

## Event metadata

Event title	WORKSHOP: Single cell RNAseq analysis in R
Event type	Workshop
Date of event	26 & 27 September 2023
Time of event	1:00 - 4:30 pm AEST
Topic description	Analysis and interpretation of single cell RNAseq (scRNAseq) data requires dedicated workflows. In this hands-on workshop we will show you how to perform single cell analysis using Seurat - an R package for QC, analysis, and exploration of single-cell RNAseq data.
	We will discuss the 'why' behind each step and cover reading in the count data, quality control, filtering, normalisation, clustering, UMAP layout and identification of cluster markers. We will also explore various ways of visualising single cell expression data.
	This workshop is presented by the <u>Australian BioCommons</u> , <u>Queensland Cyber Infrastructure Foundation (QCIF)</u> and the <u>Monash</u> <u>Genomics and Bioinformatics Platform</u> with the assistance of a network of facilitators from the national <u>Bioinformatics Training Cooperative</u> .
Format description	Workshop, online via Zoom over two three and a half hour sessions.
	The lead trainers introduced key concepts and demonstrated the steps involved in the analysis. Participants then moved into breakout rooms where they had the chance to apply these skills with support from facilitators.
	The workshop followed the tutorial linked in the 'Related work' section.
	A breakdown of timings and topics is provided in the schedule.
	Participation was free but subject to application with selection.
	Applications were reviewed by the organising committee.
	Number of participants = 59
	Breakout room size = 10 participants, 1 trainer
Identifier(s)/URL	https://www.biocommons.org.au/events/scrnaseq2023
Licence	Materials are shared under a Creative Commons Attribution 4.0 International agreement unless otherwise stated on the materials
Keywords	Bioinformatics <u>http://edamontology.org/topic_0091</u> Analysis <u>http://edamontology.org/operation_2945</u>



	Transcriptomics <u>http://edamontology.org/topic_3308</u> Single cell RNAseq Seurat R statistical software
Contact	training@biocommons.org.au
Audience	This workshop is for Australian researchers who have or will work on scRNAseq data as part of their projects. You must be associated with an Australian organisation for your application to be considered.
Prerequisites	This workshop will not teach you the basics of R. While you don't need to be an expert, you require some basic familiarity with R. For example you should be able to work with tables of data, know how to load an R package, make basic plots (ideally with ggplot2).
	You should also have basic knowledge of single cell RNA sequencing technology.
Technical	Slack was used to facilitate discussions.
requirements	<ul> <li>Access to the internet, speakers, a webcam, microphone and Zoom.</li> </ul>
	<ul> <li>Participants were provided with access to virtual machines running RStudio via a Nectar Cloud instance. Packages and data were preinstalled</li> </ul>
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It is also drawing from a similar workshop held by Monash Genomics
and Bioinformatics Platform Single-Cell-Workshop, with material here.