



Certificate of Achievement

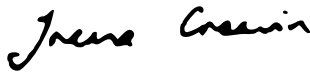
Manimozhi S

has completed the following course:

BACTERIAL GENOMES II: ACCESSING AND ANALYSING MICROBIAL GENOME DATA USING ARTEMIS WELLCOME CONNECTING SCIENCE

This online course covered the principles of microbial bioinformatics analysis and comparative genomics. Artemis, an open source genome browser, was used to investigate whole bacterial genomes, and through analysis of bacterial genes and proteins, the genomic features of pathogens were explored.

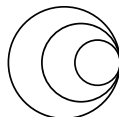
The Royal College of Pathologists (RCPATH) has accredited this course for 15 Continuing Professional Development (CPD) credits. This applies to medical staff and clinical scientists in career grade posts who are enrolled with one of the Royal Colleges for CPD purposes.



Dr Treasa Creavin
Scientific Programme Manager
Wellcome Genome Campus Advanced Courses and
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Professor Nicholas Thomson
Group Leader, Bacterial Genomics and Evolution
Wellcome Sanger Institute, Wellcome Genome Campus



**wellcome
connecting
science**

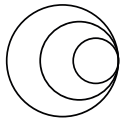
Accredited by



The person named on this certificate has completed the activities in the attached transcript. For more information about Certificates of Achievement and the effort required to become eligible, visit futurelearn.com/proof-of-learning/certificate-of-achievement.



This certificate represents proof of learning. It is not a formal qualification, degree, or part of a degree.



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93%
OVERALL
SCORE

This course covered the principles of microbial bioinformatics analysis and comparative genomics. Using Artemis, an open source genome browser, whole bacterial genomes were investigated: comparisons of reference and draft genomes were made; genome annotation and file formats were shown and together explored, GC (guanine-cytosine) content and the use of this metric to analyse functional blocks in the genome was covered; and Artemis was used to compare two bacterial genomes and their gene content.

STUDY REQUIREMENT

3 weeks, 5 hours per week

LEARNING OUTCOMES

- Collect, access, and download whole bacterial genomes from public repositories
- Investigate and navigate bacterial genomes and their annotation using Artemis
- Identify genomic regions with low/high GC (guanine-cytosine) content, often associated with virulence
- Perform simple comparative analyses between bacterial genomes

SYLLABUS

Week 1

- Multi-FASTA files
- Reference and draft bacterial genomes
- Introduction to Artemis, an open source genome browser

Week 2

- Genome annotation
- Genomic regions defined by GC (guanine-cytosine) content

- Accessing and downloading whole genome sequences
- Pathogenicity islands

Week 3

- Peer-reviewed project: Ecological niche adaptation between two *Mycobacterium* species
- Pseudogenes in *Mycobacterium* species

ACCREDITATION

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