequence for each sample.

181 Phylogenetic analysis for Illumina and Nanopore platform

Prank (v140603) was used to generate a multiple alignment of all the samples including some available reference genomes around the globe (Downloaded from RefSeq). These strains were selected based on the patients' travel history and the major geographical spread of the pandemic. We finally constructed a maximum likelihood phylogenetic tree using the General time reversible model (GTR) with IQTREE (v1.3.11.1). The Interactive Tree of Life (ITOL) (v5) was used to visualise and annotate the phylogenetic tree.

189 Results and Discussion

Whole genome sequencing data was generated from six confirmed cases from bothsequencing platforms; the additional time points from cases D and E were sequenced

only on the Nanopore GridION (Table 2). Two samples from the first case were sequenced on both platforms following ribosomal depletion, the results generated (not included) showed depletion of human sequences and the majority of the reads mapped to bacterial sequences with only 0.03% from the Illumina reads mapping to the SARS-CoV-2 reference strain. The rRT-PCR and the sequencing data generated are summarized in Table 2.

198 Table 2: Summary of COVID-19 results

Case ID	Time point		tic results -19 (Ct)	Sequencing Covid-19	Phylogenetic Inferences	
		Gene 1 (E gene)	Gene 2 (RdRP)	Illumina	Nanopore	Illumina & Nanopore
Α	Day 0	+	+	+		
	Day 4	+	+	+		Europe (UK)
	Day 7	+	+	+		
	Day 10	+	+	+	-	
В	Day 0	+	+	+		Asia (Japan)
С	Day 0	+	+	+	+	Europe (Spain)
D	Day 0	+	+	+		
	Day 11	+	+	Not		Europe (Spain)
				sequenced		
E	Day 0	+	-	+	1	
	Day 14	+	-	Not	1	Asia (Japan)
				sequenced		
F	Day 0	-	-	-	-	

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From the Illumina platform, a high quality read length of 250 bp paired-end reads was generated for each sample after 48 hrs post library preparation. Total number of sequences ranged from three to six million reads with an average mapping quality of 60 when mapped to the Wuhan reference genome. The Nanopore platform generated a read length of 400 - 17000 bp for each sample after 12 hrs of sequencing. It recorded a range of one to four million reads with average mapping quality of 58 across the
reference genome for each sample. To compare the results from both technologies,
consensus genomes were generated for each sample and a maximum likelihood
phylogenetic tree was constructed. Both platforms showed a similar topology with
1000 bootstrap clustering The Gambian isolates with the European and Asian strains
as illustrated in Figs 2 and 3.

Figure 2. A maximum likelihood phylogenetic tree of ten SARS-CoV-2 genomes isolated from The Gambia (Nanopore data) and 11 SARS-CoV-2 strains isolated in different parts of the world. The tree showed the genetic relation of strains isolated in The Gambian to the global circulating strains.

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Figure 3. A maximum likelihood phylogenetic tree of eight SARS-CoV-2 genomes isolated from The Gambia (Illumina data) and 11 SARS-CoV-2 strains isolated in different parts of the world. The tree showed the genetic relation of strains isolated in The Gambian to the global circulating strains.

Six genomes (4 samples from Case A, 1 from case D and 1 from case C) from the Gambian samples clustered with the European (Spanish and United Kingdom) SARS-CoV-2 strains. This is not surprising given that these patients had been in Europe before arriving in The Gambia. Although viruses are known to mutate and change rapidly,^{19,20} the viral genome of case A clustered on the same node at different time points indicating the patient was shedding the same virus with no observed polymorphism according to Nanopore results. Interestingly, the same samples 228 sequenced on the MiSeq suggested polymorphism at day 10, resulting in a longer 229 branch length compared to previous time points (Figure 3). Further analysis of the 230 Illumina data showed seven more SNPs in the day 10 sample compared to the other 231 time points. The SNP winked by the Nanopore phylogenetically, might have an 232 associated higher error rate compared to the Illumina. Strains from cases C and D, 233 both having travelled from France, were more closely related to the Spanish strain 234 included for comparison. Though cases D and A travelled to The Gambia on the same 235 flight, their strains had a different origin, indicating that they could have been infected 236 independently, before the start of their journeys.

237 The viral genome from case B who initiated travel from Bangladesh and then across 238 four other countries, including Senegal, before arriving in The Gambia, clustered with 239 a strain from Japan. This case may have contracted the infection in Asia and his travel 240 history suggests he could have contributed to infections in other countries. The two 241 isolates from case E at different time points clustered with strains from Japan as well. 242 Interestingly, case E samples were indeterminate by rRT-PCR diagnostics, even 243 though the outcome from multiple alignment showed no mismatch between the 244 sequences and the primer set. The indeterminate diagnostic rRT-PCR result could be 245 due to low sensitivity of the assay, an indication of low viral density of SARS-CoV-2 in 246 the sample. Therefore, subsequent follow up for such cases is essential to further 247 evaluate diagnosis and aid towards the understanding of the disease progression and 248 the evolution of this novel virus strain under different case management environments.

Although WGS data is still limited in sub-Saharan Africa, this approach has proven to be a highly sensitive, specific and confirmatory tool for SARS-CoV-2 detection. Hence, the use of second and third generation sequencing technologies coupled with

bioinformatics is quite imperative in providing data for monitoring transmissiondynamics.

254 From the two sequencing platforms, we were able to rapidly generate sequencing data 255 in 20 hours and 3 days after sample reception on the Nanopore and Illumina platforms 256 respectively. While Illumina sequencing may be more accurate in determining within-257 sample-diversity, Nanopore data can help with the understanding of the linkage 258 between SNPs within individual virions. The Nanopore platform with its flexibility for 259 number of samples per run, and the generation of data in real-time and at a reasonable 260 cost makes it most suitable for outbreaks. Therefore, with our optimised and ready-to-261 go workflow, we are set to generate data for tracking SARS-CoV-2 in The Gambia and 262 other African countries within 24 hours of sample reception. This would go a long way in providing knowledge on the molecular epidemiology of this disease, give the true 263 264 burden of the disease in this setting (as seen in the resolution of the indeterminate 265 cases) as well as provide information for African specific vaccine development and 266 inform policy makers on decisions for strategic control measures.

267 Conclusion

We have demonstrated that the Nanopore platform with the flexibility of high-end desktop sequencer (GridION) to the portable sequencer (MinION) in combination with the ARTIC protocol and workflow allows for cost-effective (wide range for the number of runs and samples per flow cell), and near real-time generation of pathogen sequence data. Our analysis has shown that the SARS-CoV-2 strains identified in The Gambia are of European and Asian origin and sequenced data matched patients'

- travel history. In addition, we were able to show that two COVID19 positive cases
 travelling in the same flight had in fact different sources of infection.
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285 Authors contribution

AK lead with Nanopore platform and the bioinformatics analysis, JM lead with Illumina

287 platform, MAK lead with viral Inactivation and purification. AK, JM, MAK, SJ, BS, MAO

288 & AB contributed to the sequencing pipeline and writing of the manuscript.

289

290 **Competing Interests**

291 The authors declare that they have no competing interests.

292

The Genomic Core facility at MRCG at LSHTM is the one and only certified service provider for the ONT GridION platform in Africa.

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296 Ethics statement:

- 297 The study has under gone under MRCG at LSTM and Gambia Government joint ethics
- committee for review.
- 299

300 Data and Materials Availability

- 301 The details of methods used in the paper is available as a supplementary document.
- 302 GISAID submission number: EPI_ISL_428856 and EPI_ISL_428857
- 303 The data from the genomes sequenced in the Gambia were submitted and available
- in Nextstrain website for real-time tracking of the pathogen evolution.
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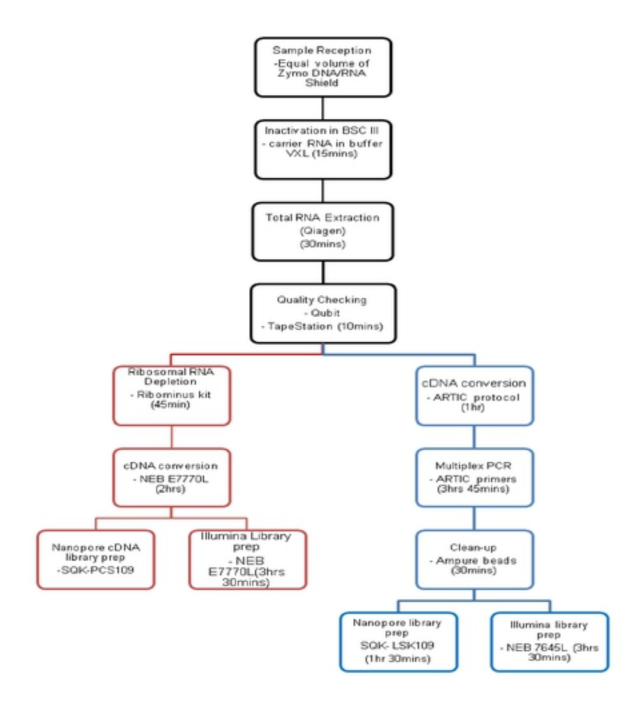


Figure 1: Summary of the Library preparation steps for Illumina and Oxford Nanopore Sequencing Technology platforms. Library preparation took ~ 8 hours for the Nanopore workflow and ~10 hours for the Illumina workflow.

Tree scale: 0.0001

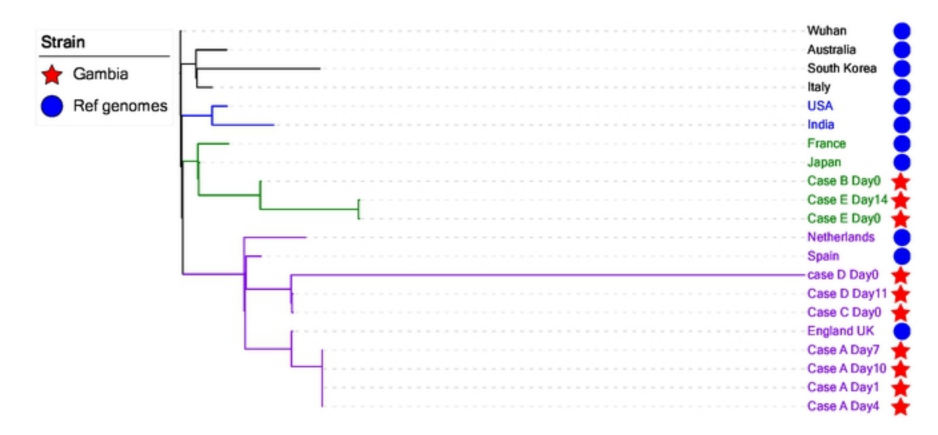


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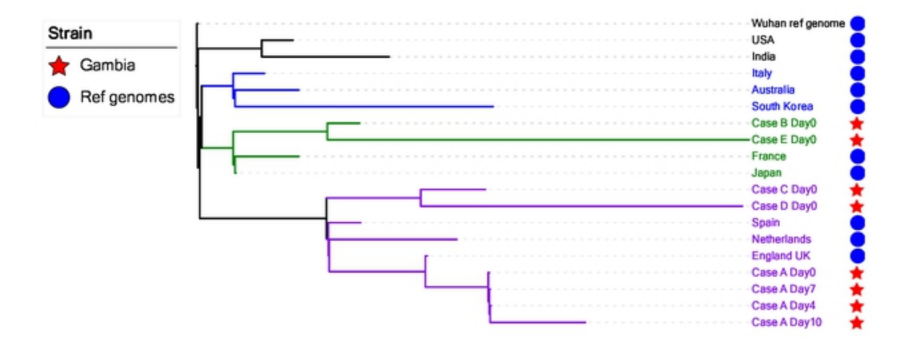


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