IGVF CRISPR Jamboree 2024: Perturb-seq Inference (R)

Gene Katsevich

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1 Overview

1.1 Perturb-seq inference

The goal of perturb-seq inference is to quantify the extent to which the perturbation of a given genomic element impacts the expression of a given gene. We allow a range of statistical interpretations of this task. In a frequentist framework, this task can be viewed as testing the null hypothesis that the perturbation of the genomic element has no effect on the gene's expression, or as estimating the effect size of the perturbation on the gene's expression. In a Bayesian framework, this task can be viewed as estimating the posterior probability of the presence of a non-zero effect, or as a posterior mean of the effect size.

1.2 Jamboree goals

The goal of the perturb-seq inference portion of the Jamboree is to implement a number of perturb-seq inference methodologies using common input and output formats. Following the Jamboree, these implementations will be added as modules to a Nextflow pipeline, which will then be used to benchmark their statistical and computational performance. This benchmarking effort will suggest best practices for perturb-seq inference, and will be used to inform the development of the IGVF perturb-seq analysis pipeline.

1.3 Data format overview

The primary input to a perturb-seq inference module is a MuData object, which contains both the perturb-seq data and a set of element-gene pairs for which the inference is to be performed. The output of each method should be the same MuData object, except with an additional table containing one or more measures of association for each element-gene pair. The MuData format is an HDF5-based language-agnostic format compatible with import into both R and Python. Each MuData object will contain a minimal set of fields required for inference, and potentially one or more optional fields that provide additional information. For the purposes of this Jamboree, we have provided MuData objects for subsets of the Gasperini et al (2019) and Papalexi et al (2021) datasets. For each dataset, we have provided a minimal MuData object that contains just the required fields, as well as a more fleshed out object that contains additional optional fields.

1.4 Requested function API, documentation, and demonstration

Please write a function in your language of choice with the following arguments:

- The first argument should be mudata_input_fp, a filepath to a MuData object.
- The second argument should be mudata_output_fp, a filepath to an output MuData object.
- There may be one or more additional arguments specific to your method.

The function should read the MuData object from mudata_input_fp, perform the inference, and write the resulting MuData object to mudata_output_fp (in R, via MuData::readH5MU()). The function should include documentation of any additional arguments used. Furthermore, please include a demonstration of the use of your function on at least one of the sample datasets provided, and a brief discussion of the results.

2 MuData format

Let us walk through the input and output format specifications, from the perspective of R, using a subset of the Gasperini et al (2019) dataset as an example.

```
library(MuData)
library(SummarizedExperiment)
```

2.1 Required input fields

We start with an example of the minimal MuData object required for perturb-seq inference.

```
mudata_input_fp <- "data/inference/gasperini_inference_input_minimal.h5mu"
input_minimal <- readH5MU(mudata_input_fp)
input_minimal</pre>
```

```
## A MultiAssayExperiment object of 2 listed
## experiments with user-defined names and respective classes.
## Containing an ExperimentList class object of length 2:
## [1] gene: SummarizedExperiment with 112 rows and 9704 columns
## [2] guide: SummarizedExperiment with 55 rows and 9704 columns
## Functionality:
## experiments() - obtain the ExperimentList instance
colData() - the primary/phenotype DataFrame
## sampleMap() - the sample coordination DataFrame
## **, `[`, `[[` - extract colData columns, subset, or experiment
## *Format() - convert into a long or wide DataFrame
## assays() - convert ExperimentList to a SimpleList of matrices
## exportClass() - save data to flat files
```

The minimal MuData object for perturb-seq inference contains two modalities: gene and guide.

2.1.1 gene modality

The minimal gene modality needs to just have one assay, whose name is the empty string, containing the RNA UMI counts:

```
input_minimal[['gene']]

## class: SummarizedExperiment
## dim: 112 9704
```

```
## dim: 112 9704
## metadata(0):
## assays(1): ''
## rownames(112): ENSG00000008853 ENSG00000104679 ... ENSG00000198899
## rowData names(0):
## colnames(9704): GCTTGAATCGAATGCT-1_1B_1_SI-GA-F2
## AGCTTGATCGAGAGCA-1_1A_2_SI-GA-E3 ... GGATTACCATGTTGAC-1_2A_4_SI-GA-G5
## GTGCTTCTCGGATGTT-1_2A_1_SI-GA-G2
## colData names(0):
```

To bring it more in line with the typical SummarizedExperiment, we can rename the assay as counts:

```
assayNames(input_minimal[['gene']]) <- 'counts'
input_minimal[['gene']]</pre>
```

```
## class: SummarizedExperiment
## dim: 112 9704
```

```
## metadata(0):
## assays(1): counts
## rownames(112): ENSG00000008853 ENSG00000104679 ... ENSG00000198899
     ENSG00000228253
## rowData names(0):
## colnames(9704): GCTTGAATCGAATGCT-1 1B 1 SI-GA-F2
     AGCTTGATCGAGAGCA-1 1A 2 SI-GA-E3 ... GGATTACCATGTTGAC-1 2A 4 SI-GA-G5
     GTGCTTCTCGGATGTT-1_2A_1_SI-GA-G2
## colData names(0):
assay(input_minimal[['gene']], 'counts')[1:2,1:2]
## 2 x 2 sparse Matrix of class "dgCMatrix"
                   GCTTGAATCGAATGCT-1_1B_1_SI-GA-F2
##
## ENSG00000008853
## ENSG0000104679
##
                   AGCTTGATCGAGAGCA-1_1A_2_SI-GA-E3
## ENSG00000008853
## ENSG0000104679
2.1.2 guide modality
After renaming the first assay of the guide modality, this modality must have assays counts and
guide_assignment containing the gRNA UMI counts and binary gRNA assignments, respectively:
assayNames(input_minimal[['guide']])[[1]] <- 'counts'</pre>
input_minimal[['guide']]
## class: SummarizedExperiment
## dim: 55 9704
## metadata(2): capture_method moi
## assays(2): counts guide_assignment
## rownames(55): ATGTAGAAGGAGACACCGGG GCGCAGAGGCGGATGTAGAG ...
     AATCCTCTAATGGACGAAGA ATATTCAGCAGCTAAAGCAT
## rowData names(2): targeting intended target name
## colnames(9704): GCTTGAATCGAATGCT-1_1B_1_SI-GA-F2
     AGCTTGATCGAGAGCA-1 1A 2 SI-GA-E3 ... GGATTACCATGTTGAC-1 2A 4 SI-GA-G5
     GTGCTTCTCGGATGTT-1_2A_1_SI-GA-G2
##
## colData names(0):
We can view a couple rows and columns of each:
cell ids <- c("GCTTGAATCGAATGCT-1 1B 1 SI-GA-F2",
             "GGTGAAGCACCAGGCT-1_1A_6_SI-GA-E7")
grna ids <- c("GCCCTGCTACCCACTTACAG",
              "ATGTAGAAGGAGACACCGGG")
assay(input_minimal[['guide']], 'counts')[grna_ids, cell_ids]
## 2 x 2 sparse Matrix of class "dgCMatrix"
                        GCTTGAATCGAATGCT-1_1B_1_SI-GA-F2
##
## GCCCTGCTACCCACTTACAG
                                                        9
## ATGTAGAAGGAGACACCGGG
                        GGTGAAGCACCAGGCT-1_1A_6_SI-GA-E7
## GCCCTGCTACCCACTTACAG
```

18

ATGTAGAAGGAGACACCGGG

assay(input_minimal[['guide']], 'guide_assignment')[grna_ids, cell_ids]

```
## 2 x 2 sparse Matrix of class "dgCMatrix"
## GCCCTGCTACCCACTTACAG
## ATGTAGAAGGAGACACCGGG
## GCCCTGCTACCCACTTACAG
## GCCCTGCTACCCACTTACAG
## GCCCTGCTACCCACTTACAG
## ATGTAGAAGGAGACACCGGG
## TGTAGAAGGAGACACCGGG
## ATGTAGAAGGAGACACCGGG
1
```

In addition to the guide UMI counts and assignments, the guide modality must contain certain metadata information. This includes a rowData() data frame containing at least the binary variable targeting (TRUE if the guide targets a genomic element of interest or FALSE if it is safe- or non-targeting) and the string intended_target_name (the name of the genomic element targeted by the guide).

```
rowData(input_minimal[['guide']])[c(1, 2, 21, 22, 31, 32),]
```

```
## DataFrame with 6 rows and 2 columns
##
                           targeting intended_target_name
##
                         <character>
                                               <character>
## ATGTAGAAGGAGACACCGGG
                                TRUE
                                          ENSG00000012660
## GCGCAGAGGCGGATGTAGAG
                                TRUE
                                          ENSG00000012660
## ACACCCTCATTAGAACCCAG
                                TRUE
                                          candidate_enh_1
## TTAAGAGCCTCGGTTCCCCT
                                TRUE
                                          candidate_enh_1
## GACCTCCTGTGATCAGGTGG
                               FALSE
                                            non-targeting
## ATTGGTATCCGTATAAGCAG
                               FALSE
                                            non-targeting
```

Note that the targeting column is a string rather than a Boolean due to type compatibility issues involving R, Python, and HDF5. It can be cast to a Boolean if desired.

Finally, the guide modality must contain metadata() fields called moi (low or high) and capture_method ("CROP-seq" or "direct capture"):

```
metadata(input_minimal[['guide']])
```

```
## $capture_method
## [1] "CROP-seq"
##
## $moi
## [1] "high"
```

2.1.3 Global metadata

The input MuData object is also required to have a global metadata field named pairs_to_test, which is a data frame containing the pairs of elements (specified via intended_target_name) and genes (specified via gene_id) for which the inference is to be performed.

```
metadata(input_minimal)$pairs_to_test |> as.data.frame() |> head()
```

```
## gene_id intended_target_name
## 1 ENSG00000187109 ENSG00000187109
## 2 ENSG00000114850 ENSG00000114850
## 3 ENSG00000134851 ENSG00000134851
## 4 ENSG00000163866 ENSG00000163866
## 5 ENSG00000181610 ENSG00000181610
## 6 ENSG00000113552 ENSG00000113552
```

2.2 Optional input fields

Next we consider optional fields that can be included in the input MuData object.

```
mudata_input_fp = "data/inference/gasperini_inference_input.h5mu"
input_optional = readH5MU(mudata_input_fp)
input_optional

## A MultiAssayExperiment object of 2 listed
## experiments with user-defined names and respective classes.
## Containing an ExperimentList class object of length 2:
## [1] gene: SummarizedExperiment with 112 rows and 9704 columns
```

```
## Functionality:
## experiments() - obtain the ExperimentList instance
```

colData() - the primary/phenotype DataFrame

sampleMap() - the sample coordination DataFrame

`\$`, `[`, `[[` - extract colData columns, subset, or experiment

*Format() - convert into a long or wide DataFrame

assays() - convert ExperimentList to a SimpleList of matrices

[2] guide: SummarizedExperiment with 55 rows and 9704 columns

exportClass() - save data to flat files

2.2.1 gene modality

The MuData object may include cellwise covariates for the gene modality in colData(), such as number of genes with nonzero UMI counts (num_expressed_genes) and total RNA UMIs (total_gene_umis):

```
colData(input_optional[['gene']])
```

```
## DataFrame with 9704 rows and 2 columns
##
                                     num_expressed_genes total_gene_umis
##
                                                <integer>
                                                                 <numeric>
## GCTTGAATCGAATGCT-1_1B_1_SI-GA-F2
                                                                       280
## AGCTTGATCGAGAGCA-1 1A 2 SI-GA-E3
                                                       35
                                                                       192
## CCCAATCTCCTCAATT-1_1B_1_SI-GA-F2
                                                                       781
                                                       41
## CGCGGTACACTGTCGG-1_1A_2_SI-GA-E3
                                                       37
                                                                       189
## GGACGTCTCATGTCTT-1_1B_8_SI-GA-F9
                                                       32
                                                                       262
                                                                       . . .
## CGCTATCTCTATCGCC-1_2A_4_SI-GA-G5
                                                       23
                                                                       203
                                                                       173
## TCACAAGCAGCCTTGG-1_2A_6_SI-GA-G7
                                                       30
## GCTGCAGGTGAAGGCT-1 2B 6 SI-GA-H7
                                                       37
                                                                       428
## GGATTACCATGTTGAC-1_2A_4_SI-GA-G5
                                                       47
                                                                       658
## GTGCTTCTCGGATGTT-1_2A_1_SI-GA-G2
                                                       23
                                                                       166
```

The MuData object may include per-gene metadata in rowData(), such as the HGNC gene symbol (symbol), the gene chromosome (chr), start (gene_start), and end (gene_end) coordinates:

```
rowData(input_optional[['gene']])
```

```
## DataFrame with 112 rows and 4 columns
##
                                 gene_chr gene_start gene_end
                       symbol
                  <character> <character>
                                           <numeric> <numeric>
                                            22844930 22844931
## ENSG0000008853
                      RHOBTB2
                                     chr8
## ENSG0000104679
                       R3HCC1
                                     chr8
                                            23145421 23145422
## ENSG0000104689
                    TNFRSF10A
                                     chr8
                                            23082573 23082574
## ENSG0000120889
                    TNFRSF10B
                                     chr8
                                            22926533 22926534
## ENSG0000120896
                       SORBS3
                                     chr8
                                            22409208 22409209
```

```
## ...
                             . . .
                                         . . .
                                                     . . .
## ENSG0000114850
                           SSR3
                                        chr3 156271913 156271914
                                              195808960 195808961
## ENSG00000072274
                           TFRC
                                        chr3
## ENSG0000134851
                        TMEM165
                                        chr4
                                                56262124
                                                          56262125
## ENSG0000198899
                                                     NaN
                                                                NaN
## ENSG00000228253
                                                     NaN
                                                                NaN
```

2.2.2 guide modality

The MuData object may include cellwise covariates for the guide modality in colData(), such as number of guides with nonzero UMI counts (num_expressed_guides) and total guide UMIs (total_guide_umis):

```
colData(input_optional[['guide']])
```

```
## DataFrame with 9704 rows and 2 columns
##
                                     num_expressed_guides total_guide_umis
##
                                                 <integer>
                                                                   <numeric>
## GCTTGAATCGAATGCT-1_1B_1_SI-GA-F2
                                                                           9
                                                         1
                                                                          18
## AGCTTGATCGAGAGCA-1_1A_2_SI-GA-E3
                                                         1
## CCCAATCTCCTCAATT-1 1B 1 SI-GA-F2
                                                         1
                                                                          24
## CGCGGTACACTGTCGG-1_1A_2_SI-GA-E3
                                                                          26
                                                         1
## GGACGTCTCATGTCTT-1 1B 8 SI-GA-F9
                                                         1
                                                                          12
## CGCTATCTCTATCGCC-1_2A_4_SI-GA-G5
                                                                           5
                                                         1
## TCACAAGCAGCCTTGG-1 2A 6 SI-GA-G7
                                                         1
                                                                          39
## GCTGCAGGTGAAGGCT-1_2B_6_SI-GA-H7
                                                                          21
                                                         1
## GGATTACCATGTTGAC-1_2A_4_SI-GA-G5
                                                         1
                                                                          73
## GTGCTTCTCGGATGTT-1_2A_1_SI-GA-G2
                                                         1
                                                                          12
```

The MuData object may include per-guide metadata in rowData() in addition to the required targeting and intended_target_name fields, such as the chromosome (intended_target_chr), start (intended_target_start), and end (intended_target_end) of the targeted element:

```
rowData(input_optional[['guide']])[c(1, 2, 21, 22, 31, 32),]
```

```
## DataFrame with 6 rows and 5 columns
##
                           targeting intended_target_name intended_target_chr
##
                         <character>
                                               <character>
                                                                    <character>
## ATGTAGAAGGAGACACCGGG
                                TRUE
                                           ENSG00000012660
                                                                           chr6
## GCGCAGAGGCGGATGTAGAG
                                TRUE
                                           ENSG00000012660
                                                                           chr6
## ACACCCTCATTAGAACCCAG
                                TRUE
                                           candidate enh 1
                                                                           chr8
## TTAAGAGCCTCGGTTCCCCT
                                TRUE
                                           candidate enh 1
                                                                           chr8
## GACCTCCTGTGATCAGGTGG
                               FALSE
                                            non-targeting
## ATTGGTATCCGTATAAGCAG
                               FALSE
                                            non-targeting
##
                         intended_target_start intended_target_end
##
                                     <numeric>
                                                          <numeric>
## ATGTAGAAGGAGACACCGGG
                                      53213723
                                                           53213738
## GCGCAGAGGCGGATGTAGAG
                                      53213738
                                                           53213754
## ACACCCTCATTAGAACCCAG
                                      23366136
                                                           23366564
## TTAAGAGCCTCGGTTCCCCT
                                      23366564
                                                           23366992
## GACCTCCTGTGATCAGGTGG
                                             -9
                                                                  -9
## ATTGGTATCCGTATAAGCAG
                                             -9
                                                                  -9
```

2.2.3 Global metadata

Optionally, the MuData input object can contain a global colData() containing cell-level information that is not specific to modality, such as batch information. Here is what it looks like for the Gasperini data:

colData(input_optional)

```
## DataFrame with 9704 rows and 3 columns
##
                                      prep_batch
                                                   within_batch_chip
##
                                     <character>
                                                         <character>
## GCTTGAATCGAATGCT-1_1B_1_SI-GA-F2 prep_batch_1 within_batch_chip_B
## AGCTTGATCGAGAGCA-1_1A_2_SI-GA-E3 prep_batch_1 within_batch_chip_A
## CCCAATCTCCTCAATT-1_1B_1_SI-GA-F2 prep_batch_1 within_batch_chip_B
## CGCGGTACACTGTCGG-1_1A_2_SI-GA-E3 prep_batch_1 within_batch_chip_A
## GGACGTCTCATGTCTT-1 1B 8 SI-GA-F9 prep batch 1 within batch chip B
## ...
## CGCTATCTCTATCGCC-1 2A 4 SI-GA-G5 prep batch 2 within batch chip A
## TCACAAGCAGCCTTGG-1_2A_6_SI-GA-G7 prep_batch_2 within_batch_chip_A
## GCTGCAGGTGAAGGCT-1_2B_6_SI-GA-H7 prep_batch_2 within_batch_chip_B
## GGATTACCATGTTGAC-1_2A_4_SI-GA-G5 prep_batch_2 within_batch_chip_A
## GTGCTTCTCGGATGTT-1_2A_1_SI-GA-G2 prep_batch_2 within_batch_chip_A
##
                                      within_chip_lane
                                           <character>
##
## GCTTGAATCGAATGCT-1_1B_1_SI-GA-F2 within_chip_lane_1
## AGCTTGATCGAGAGCA-1_1A_2_SI-GA-E3 within_chip_lane_2
## CCCAATCTCCTCAATT-1_1B_1_SI-GA-F2 within_chip_lane_1
## CGCGGTACACTGTCGG-1_1A_2_SI-GA-E3 within_chip_lane_2
## GGACGTCTCATGTCTT-1_1B_8_SI-GA-F9 within_chip_lane_8
## CGCTATCTCTATCGCC-1_2A_4_SI-GA-G5 within_chip_lane_4
## TCACAAGCAGCCTTGG-1_2A_6_SI-GA-G7 within_chip_lane_6
## GCTGCAGGTGAAGGCT-1 2B 6 SI-GA-H7 within chip lane 6
## GGATTACCATGTTGAC-1_2A_4_SI-GA-G5 within_chip_lane_4
## GTGCTTCTCGGATGTT-1_2A_1_SI-GA-G2 within_chip_lane_1
```

2.2.4 Pairs to test

Optionally, the pairs_to_test field of the global metadata can have a third column: pair_type:

```
metadata(input_optional)$pairs_to_test |> as.data.frame() |> head()
```

```
## gene_id intended_target_name pair_type
## 1 ENSG00000187109 ENSG00000187109 positive_control
## 2 ENSG00000114850 ENSG00000114850 positive_control
## 3 ENSG00000134851 ENSG00000134851 positive_control
## 4 ENSG00000163866 ENSG00000163866 positive_control
## 5 ENSG00000181610 ENSG00000181610 positive_control
## 6 ENSG00000113552 ENSG00000113552 positive_control
```

This optional column classifies pairs based on whether they are intended to be positive controls (an association is known to exist), negative controls (an association is known not to exist), or discovery pairs (pairs where it is unknown whether an association exists). This information need not be used by the inference module, but it is useful for downstream analysis.

2.3 Output fields

The output should be the same MuData object as the input, with the addition of a test_results field to the global metadata:

```
mudata_output_fp <- "data/inference/gasperini_inference_output.h5mu"
output_optional <- readH5MU(mudata_output_fp)</pre>
```

```
output_optional
```

```
## A MultiAssayExperiment object of 2 listed
   experiments with user-defined names and respective classes.
## Containing an ExperimentList class object of length 2:
## [1] gene: SummarizedExperiment with 112 rows and 9704 columns
## [2] guide: SummarizedExperiment with 55 rows and 9704 columns
## Functionality:
## experiments() - obtain the ExperimentList instance
## colData() - the primary/phenotype DataFrame
## sampleMap() - the sample coordination DataFrame
## `$`, `[`, `[[` - extract colData columns, subset, or experiment
## *Format() - convert into a long or wide DataFrame
## assays() - convert ExperimentList to a SimpleList of matrices
## exportClass() - save data to flat files
metadata(output_optional)$test_results |> as.data.frame() |> head()
            gene id intended target name
                                             log2_fc
                                                          p_value
                                                                         pair_type
## 1 ENSG00000187109
                          ENSG00000187109 -0.7743672 3.217223e-85 positive_control
## 2 ENSG00000114850
                          ENSG00000114850 -1.8495718 2.414163e-79 positive_control
## 3 ENSG00000134851
                          ENSG00000134851 -0.8938597 4.309833e-50 positive_control
## 4 ENSG0000163866
                          ENSG00000163866 -1.2236996 4.704066e-49 positive control
## 5 ENSG00000181610
                          ENSG00000181610 -1.3142850 3.766690e-42 positive_control
## 6 ENSG00000113552
                          ENSG00000113552 -1.6785978 2.554175e-39 positive control
```

This is a data frame containing the same columns as the pairs_to_test data frame, plus at least one column containing a measure of the association for each pair. These columns can be p_value, log2_fc, posterior_probability, or any other measure of association.

3 Sample submission

Here we present a sample Jamboree submission.

3.1 Function

Here is a sample function that computes a p-value based on a Wilcoxon test:

```
#' Wilcoxon test
#'

#' @param mudata_input_fp Path to input MuData
#' @param mudata_output_fp Path to output MuData
#' @param side The sidedness of the test (`left`, `right`, or `both`)
#'

#' @return This function is called for its side effect of writing the output MuData.
compute_wilcoxon_test <- function(mudata_input_fp, mudata_output_fp, side) {
    # Read input MuData
    mudata <- MuData::readH5MU(mudata_input_fp)
    # Rename primary assay to 'counts'
    if(is.null(SummarizedExperiment::assayNames(mudata[['gene']]))){
        SummarizedExperiment::assayNames(mudata[['gene']]))[[1]] <- 'counts'
    }
    SummarizedExperiment::assayNames(mudata[['gene']])[[1]] <- 'counts'
}</pre>
```

```
# Extract pairs to test and MOI
pairs_to_test <- MultiAssayExperiment::metadata(input_minimal)$pairs_to_test |>
  as.data.frame()
moi <- MultiAssayExperiment::metadata(input minimal[["guide"]])$moi</pre>
# In low-MOI case, extract control cells as those containing an NT qRNA
if (moi == "low") {
 non_targeting_guides <- SummarizedExperiment::rowData(mudata[["guide"]]) |>
    as.data.frame() |>
    dplyr::filter(targeting == "FALSE") |>
    rownames()
 nt_grna_presence <- SummarizedExperiment::assay(</pre>
    mudata[["guide"]],
    "guide_assignment"
 )[non_targeting_guides, ] |>
    apply(MARGIN = 2, FUN = max)
  control_cells <- names(nt_grna_presence)[nt_grna_presence == 1]</pre>
# Initialize test results data frame based on pairs to test
test_results <- pairs_to_test |>
 dplyr::mutate(p_value = NA_real_)
# Carry out the Wilcoxon test for each pair
for (pair_idx in 1:nrow(pairs_to_test)) {
  # Extract the gene and element to be tested
 gene_id <- pairs_to_test[pair_idx, "gene_id"]</pre>
  intended_target_name <- pairs_to_test[pair_idx, "intended_target_name"]</pre>
  # Find the set of treatment cells (i.e. cells with element targeted)
  grnas_targeting_element <- SummarizedExperiment::rowData(mudata[["guide"]]) |>
    as.data.frame() |>
    dplyr::filter(intended_target_name == !!intended_target_name) |>
    rownames()
  element_targeted <- assay(</pre>
    mudata[["guide"]],
    "guide_assignment"
 )[grnas_targeting_element, ] |>
    apply(MARGIN = 2, FUN = max)
  treatment_cells <- names(element_targeted)[element_targeted == 1]</pre>
  # Set controls cells using the complement set in high MOI
 if (moi == "high") {
    control_cells <- names(element_targeted)[element_targeted == 0]</pre>
 }
  # extract expressions for treatment and control cells
 treatment_expressions <- SummarizedExperiment::assay(</pre>
    mudata[["gene"]],
    "counts"
  )[gene_id, treatment_cells]
  control_expressions <- SummarizedExperiment::assay(</pre>
    mudata[["gene"]],
    "counts"
 )[gene_id, control_cells]
  # Map `side` argument to `alternative` argument required by `wilcox.test()`
  alternative <- switch(side,
    left = "less",
    right = "greater",
```

```
both = "two.sided"
)

# Carry out the Wilcoxon test

test_results[pair_idx, "p_value"] <- stats::wilcox.test(
    x = treatment_expressions,
    y = control_expressions,
    alternative = alternative
) $p.value
}

# Add output to MuData and write to disk
mudata_output <- mudata
MultiAssayExperiment::metadata(mudata_output)$test_results <- test_results
MuData::writeH5MU(mudata_output, mudata_output_fp)
}</pre>
```

3.2 Demonstration

Here is a demonstration of this function on the Gasperini data:

```
# Compute Wilcoxon test on the Gasperini data
compute_wilcoxon_test(
 mudata_input_fp = "data/inference/gasperini_inference_input.h5mu",
 mudata_output_fp = "data/inference/gasperini_inference_output_wilcoxon.h5mu",
  side = "left"
)
# Read results from disk
output_wilcoxon <- MuData::readH5MU("data/inference/gasperini_inference_output_wilcoxon.h5mu")
# Print results
metadata(output_wilcoxon)$test_results |> as.data.frame() |> head()
            gene_id intended_target_name
                                                p_value
## 1 ENSG00000187109
                          ENSG00000187109 3.030394e-72
                          ENSG00000114850 4.672505e-128
## 2 ENSG00000114850
## 3 ENSG00000134851
                         ENSG00000134851 6.398416e-49
## 4 ENSG00000163866
                         ENSG00000163866 4.183842e-68
                         ENSG00000181610 3.903196e-65
## 5 ENSG00000181610
## 6 ENSG00000113552
                         ENSG00000113552 1.360795e-60
```