

SAGESA Research and Training Network Activities

1. Establishment of the SAGESA Research and Training Network

In 2021, the SAGESA project established a research and training network, to facilitate capacity building for pathogen genomics and surveillance of anti-microbial resistance (AMR) in Africa. The SAGESA project is funded by the Cambridge Africa ALBORADA Research Fund, and is a collaboration by scientists from the National Institute of Communicable Diseases in South Africa, the Faculty of Medicine and Health Sciences, University of Zimbabwe, Wellcome Connecting Science and the Centre for Genomic Pathogen Surveillance.

Project goal

The goal of this project was to establish a peer network platform which provides access to training and expertise to scientists interested in genomics of disease pathogens and surveillance of antimicrobial resistance (AMR).

Objectives

- Establish the training and research network for genomic surveillance of AMR in Africa
- Provide training for bioinformatics in AMR genomics in the SADC region
- Establish collaborative research and training projects for AMR surveillance

Target audience

Scientists based anywhere in the world interested in developing skills and resources for AMR genomics in Africa and globally.

Joining the network

To join the network simply <u>contact us</u> or access an <u>online form</u> for providing basic personal information, country of origin and consent for use of media for promotional purposes by the collaborating organisations.

Other network scientists can be found on this Padlet.

Additional interactivity is available on the Slack channel via this link

THE SAGESA Core Team developed activities encompassing virtual meetings and training and facilitated a network of scientists interested in sharing knowledge and skills in genomics of antimicrobial resistance (AMR) Africa.



2. SAGESA Launch and Needs Assessment Workshop

The SAGESA launch meeting, held in November 2021, facilitated the introduction of initial network members – approximately 50 at the time. This was followed by a needs assessment workshop to explore network, research and training needs in Africa.

Workshop objectives

- Official launch of SAGESA Network the first Africa AMR genomics research and training network.
- Discuss needs and gaps within the region with regards to application of genomics for AMR surveillance - a focus on training needs and network development.
- Provide an overview of the planned/proposed SAGESA activities and discuss the future of the network.

A virtual hybrid model used live discussion of topics and the Padlet platform where participants provided detailed information about - the current status of training and research, training needs and research interests, and expectations of SAGESA network.



3. SAGESA Network Curriculum

Needs assessment analysis from the SAGESA Launch Workshop guided the development of the SAGESA Curriculum consisting of 5 webinars and a 3 Day bioinformatics workshop. Feedback was collected after each event. Click on links below for:

Webinar schedule

AMR Bioinformatics Course

Media and Networking Platforms

Join Slack Workspace
Visit SAGESA website
Follow us on Twitter and keep growing the network
Consent SAGESA consent form

SAGESA Core Team

Alice Matimba - WCS Stanford Kwenda - NICD Monica Abrudan - CGPS Isabela Malta - WCS Dusanka Nikolic - WCS Joconiah Chirenda - UZ

Acknowledgements

Jorge Batista da Rocha (WCS) Martin Aslett (WCS) Melanie Sharpe (WCS



SAGESA Webinar Schedule

February - July 2022

Webinar Title	Title and objective	Session outline (All sessions were held virtually and CAT timezone)
Data analysis pipelines and Galaxy, Nextrain Organised by: Stanford Kwenda Isabela Malta Monica Abrudan Dusanka Nikolic	To provide an overview of workflows and tools used for WGS analysis and AMR genomic surveillance	Date: 4 february 2022 11:00 Welcome - Stanford Kwenda 11:05 Data analysis pipelines and Galaxy - Peter Van Heusdsen 12:05 Coffee break 12:10 Data analysis pipelines, Nextrain, Q&A - Moderation by Stanford Kwenda 12:40 SAGESA announcements - Alice Matimba 12:50 Wrap-up & evaluation poll - Dusanka Nikolic
Samples to Sequencing Organised by: Joconiah Chirenda Isabela Malta Dusanka Nikolic	Discuss issues around sample and data collection, sample processing, sequencing technologies and procurement issues	Date: 25 February 2022 11:00 Welcome and meeting opening – <i>Joconiah Chirenda</i> 11:05 Sample preparation: Lessons from the field – <i>Tapfumanei Mashe/David Amini</i> 11:35 Experiences from the SAGESA community – <i>ALL</i> 11:45 Sequencing tools – <i>Taya Forde</i> 12:15 Q&A plus experiences from the community – <i>Moderation by Taya Forde</i> 12:25 Break 12:30 Short case studies on Samples to Sequencing – <i>Sabelle Jallow</i> 12:50 Announcements – <i>Joconiah Chirenda</i> 12:55 Wrap-up & evaluation poll – <i>Dusanka Nikolic</i> 13:00 End



AMR genomics tools and databases Organised by: Stanford Kwenda Isabela Malta Monica Abrudan Dusanka Nikolic	Discuss genomics tools and resources available for AMR prediction and global pathogen surveillance	Date: 18 March 2022 11:00 Welcome 11:05 AMR prediction tools & databases overview - Narender Kumar 11:40 rMAP + Q&As - Ivan Sserwadda 12:05 Question and answer and experiences from the community - Moderation by S Kwenda 12:10 Coffee break 12:15 Pathogenwatch + Q&As - Silvia Argimon 12:50 SAGESA Announcements - Alice Matimba 12:55 Wrap-up & evaluation poll - Dusanka Nikolic 13:00 End
Skills lab 1: Study design and grant writing Organised by Isabela Malta Alice Matimba Joconiah Chirenda Dusanka Nikolic Stanford Kwenda	To provide tools and resources on how to get started with grant writing, project design and management, and collaborating on a grant project team	Date: 8 April 2022 11.00 Welcome - Isabela Malta 11.05 Key elements for grant writing - presentation and Q&A - Helena Kuivaniemi 11.45 Logistics and Project Management, How to design a project - Vicky Nembaware 12.10 Break 12.15 Successes and failures Part 1 - Boua Palwendé Romuald 12.25 Successes and failures Part 2 - John Ndemi Kiiru 12.35 Success and failures - Open discussion - Joconiah Chirenda 12.50 Activity for Session 2 - Samantha Sampson 12.55 Feedback and wrap-up - Stanford Kwenda
Skills lab 2: Grant writing Project Organised by Isabela Malta Alice Matimba Joconiah Chirenda Dusanka Nikolic Stanford Kwenda	To discuss grant project assignments and provide feedback	Date: 8 July 2022 10:00 Welcome - Isabela 10:05 Aims Page Presentation Group 1: John, Sheila, Geoffrey, Lesley, Richmond 10:10 Aims Page Presentation Group 2: Mulemba, Musa, George, Oscar 10:15 Aims Page Presentation Group 3: Azra, Emmanuel, James, Shamsudin, Rhama 10:20 General feedback and remarks - Helena and Sam 10:30 NIH Biosketches - Gerard 10:50 New assignment plan - Helena and Sam 11:00 Break 11:05 Network discussions - Dusanka 11:50 Wrap-up - Isabela



AMR Bioinformatics Course - (Virtual)

25-27 May 2022

Links in this document

Course overview
Target Audience

Learning outcomes

Content and activities overview

Timetable

Organising and Training team

Course materials

Software and datasets

Planning and development

<u>Acknowledgements</u>

Course overview

Course website

In collaboration with the <u>National Institute of Communicable Diseases (NICD)</u>, South Africa, we are pleased to announce the AMR Bioinformatics course as part of the <u>SAGESA network</u> aimed at training and building capacity for genomic surveillance of AMR in Africa.

Antimicrobial resistance (AMR) is a global public health emergency and threatens the safe delivery of modern medical care. Global projections of the negative impact of AMR predict the loss of nearly 10 million lives and a cumulative loss of global production up to 100 trillion US dollars by 2050. AMR requires immediate, concerted, international, collaborative action to monitor its prevalence and spread throughout the world. The importance of this has been recognised both by the United Nations and by the World Health Organisation; the latter has published a list of priority pathogens and a global strategy for AMR surveillance.

The challenges of controlling AMR in low- and middle-income countries (LMICs) may be considerable for several reasons. Firstly, antibiotics may be available for human and animal use without a prescription, resulting in unregulated use. Secondly, accurate data on antimicrobial prescription or consumption may not be readily available. Thirdly, there may be limited knowledge about the appropriate use of antibiotics and a lack of antimicrobial policies or stewardship programmes. Finally, there may be limitations in healthcare facilities and laboratory capacity for the detection and management of AMR.

This 3-day workshop aims to address some of these challenges by improving the knowledge and skills of individuals interested in antimicrobial resistance in bacterial pathogens in Africa. It will contain hands-on WGS-based bacterial typing methods such as AMR detection, cgMLST, MLST, core genome SNP based phylogenetic analysis.



Target audience

This workshop is free to attend and open to <u>SAGESA</u> network members based in Africa who are actively engaged in, or soon to commence, research, clinical practice, provision of clinical services, or policy development related to AMR in bacteria. To join the network <u>please click on this link</u>.

Learning outcomes

At the end of the course participants should be able to

- Implement data workflows and bioinformatic analysis of genomic sequence data, including risk and resistance prediction for bacterial epi and pandemic response
- Explain Genomic surveillance and linking epidemiological data, outbreak analyses and global contexts, geographical contexts
- Demonstrate interpretation and application in public health and epidemic/pandemic response

Content and activities overview

- Genomic data analysis
- Interpretation for various applications
- Training support mentorship and networking

Organisers and developers

Wellcome Connecting Science - Isabela Malta, Alice Matimba, Dusanka Nikolic, Melanie Sharpe, Martin Aslett

National Institutes of Communicable Diseases - Stanford Kwenda University of Zimbabwe - Joconiah Chirenda

Module leads

- Annie Chan, National Institute for Communicable Diseases, South Africa
- Arun Decano, St. Andrews University, UK
- Erikson Odih, University of Ibadan, Nigeria
- Narendar Kumar, Wellcome Sanger Institute, UK
- <u>Pimlapas Leekitcharoenphon (Shinny)</u>, National Food Institute, Denmark
- Stan Kwenda, National Institute for Communicable Diseases. South Africa

Guest Speakers

- Plenary Lecture: Rene Hendriksen, National Food Institute, Denmark
- Lecture 1: Zamantungwa Khumalo, ClinVet, South Africa
- Lecture 2: Joconiah Chirenda, University of Zimbabwe, Zimbabwe
- Lecture 3: Sabelle Jallow, National Institute for Communicable Diseases, South Africa
- Lecture 4: Lavania Joseph, National Institute for Communicable Diseases, South Africa
- Lecture 5: Michelle Lowe, National Institute for Communicable Diseases, South Africa



Timetable

Pre Course	AMR Bioinformatics Workshop - Africa. 25th to 27th of April				
Pre-course intro	Time	Wednesday	Thursday	Friday	Time
Participants Intro	Time	Wednesday 25/5/2022	Thursday 26/5/2022	Friday 27/5/2022	i iiie
Familipants intro	08:30	Genomic surveillance Over	20/3/2022	21/3/2022	08:30
Overview of lab	00.50	Genomic surveillance Over			00.30
workflow,	09:00	Intro to sequence data;	AMR detection &	Phylogenetic analysis	00.00
Zamantungwa	00.00	QC of raw sequence data	surveillance (seminar)	(SNP-based and	05.00
Khumalo, ClinVet		Contamination & species	Rene Hendriksen	cgMLST) (lecture &	
Taniaio, Giirivat		ID (lecture & practical 1)	7,0710 110714711.0077	practical 5)	
	09:30	Annie Chan		Pimlapas	09:30
	10.00	-		Leekitcharoenphon	10.00
	10:00	-	Intro to bacterial		10:00
	40.00	-	subtyping approaches		40.00
	10:30		(lecture) Erikson Odih		10:30
	14.00	Break Lab based AMR typing using	Break	Break	44.00
	11:00	ASTs (Lecture)	Bacterial subtyping	Phylogenetic analysis	11:00
	44.00	Sabelle Jallow	approaches (practical 4)	(SNP-based tree	44.00
	11:30	-	Erikson Odih	construction) (group	11:30
	40.00		Levenh Dennik	project 2) <i>Pimlapas</i>	40.00
	12:00	-	Lunch Break		12:00
	12:30	De novo assembly vs. ref	Case studies: genomic	Case studies:	12:30
		based mapping (lecture &	surveillance in public	genomic surveillance	
	13:00	practical 2)	health (TB) Lavania	in public health	13:00
		Arun Decano	Joseph	Michelle Lowe (pre-	
	13:30		AMR typing - WGS data	Transmission analysis	13:30
			(lecture)	(lecture & practical 6)	
	14:00		Arun Decano	Narender Kumar	14:00
	14:30				14:30
	15:00	Break	Break	Break	15:00
		Genome annotation	AMR typing using WGS		
	15:30	(lecture & practical 3)	data (group project 1)		15:30
		Stan Kwenda	Arun Decano,		
	16:00		Stan Kwenda,		16:00
			Narender Kumar	discussion	
	16:30				16:30



Course materials

accessible via Github repository - https://github.com/WCSCourses/AMR-Bio-Africa-2022

Introductory session

- Welcome and overview
- Reading material

Session 1: Intro to sequence data & QC

- <u>Introductory presentation</u>
- Hands-on exercises manual
- Hands-on exercises example commands

Session 2: Genome assembly

- Introductory presentation
- Hands-on exercises manual

Session 3: Genome annotation

- Introductory presentation
- Hands-on exercises manual
- Hands-on exercises online version

Session 4: Bacteria subtyping

- <u>Introductory presentation</u>
- Hands-on exercises manual

Session 5: AMR detection

- <u>Introductory presentation</u>
- Hands-on exercises manual

Session 6: Phylogenetic analysis

- <u>Introductory presentation</u>
- Visualization presentation
- Hands-on exercises manual

Session 7: Transmission analysis

- <u>Introductory presentation</u>
- Hands-on exercises manual
- Use this link to navigate to the microreact instance for this tutorial

Session 8: SAGESA Network discussion

Course feedback

^{*}Some content was adapted from the WCS AMR for Bacterial Pathogens Course



Software and datasets

Software Name	Version*	Link	Session
Anaconda3			Raw sequence data QC
FastQC	0.11.9		De novo assembly
MultiQC	1.6		AMR prediction
Trim-galore	0.6.2		Genome annotation
BactInspector	0.1.3		Reference based mapping
Shovill	1.1.0		General tools
QUAST	5.0.2		
abricate + (preloaded resfinder DB & NCBI DB)	1.0.1		
prokka			
snippy			
bwa			
bowtie2			
samtools	1.11		
R + TidyVerse			



Planning and development

Technical resources and platforms

All participants and training team members will use own laptops, and should have access internet.

- Minimum requirements for the computer should be 8GB RAM, i5 processor and 80GB hard disk space
- All have to install Virtual Box to use a Virtual Machine containing all software and datasets
- Zoom as the meeting platform, participants pre-register
- Slack Workspace for interactions, Q&A and networking
- All courses materials compiled on Github repository

Advertising and marketing

- Directly to SAGESA members via emails and Slack
- WCS Newletters, Twitter and LinkedIn Platforms
- SAGESA Twitter
- Personal networks in Africa region

Application and selection

- Application set up in V1 platform
- Based on justification and relevance of work or research; Benefit to current work and to colleagues
- Application received over 300
- No. of applications accepted 43

Application form questions:

- 1. First name
- 2. Last name
- 3. Current role
- 4. Department, institute
- 5. Country6. Nationality
- 7. Highest level of qualification
- 8. Subject area of work
- 9. Previous jobs (please list up to a maximum of 3)
- 10. Justification for attending this course. Please note that our courses are frequently oversubscribed and the selection committee relies heavily on this section to make its decision. Please provide an outline of your work (research project(s) and/or clinical/healthcare activities)
- 11. How will you use the training to benefit your work?
- 12. Are you currently generating or analysing genome sequence data?
- 13. How would you rate your skills in command line/Linux?
- 14. What is your computer processor (e.g i3, i5, i7...)
- 15. Do you have a stable internet connection?

Post-course feedback questions

Refer to attachments separately

Timelines Overview



Item	Activity	Dates	Respons.
Finalise course description	Programme outline; target audience, pre-requisites	Done	AM/IM
Development and delivery teams	Finalise module leads, instructors, facilitators, assistants	Get confirmation by 28/02	IM/SK
Curriculum	Course plan - Develop module outlines, objectives, activities	18/03	AM/IM/SK
Call for participant applications, review and selection	Application forms	Open: 16/03 Close: 15/04 Selection: 25/04	IM/MS
Develop training materials	Lecture videos, resources and contact session guides	15/04	Training team
Reviewing and testing	All materials including VM and software and datasets to be tested and approved	29/04	MA/SK/ Training team
Course delivery	Virtual delivery	25 - 27 May	All
Feedback	Participant and training team feedback	10/06	All

Acknowledgements

Wellcome Connecting Science (WCS) – Learning and Training Team, UK National Institute of Communicable Diseases (NICD) Staff, South Africa Centre for Genomic Pathgen Surveillance (CGPS), UK

University of Zimbabwe

Funding: Cambridge Africa ALBORARA Fund

*Some content of this course was adapted from WCS AMR for Bacterial Pathogens Course