snRNA-seq snATAC-seq scRNA and scATAC co-assays

Gene expression matrix



Open chromatin intensity matrix



Embedded scRNA and scATAC datasets

The cell counts **must** be identical in scRNA and scATAC dataset.

- Using scJoint to unify cell type annotations, and pairing individual cells using scOptmatch is highly recommended.
- Make sure the samples can be embedded reasonably,



