

# **BestPracticesSTBiocBook**

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# Welcome

**Package:** BestPracticesSTBiocBook **Authors:** First Last [aut, cre] **Compiled:** 2024-09-25  
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**License:** MIT + file LICENSE

This is the website for the online book ‘**Best Practices for Spatial Transcriptomics Analysis with Bioconductor**’.

This book provides discussion and interactive examples on best practices for computational analysis workflows for spatial transcriptomics data, using the [Bioconductor](#) framework within R. The chapters contain details on individual analysis steps as well as complete example workflows, with interactive example datasets and R code.

The book is organized into several parts, including introductory materials, analysis steps, and example workflows.

Additional details on analysis workflows for non-spatial single-cell data as well as further introductory materials on R and Bioconductor can be found in the related book [Orchestrating Single-Cell Analysis with Bioconductor \(OSCA\)](#).

# Docker image

A Docker image built from this repository is available here:

[ghcr.io/lmweber/bestpracticesstbiocbook](https://ghcr.io/lmweber/bestpracticesstbiocbook)

 Get started now

You can get access to all the packages used in this book in < 1 minute, using this command in a terminal:

---

**Listing 0.1** bash

---

```
docker run -it ghcr.io/lmweber/bestpracticesstbiocbook:devel R
```

---

# RStudio Server

An RStudio Server instance can be initiated from the **Docker** image as follows:

---

**Listing 0.2** bash

---

```
docker run \  
  --volume <local_folder>:<destination_folder> \  
  -e PASSWORD=OHCA \  
  -p 8787:8787 \  
  ghcr.io/lmweber/bestpracticesstbiocbook:devel
```

---

The initiated RStudio Server instance will be available at <https://localhost:8787>.

## Session info

 Click to expand





# **Part I**

## **Introduction**

# 1 Introduction

## 1.1 Overview

Bioconductor

## 1.2 Contents

- 
- 
- 
- 

## 1.3 Scope and who this book is for

Preprocessing

Visium Data

## 1.4 Bioconductor

[Bioconductor](#)

## 1.5 Additional resources

- [Orchestrating Single-Cell Analysis with Bioconductor \(OSCA\)](#)
- [R for Data Science](#)
- [Data Carpentry](#)     [Software Carpentry](#)

- [detailed guide](#)  
[YouTube videos](#)
- [Visium Data Preprocessing](#)

## 1.6 Contributions

[GitHub issues](#)

## References

## 2 Spatial transcriptomics

### 2.1 Overview

of the Year 2020

Method

### 2.2 Sequencing-based platforms

### 2.2.1 10x Genomics Visium

#### 10x Genomics Visium

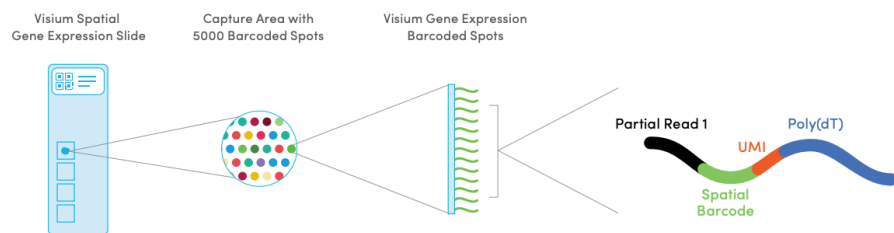


Figure 2.1: Schematic illustrating the 10x Genomics Visium platform. Source: [10x Genomics Visium](#)

### **2.2.2 10x Genomics Visium HD**

[10x Genomics Visium HD](#)

### **2.2.3 Curio Seeker**

[Curio Seeker](#)

## **2.3 Molecule-based platforms**

### **2.3.1 10x Genomics Xenium**

[10x Genomics](#)



### 2.3.2 Vizgen MERSCOPE

Vizgen

### 2.3.3 NanoString CosMx

NanoString

## References

## 3 Bioconductor data classes

### 3.1 Overview

### 3.2 SpatialExperiment class

[SpatialExperiment](#)

[SingleCellExperiment](#)

[Bioconductor vignette](#)

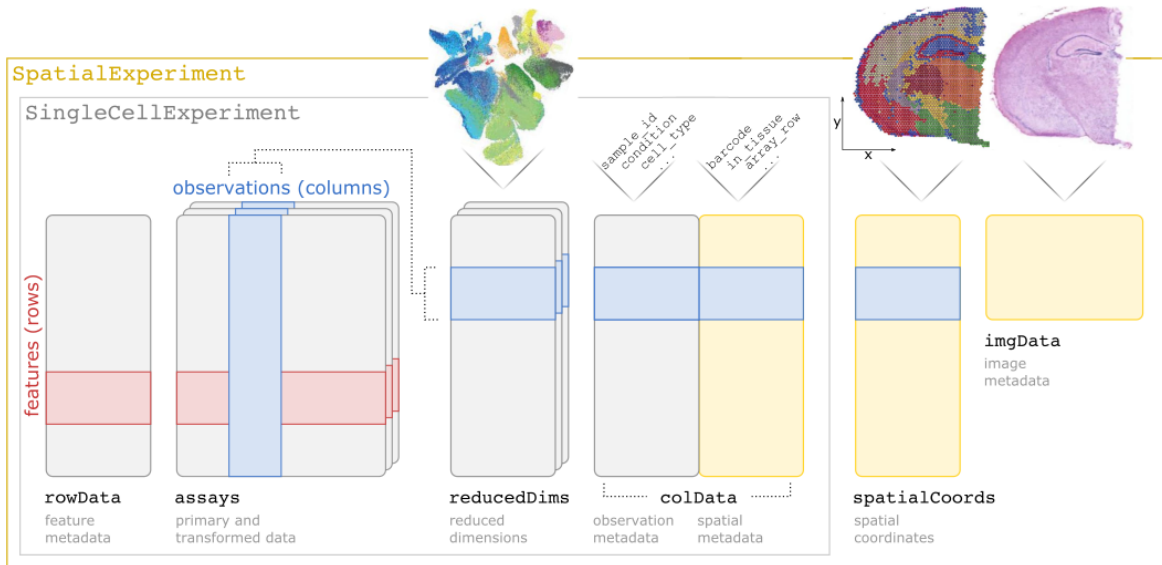


Figure 3.1: Overview of the `SpatialExperiment` data class for storing and manipulating spatial transcriptomics datasets within the Bioconductor framework.

### 3.3 Molecule-based data

#### 3.3.1 MoleculeExperiment

[Bioconductor package](#)

#### 3.3.2 SpatialFeatureExperiment

[Bioconductor package](#)

## References

## **Part II**

# **Analysis steps**

## 4 Analysis steps

3

### 4.1 Save data objects for re-use in later chapters

#### 4.1.1 Human DLPFC dataset

```
# LOAD DATA

library(SpatialExperiment)
library(STexampleData)
spe <- Visium_humanDLPFC()

# save object
library(here)
# if (!dir.exists(here("outputs"))) dir.create(here("outputs"))
# saveRDS(spe, file = here("outputs/spe_load.rds"))
saveRDS(spe, file = here("spe_load.rds"))
```

```

# QUALITY CONTROL (QC)

library(scater)
# subset to keep only spots over tissue
spe <- spe[, colData(spe)$in_tissue == 1]
# identify mitochondrial genes
is_mito <- grepl("(^MT-)|(^mt-)", rowData(spe)$gene_name)
# calculate per-spot QC metrics
spe <- addPerCellQC(spe, subsets = list(mito = is_mito))
# select QC thresholds
qc_lib_size <- colData(spe)$sum < 600
qc_detected <- colData(spe)$detected < 400
qc_mito <- colData(spe)$subsets_mito_percent > 28
qc_cell_count <- colData(spe)$cell_count > 10
# combined set of discarded spots
discard <- qc_lib_size | qc_detected | qc_mito | qc_cell_count
colData(spe)$discard <- discard
# filter low-quality spots
spe <- spe[, !colData(spe)$discard]

# save object
# saveRDS(spe, file = here("outputs/spe_qc.rds"))
saveRDS(spe, file = here("spe_qc.rds"))

```

```

# NORMALIZATION

library(scraper)
# calculate logcounts using library size factors
spe <- logNormCounts(spe)

# save object
# saveRDS(spe, file = here("outputs/spe_logcounts.rds"))
saveRDS(spe, file = here("spe_logcounts.rds"))

```

```

# FEATURE SELECTION

# remove mitochondrial genes
spe <- spe[!is_mito, ]
# fit mean-variance relationship
dec <- modelGeneVar(spe)
# select top HVGs
top_hvgs <- getTopHVGs(dec, prop = 0.1)

```

```
# save object
# saveRDS(spe, file = here("outputs/spe_hvgs.rds"))
# saveRDS(top_hvgs, file = here("outputs/top_hvgs.rds"))
saveRDS(spe, file = here("spe_hvgs.rds"))
saveRDS(top_hvgs, file = here("top_hvgs.rds"))
```

#### # DIMENSIONALITY REDUCTION

```
# compute PCA
set.seed(123)
spe <- runPCA(spe, subset_row = top_hvgs)
# compute UMAP on top 50 PCs
set.seed(123)
spe <- runUMAP(spe, dimred = "PCA")
# update column names
colnames(reducedDim(spe, "UMAP")) <- paste0("UMAP", 1:2)

# save object
# saveRDS(spe, file = here("outputs/spe_reduceddims.rds"))
saveRDS(spe, file = here("spe_reduceddims.rds"))
```

#### # CLUSTERING

```
# graph-based clustering
set.seed(123)
k <- 10
g <- buildSNNGraph(spe, k = k, use.dimred = "PCA")
g_walk <- igraph::cluster_walktrap(g)
clus <- g_walk$membership
colLabels(spe) <- factor(clus)

# save object
# saveRDS(spe, file = here("outputs/spe_cluster.rds"))
saveRDS(spe, file = here("spe_cluster.rds"))
```

## References



## 5 Load data

### 5.1 Overview

3

Preprocessing

Visium Data

STexampleData

### 5.2 Dataset

### 5.3 Load data

STexampleData

```
library(SpatialExperiment)
library(STexampleData)

# load object
spe <- Visium_humanDLPFC()
```

## 5.4 SpatialExperiment object

3

```
# check object
spe
## class: SpatialExperiment
## dim: 33538 4992
## metadata(0):
## assays(1): counts
## rownames(33538): ENSG00000243485 ENSG00000237613 ... ENSG00000277475
##      ENSG00000268674
## rowData names(3): gene_id gene_name feature_type
## colnames(4992): AAACAACGAATAGTTC-1 AAACAAGTATCTCCCA-1 ...
##      TTGTTTGTATTACACG-1 TTGTTTGTGTAAATTC-1
## colData names(8): barcode_id sample_id ... reference cell_count
## reducedDimNames(0):
## mainExpName: NULL
## altExpNames(0):
## spatialCoords names(2) : pxl_col_in_fullres pxl_row_in_fullres
## imgData names(4): sample_id image_id data scaleFactor

# number of genes (rows) and spots (columns)
dim(spe)
## [1] 33538 4992

# names of 'assays'
assayNames(spe)
## [1] "counts"

# row (gene) data
head(rowData(spe))
## DataFrame with 6 rows and 3 columns
```

```
##           gene_id   gene_name   feature_type
##           <character> <character>   <character>
## ENSG00000243485 ENSG00000243485 MIR1302-2HG Gene Expression
## ENSG00000237613 ENSG00000237613   FAM138A Gene Expression
## ENSG00000186092 ENSG00000186092   OR4F5  Gene Expression
## ENSG00000238009 ENSG00000238009 AL627309.1 Gene Expression
## ENSG00000239945 ENSG00000239945 AL627309.3 Gene Expression
## ENSG00000239906 ENSG00000239906 AL627309.2 Gene Expression

# column (spot) data
head(colData(spe))
## DataFrame with 6 rows and 8 columns
##           barcode_id   sample_id in_tissue array_row
##           <character> <character> <integer> <integer>
## AAACAACGAATAGTTC-1 AAACAACGAATAGTTC-1 sample_151673      0      0
## AAACAAGTATCTCCCA-1 AAACAAGTATCTCCCA-1 sample_151673      1     50
## AAACAATCTACTAGCA-1 AAACAATCTACTAGCA-1 sample_151673      1      3
## AAACACCAATAACTGC-1 AAACACCAATAACTGC-1 sample_151673      1     59
## AAACAGAGCGACTCCT-1 AAACAGAGCGACTCCT-1 sample_151673      1     14
## AAACAGCTTTCAGAAG-1 AAACAGCTTTCAGAAG-1 sample_151673      1     43
##           array_col ground_truth   reference cell_count
##           <integer> <character> <character> <integer>
## AAACAACGAATAGTTC-1      16         NA         NA         NA
## AAACAAGTATCTCCCA-1     102      Layer3      Layer3         6
## AAACAATCTACTAGCA-1      43      Layer1      Layer1        16
## AAACACCAATAACTGC-1      19         WM         WM         5
## AAACAGAGCGACTCCT-1      94      Layer3      Layer3         2
## AAACAGCTTTCAGAAG-1       9      Layer5      Layer5         4

# spatial coordinates
head(spatialCoords(spe))
##           pxl_col_in_fullres pxl_row_in_fullres
## AAACAACGAATAGTTC-1          3913          2435
## AAACAAGTATCTCCCA-1          9791          8468
## AAACAATCTACTAGCA-1          5769          2807
## AAACACCAATAACTGC-1          4068          9505
## AAACAGAGCGACTCCT-1          9271          4151
## AAACAGCTTTCAGAAG-1          3393          7583

# image data
imgData(spe)
## DataFrame with 2 rows and 4 columns
```

```
##      sample_id  image_id  data scaleFactor
##      <character> <character> <list>   <numeric>
##  1 sample_151673      lowres  #####    0.0450045
##  2 sample_151673      hires   #####    0.1500150
```

## 5.5 Build object

SpatialExperiment

```
# create data
n_genes <- 200
n_spots <- 100

counts <- matrix(0, nrow = n_genes, ncol = n_spots)

row_data <- DataFrame(
  gene_name = paste0("gene", sprintf("%03d", seq_len(n_genes)))
)

col_data <- DataFrame(
  sample_id = rep("sample01", n_spots)
)

spatial_coords <- matrix(0, nrow = n_spots, ncol = 2)
colnames(spatial_coords) <- c("x", "y")

# create SpatialExperiment object
spe <- SpatialExperiment(
  assays = list(counts = counts),
  colData = col_data,
  rowData = row_data,
  spatialCoords = spatial_coords
)
```

## 5.6 Molecule-based data

3

## References