Overcoming Data Bottlenecks in Genomic Pathogen Surveillance

Ayorinde O. Afolayan,1, Johan Fabian Bernal,2 June M. Gayeta,3 Melissa L. Masim,3 Varun Shamanna,4 Monica Abrudan,5 Khalil Abudahab,5 Silvia Argimón,5 Celia C. Carlos,3 Sonia Sia,1 Kadahalli L. Ravikumar,1 Iruka N. Okeko,5 Pilar Donado-Godoy,7 David M. Aanensen,5 and Anthony Underwood5; for the NIHR Global Health Research Unit on Genomic Surveillance of Antimicrobial Resistance

1Department of Pharmaceutical Microbiology, Faculty of Pharmacy, University of Ibadan, Oyo State, Nigeria; 2Colombian Integrated Program for Antimicrobial Resistance Surveillance, Centro de Investigación Tíbaitatá, Corporación Colombiana de Investigación Agropecuaria, Tíbaitatá, Mosquera, Cundinamarca, Colombia; 3Antimicrobial Resistance Surveillance Reference Laboratory, Research Institute for Tropical Medicine, Muntinlupa, Philippines; 4Central Research Laboratory, Kempegowda Institute of Medical Sciences, Bengaluru, India; 5Centre for Genomic Pathogen Surveillance, Wellcome Genome Campus, Hinxton, Cambridge, United Kingdom; and 6The NIHR Global Health Research Unit for the Genomic Surveillance of Antimicrobial Resistance, Faculty of Medicine, University of Oxford, UK (anthony.underwood@cgps.group).

Performing whole genome sequencing (WGS) for the surveillance of antimicrobial resistance offers the ability to determine not only the antimicrobials to which rates of resistance are increasing, but also the evolutionary mechanisms and transmission routes responsible for the increase at local, national, and global scales. To derive WGS-based outputs, a series of processes are required, beginning with sample and metadata collection, followed by nucleic acid extraction, library preparation, sequencing, and analysis. Throughout this pathway there are many data-related operations required (informatics) combined with more biologically focused procedures (bioinformatics). For a laboratory aiming to implement pathogen genomics, the informatics and bioinformatics activities can be a barrier to starting on the journey; for a laboratory that has already started, these activities may become overwhelming. Here we describe these data bottlenecks and how they have been addressed in laboratories in India, Colombia, Nigeria, and the Philippines, as part of the National Institute for Health Research Global Health Research Unit on Genomic Surveillance of Antimicrobial Resistance. The approaches taken include the use of reproducible data parsing pipelines and genome sequence analysis workflows, using technologies such as Data-flo, the Nextflow workflow manager, and containerization of software dependencies. By overcoming barriers to WGS implementation in countries where genome sampling for some species may be underrepresented, a body of evidence can be built to determine the concordance of antimicrobial sensitivity testing and genome-derived resistance, and novel high-risk clones and unknown mechanisms of resistance can be discovered.

Keywords. bioinformatics; metadata; whole genome sequencing; WGS; antimicrobial resistance.

The utility of whole genome sequencing (WGS) for public health purposes has been proposed and implemented in a few countries, mostly to supplement existing methodologies with the expectation that it will become more widely used in the future [1]. The application of bacterial WGS specifically for the purposes of antimicrobial resistance (AMR) surveillance has also been the subject of scientific review [2, 3]. On one hand, these reviews are careful to highlight how WGS is not currently able to completely replace phenotypic antimicrobial sensitivity testing (AST), due to factors such as cost and the lack of concordance between the genotypic- and phenotypic-derived results for some bug-drug combinations.

For some species, such as Salmonella spp, Staphylococcus aureus, and Mycobacterium tuberculosis, the concordance is very good, but for others, such as Pseudomonas aeruginosa or Acinetobacter baumannii, the concordance is lower due to an incomplete understanding of mechanisms responsible for resistance [4, 5]. However, while a complete switch from phenotypic to WGS-based AMR surveillance is not likely in the near future, performing WGS of targeted pathogen samples has many benefits. WGS can enhance surveillance by providing information about the determinants responsible for resistance, the “vehicles” that carry them, such as plasmids and other mobile genetic elements, and the clonal lineages within which they are found. This enables the study of the emergence and expansion of AMR with a One Health approach, by facilitating the investigation of possible transfer of AMR between different reservoirs, such as humans, animals, and the environment. Because of this, many public health organizations are enthusiastic to adopt WGS in a targeted way to complement existing phenotypic-based surveillance, as exemplified in the Philippines [6]. Balancing the enthusiasm for adoption are the challenges in the implementation. These include challenges related to laboratory aspects of the process, such as cost and turnaround, and are
covered in more detail in [4, 7, 8]. The processes required to transform raw sequence data into human-interpretable results are often grouped under the umbrella term “bioinformatics.” Bioinformatics has its own set of challenges, which may also be barriers to introduction. Several manuscripts detail these, including a review of the steps required for the implementation of bacterial WGS, a recent report by the World Health Organization (WHO) on WGS for AMR surveillance, and others [2–4, 9]. These include availability of computing resources to run analytical pipelines; the wide range of software and associated catalogues of AMR determinants available to determine their presence, often confounded by lack of comprehensive benchmarking; standardization of bioinformatics pipelines used to produce the predictions based on WGS so that results are reproducible, including the use of standard operating procedures; requirements for quality assessment throughout the process from the raw sequence data through to the final interpretation; and trained personnel who are able to run and interpret the analyses.

For any institute starting on the journey toward the implementation of genome-based surveillance, these factors can become bottlenecks that impede adoption. In this manuscript we will describe the barriers that may prevent implementation of the bioinformatics required for WGS-based AMR prediction. Through examples derived from the implementation of WGS within the National Institute for Health Research Global Health Research Unit on Genomic Surveillance of Antimicrobial Resistance (GHRU), we illustrate possible avenues to overcome them. Crucially, and in addition to implementation, the final step in the pathway to WGS adoption is the interpretation of the bioinformatics outputs. This gap is primarily filled by hands-on training to analyze and interpret "real" data produced by bioinformatics processes [10].

**BIOINFORMATICS IMPLEMENTATION CHALLENGES**

Laboratory and bioinformatics processes generate various data types when performing pathogen genomics for AMR surveillance as illustrated by an example workflow (Figure 1).

**Data Collection and Integration**

Metadata describing samples are critical to contextualize genomics outputs by describing epidemiological and clinical background of

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**Figure 1.** An overview of a potential pathway from sample to phenotypic and genomic outputs. The bottleneck icons represent some of the steps in the process that can cause particular implementation challenges. ① Sample metadata cleaning and validation. ② Conversion of antimicrobial sensitivity testing minimum inhibitory concentration data into standardized formats for downstream processing and interpretation. ③ Quantitative quality assessment of raw reads and assemblies. ④ Processing raw reads to detect the presence or absence of genetic loci, genes, specific nonsynonymous mutations, and variants. ⑤ Aggregating results to produce human readable reports. Abbreviations: AMR, antimicrobial resistance; AST, antimicrobial sensitivity testing; MLST, multilocus sequence typing; ST, sequence type.
samples, allowing association between sample features and genomic markers. Data may be collected at a central site such as a public health laboratory, or by local sample collection sites such as hospitals, and sent to the center performing WGS. In the latter case, data may be stored in a variety of file formats, each structured in different ways. Once AST has been performed, results require merging with sample metadata. Further integration of data will be required later in the process to join these data to the genomic outputs. These data parsing steps can be performed manually but are person-hour expensive and subject to human error. In order to make these processes efficient and minimize turnaround time, automation is crucial.

Sequence Analysis
Once raw WGS data have been generated, sequence reads need to be processed to derive human interpretable results. This process has often been the hardest to implement, requiring specialist training (see next section), computing infrastructure, and software. Important considerations include using well-defined pipelines so that the outputs are reproducible, and utilizing a solution that is scalable so the hands-on time taken to generate results is, in general, independent of the numbers of samples processed.

Training
The specialist knowledge required to process WGS data, and understand the databases underpinning interpretation and the principles required when working with big data, is not often a component of the education of many healthcare professionals. If the WGS analysis software has a familiar web-based interface, such as those found on the Galaxy, Centre for Genomic Epidemiology, and Pathogenwatch web applications, less training is required with a focus being on interpretation of the data rather than performing the analysis [11–14]. However, this limits users to the analyses available on the web applications and does not prepare those wanting to perform more in-depth bioinformatics analyses. This requires training in running Linux command-line tools and interpreting the outputs.

IMPLEMENTATION SOLUTIONS
The GHRU project has sought to address the challenges described in the previous section. We present these as examples of how bottlenecks can be overcome, rather than promoting them as the only or best methods (Figure 2). In addition, the 4 units that have implemented a WGS AMR surveillance pathway present their experiences, unique circumstances, and implementation journeys.

Data Collection and Integration
Although there are multiple data files and many formats describing metadata, it is usually possible to describe the steps to turn them into consistent formats as a series of defined transformations. There are many alternative data science software solutions for this. We chose to use Data-flo software (https://data-flo.io/), which builds transformation dataflows by combining ready-to-use data adaptors. This technique is very similar to command-line approaches, such as R, or the Python pandas library, but allows users to construct and debug the data flow in a visual way [15, 16]. This enables data science specialists to construct a complex data parsing pipeline and share it with non-bioinformatics staff, such as those responsible for sample receipt, who are then able to run Data-flo without command-line experience. Within the project, we used Data-flo for several purposes: cleaning and transforming sample metadata data before combining with AST outputs (Figures 1 and 2, bottleneck ③; Supplementary Figure 1); converting different VITEK outputs into a single format that could be used as the input source for the WHONET AMR software (Figures 1 and 2, bottleneck ④) [17]; transforming bioinformatics outputs into a readable tabular format before uploading to a Google Sheet for ongoing storage of the results; and reading data from multiple sheets in different Google Sheets and joining them, in order to produce an aggregated table of epidemiological, laboratory, and bioinformatics results.

For example, a downstream Data-flo workflow was constructed to combine epidemiological metadata, AST data, and genomics outputs into the text format required as the source for visualization in Microreact (Figures 1 and 2, bottleneck ①); Supplementary Figure 1); converting different AST data before combining with AST outputs (Figures 1 and 2, bottleneck ②) [18]. A suite of programmatic functions was compiled within the grhuR package [19]). However, an important difference is that this did not require a relational database management system to be installed, backed up, and maintained. This would have represented a barrier to sustainability of the data storage solution.

Sequence Analysis
The analytical procedures to transform raw sequence into meaningful results are complex, often involving multiple steps and software. In order to efficiently parallelize this, so that multiple samples can be processed simultaneously on multicore workstations or in high-performance computing clusters, the use of workflow managers is recommended [20]. Installation of software with specific versions that may differ in different workflows can be one of the main hindrances to running bioinformatics analyses, especially as in many settings a UNIX system administrator with the necessary skills is not available. The use of containerization technology to bundle together specific software versions in a sandboxed environment is the most common solution for this [20–22]. For the GHRU project we took both these approaches, using workflows written for the Nextflow workflow manager, and Docker or Singularity containers to analyze the raw sequence data and produce de novo assemblies, AMR predictions, the 7-locus multilocus sequence type, and mapping-based single-nucleotide polymorphism phylogenies (Figures 1 and 2, bottleneck ②; Supplementary Table 1) [23–25].
Figure 2. Diagram showing the flow of data from sample receipt to final outputs and highlighting the solutions used for each step. The numbers refer to the same data bottlenecks described in Figure 1. The diagram starts when each bacterial sample is submitted accompanied by associated metadata. The sample is processed by traditional phenotypic antimicrobial sensitivity testing to produce minimum inhibitory concentration data. In parallel, genomic DNA from the sample is extracted and sequenced and whole genome sequencing data are processed through reproducible bioinformatics pipelines to produce multiple outputs such as multilocus sequence type, antimicrobial resistance determinant prediction, and single-nucleotide polymorphism–based phylogenies. These data are aggregated using Data-flo and stored in Google Sheets where they can be combined and manipulated using downstream processes such as R scripts or Data-flo pipeline to make final visualizations or reports. Abbreviations: AMR, antimicrobial resistance; AST, antimicrobial sensitivity testing; MIC, minimum inhibitory concentration; MLST, multilocus sequence typing; QC, quality control; RIS, Resistant, Intermediate, Susceptible; SNP, single-nucleotide polymorphism; WGS, whole genome sequencing.
To further simplify the software build process, the containers were built using the Conda package management software [26, 27]. This has allowed a total of 5979 genomes from species listed by the WHO as requiring priority research for AMR [28] to be processed across the GHRU partners in Colombia, India, Nigeria, and the Philippines (Supplementary Figure 2), and in parallel at the Wellcome Sanger Institute to confirm reproducibility.

The computational setup required for this was minimal. In each partner center, samples were processed on either high-specification laptops, or a workstation only requiring the installation of Ubuntu Linux, Java RE, Nextflow, and Docker, with either a simple command-line recipe or automated with ansible [29]. An example of a workstation specification is included in Supplementary Table 2. At Wellcome Sanger Institute, Docker images were converted to Singularity images and run on the high-performance computing Load Sharing Facility (LSF) cluster for a combined total of <24 hours. Analytical outputs were compared and shown to be identical, demonstrating the reproducible nature of the pipelines. A critical aspect to automate the processing of samples for WGS is a simple quality control procedure whereby samples that meet the appropriate studies can be triaged. We developed a Python package named Qualifyr to allow sorting of sample outputs into a red/amber/green status based on quality metrics derived from the fastqc read quality assessment package, confindr, and quast [30–33]. This produces a web-based graphical report as well as a text version for computational parsing, so samples that should be taken for further analysis can be easily assessed (Figure 1 and 2, bottleneck ©; Supplementary Figure 3).

Training

Bridging the knowledge gap is perhaps the most important requirement for effective implementation of WGS. It is important to consider trainee requirements. When processing pathogen WGS data for public health, there are 2 broad use cases: routine interpretation of the data for epidemiological investigation and intervention, and basic research to extrapolate the routine results and explore hypotheses. The first use case requires training to be able to run analytical pipelines and interpret the results, whereas the second use case requires additional training to be able to run more in-depth analyses. In this project, a tiered training program was developed. An example of a training stream for genome assembly is shown in Table 1 [29].

In this example, a public health bioinformatician would likely take the first and last tiers if they are going to run pipelines. An academic research bioinformatician would take the second tier in addition, in order to understand the commands used so they could make edits to the pipeline if required.

The training was enhanced logistically by delivering some of the command-line tutorial through shared terminal sessions (eg, with Tmux), allowing interactive training. This was supported by using a forum style chat application to allow trainees to talk and assist one another across different time zones [34]. A relatively stable internet connection is therefore a crucial requirement for collaboration in this kind of project. Once training was completed, a series of versioned standard operating procedures were designed to help the project users follow uniform procedures when generating results [35]. An important part of the training aimed to equip local bioinformaticians to train others [10]. Use of these training materials combined with application of the pedagogical techniques taught in the train-the-trainer course has allowed the GHRU units to train public health scientists and researchers outside of their own teams to run the same reproducible analyses [36].

IMPLEMENTATION VIGNETTES

One of the GHRU’s primary objectives is to build sustainable capacity in AMR laboratories. The goal is for each laboratory to be self-sufficient in the processes of laboratory sequencing and WGS analysis of samples collected for the purposes of AMR surveillance. Each laboratory has a unique set of local circumstances and objectives. These are described in Figure 3, in conjunction with the major milestones achieved.

DISCUSSION

There are many significant potential hurdles to overcome when implementing genomics for routine pathogen surveillance.

Table 1. Genome Assembly Training Stream

| Tier Outcome | Tier Title | User Proficiency | Notes |
|--------------|------------|------------------|-------|
| Understand the principles and be able to perform hands-on analysis using web tools | Genome assembly tutorial: principles and web-based analysis | Genomic scientist | Use the Galaxy web platform to run examples samples and assess output |
| Be able to implement analysis using command line | Genome assembly tutorial: command-line analysis | Command-line user | Run assembly with the command-line tools underpinning the Galaxy interface. The same parameters are employed so that the connection between running the assembly via the website and on the command line is apparent. |
| Be able to run reproducible high-throughput analysis and interpret the results | Genome assembly tutorial: reproducible batch processing | Command-line user | An in-depth knowledge of the command line is not required for this training. |
Through the use of standardized bioinformatics and a structured training program, 4 institutes in different countries and at different starting points along the genomics journey have been able to process hundreds of pathogen genomes from many clinically significant species. Further developments are still required to ensure sustainable and reliable analysis and interpretation of genome data for AMR surveillance. To check competency, running regular internal quality assessments and taking part in an external quality assessment (EQA) scheme will be essential. The GHRU has designed and implemented a pilot EQA that is currently in progress in the low- and middle-income country partner units. It has been reported that multiple bioinformatics processes for prediction of AMR are often discordant with each other and with phenotypic result, emphasizing the need for quality control standards, which an accredited EQA will partially address [37]. This also highlights the need for standardization in AMR databases. Currently, there are several widely used databases containing catalogues of AMR determinants. The incompatibility between these databases in terms of the nomenclature, use of different reference sequences for allelic
variants, and nonstandardized assignment of determinants to resistance to a specific antimicrobial is one of the remaining challenges facing those working in the field of genomic AMR surveillance. The hAMRonization software aims to rationalize the outputs from different AMR software, but a greater need remains the harmonization of the nomenclature and assignments of AMR determinants [38, 39]. This will be crucial for constructing a globally viewable database that stores genome-based AMR data, which is one of the future aims of the WHO Global Antimicrobial Resistance Surveillance System (GLASS) [40].

Future directions for the GHRU project include implementing cloud-based pipelines [41]. The advantages include no requirement for physical workstation procurement, installation, or maintenance, resulting in overall cost savings. However, slow and intermittent internet bandwidth may prove an insurmountable barrier in some locations. The workflow manager Nextflow that was used in the project already has an interface to launch pipelines in cloud computing environments [42]. Training will also be improved using online notebooks such as Google Colaboratory and Kaggle, so that trainees can take advantage of command-line training without requiring a Linux machine and with interactive examples [43, 44]. Additional training in interpretation of results as well as running analyses will be implemented, because the genetic factors contributing to phenotypic AMR are complex. Furthermore, our understanding of the genetics of AMR is constantly changing, and therefore training to build skills that allow the assessment and incorporation of new software into pipelines will be required to ensure sustainability. Together, these iterative improvements will further lower the barriers for bioinformatics implementation for WGS-based surveillance.

CONCLUSIONS

There are several alternate routes to implementing pathogen genome sequencing for surveillance of AMR, each of which has its own strengths. Here we have set out principles for overcoming some of the challenges. The fundamental process is running reproducible analytical pipelines so that consistent results are obtained from multiple centers, as measured by the text/digital output from the pipelines, prior to interpretation. Just as important are the processes to clean incoming data pertaining to the samples, and the downstream processes to amalgamate these with the genomic outputs. We chose to use online spreadsheets as a proxy for database tables and programmatic means to fetch, join, and filter these to produce aggregated data that could be tabulated or visualized. Training was crucial for the implementation and allowed bioinformatics scientists to both run standardized pipelines and perform more specialized, species-directed analyses. For those wanting to run consistent pipelines, online platforms offer capacity without requiring more in-depth bioinformatics expertise. Pathogenwatch offers a web-based platform for AMR analysis and phylogeny generation of Campylobacter, Klebsiella, Neisseria gonorrhoeae, Staphylococcus aureus, and Salmonella Typhi [13, 14]. The Center for Genomic Epidemiology website offers services for phylogenetic tree building and AMR prediction [45]. However, when using these services it is important to realize that the underlying analyses may change and recording the exact version of the analysis software becomes more challenging.

In the long term, the WHO GLASS system will accept genomic-based AMR prediction and supporting information [40]. Several factors will be important for its success: a mechanism to check the quality of incoming data; a standardized data format; and a minimal data specification. The implementation described here will help inform these factors.

Supplementary Data

Supplementary materials are available at Clinical Infectious Diseases online. Consisting of data provided by the authors to benefit the reader, the posted materials are not copyedited and are the sole responsibility of the authors, so questions or comments should be addressed to the corresponding author.

Notes

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