## CRISPR Jamboree 2024: Inference

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# 1 Overview of datasets, inputs, and outputs

I propose the following for the inference module:

- Datasets: We should test on at least one high-MOI and one low-MOI dataset. I propose we try subsets of the Gasperini (high-MOI) and Papalexi (low-MOI) datasets, which are built into the sceptre package. These datasets are available on GitHub.
- Input: The input will be a MuData object with a certain set of minimum required fields (see more below), the sidedness of the test, and any additional arguments to the analysis method.
- Output: The output will be an updated MuData object, with the p-values and log fold-change estimates added in the uns field.

#### 2 MuData structure

We will need to specify precisely how the MuData object should be formatted. The current MuData formatting guidelines do not specify naming conventions precisely enough. In what follows, I propose an example of a more precise specification of the MuData object structure. To illustrate, I will use the subset of the Gasperini dataset.

```
import mudata as md
import pandas as pd
gasperini data = md.read h5mu("data/gasperini subset.h5mu")
gasperini_data
## MuData object with n_obs \times n_vars = 44308 \times 716
##
     obs:
             'batch'
##
             'inference_results', 'moi'
     uns:
##
     3 modalities
##
                 44308 x 526
       gene:
                 'symbol'
##
         var:
                 44308 x 95
##
       guide:
##
                 'targeting', 'guide_target', 'guide_chr', 'guide_start', 'guide_end'
##
       guide assignment:
                              44308 x 95
```

We can see that this object has three modalities, named gene (for gene expression), guide (for gRNA expression), and guide\_assignment (for binary gRNA-to-cell assignments). We need to make sure these names are standardized.

#### 2.1 Gene expression modality

The gene expression modality (gene) should contain the raw RNA UMI counts; we will leave any normalization to each of the individual analysis methods. The variable names are the ENSEMBL gene IDs:

```
gasperini_data['gene'].var_names[:5].tolist()

## ['ENSG00000069275', 'ENSG00000117222', 'ENSG00000117266', 'ENSG00000117280', 'ENSG00000133059']

The gene names can be optionally be provided in a variable called symbol:

gasperini_data['gene'].var

## symbol
```

```
## ENSG00000069275
                     NUCKS1
## ENSG0000117222
                      RBBP5
## ENSG0000117266
                      CDK18
## ENSG0000117280
                      RAB29
## ENSG0000133059
                      DSTYK
## ENSG0000155380
                    SLC16A1
## ENSG0000196683
                      TOMM7
## ENSG0000176890
                       TYMS
## ENSG0000198786
                     MT-ND5
## ENSG0000198840
                     MT-ND3
##
## [526 rows x 1 columns]
```

We need to choose a standardized name for the variable storing the gene names; I chose symbol because this is what DACC appears to use.

## 2.2 gRNA expression modality

The gRNA expression modality (guide) should contain the raw gRNA UMI counts; we will leave any normalization to each of the individual analysis methods. Within the guide modality, the variable names are the gRNA IDs:

```
gasperini_data['guide'].var_names[:5].tolist()
```

```
## ['grna_CCGGGCG', 'grna_TGGCGGC', 'grna_AAGGCCG', 'grna_GACGCCG', 'grna_CACACCC']
```

The variables in this modality are similar to the per-guide metadata format we developed:

```
gasperini_data['guide'].var
```

```
##
                  targeting
                                 guide target guide chr
                                                                          guide end
                                                           guide start
## grna_CCGGGCG
                              ENSG00000069482
                                                   chr11
                                                            68451943.0
                                                                         68451958.0
                           1
## grna_TGGCGGC
                              ENSG00000069482
                                                    chr11
                                                            68451958.0
                                                                         68451974.0
                           1
## grna_AAGGCCG
                           1
                              ENSG0000100316
                                                    chr22
                                                            39715775.0
                                                                         39715790.0
## grna_GACGCCG
                           1
                              ENSG00000100316
                                                    chr22
                                                             39715790.0
                                                                         39715806.0
                                                            44829255.0
## grna_CACACCC
                              ENSG00000104131
                                                                         44829270.0
                           1
                                                    chr15
## ...
                         . . .
                                                      . . .
                                                                    . . .
                                                                                 . . .
## grna_ATTAGCA
                           0
                                                                   -9.0
                                                                                -9.0
                                non-targeting
                                                                                -9.0
## grna_AGATACC
                           0
                                non-targeting
                                                                   -9.0
## grna_ATATGTA
                           0
                                                                                -9.0
                                non-targeting
                                                                   -9.0
## grna_GTAGCCT
                           0
                                non-targeting
                                                                   -9.0
                                                                                -9.0
                           0
                                                                                -9.0
## grna_TTAGCTT
                                non-targeting
                                                                   -9.0
##
## [95 rows x 5 columns]
```

First, we must specify for each gRNA whether it is targeting. (Here this boolean variable shows up as 0/1; probably we should change this.) Next, we must specify for each gRNA what exactly it targets (which gene, which putative enhancer, etc). This is probably similar to intended\_target\_name in the per-guide metadata

format but I wasn't sure, so I named it <code>guide\_target</code> (we should come to a consensus on this). The reason this is required is because we it is most meaningful to test for associations between targeted elements and gene expression rather than between individual <code>gRNAs</code> and gene expression. Therefore, we must know which guides target the same element. We should have a reserved string for the <code>guide\_target</code> for non-targeting guides; I propose non-targeting. Optionally, we can specify the genomic coordinates of the region targeted by the guide. We should have standard placeholder values genomic coordinates of non-targeting guides. I propose the empty string for chromosome and <code>-9</code> for start and end coordinates.

### 2.3 gRNA assignment modality

The gRNA assignment modality (guide\_assignment) should contain the binary gRNA-to-cell assignments. The variable names are again the gRNA IDs:

```
gasperini_data['guide_assignment'].var_names[:5].tolist()
```

```
## ['grna_CCGGGCG', 'grna_TGGCGGC', 'grna_AAGGCCG', 'grna_GACGCCG', 'grna_CACACCC']
```

There are no required vars for this modality, because the relevant metadata are already in the guide modality.

#### 2.4 Other fields

There are two other fields of the gasperini\_data object, obs and uns. The obs field contains cell metadata not specific to any modality. The most important such metadata is batch.

```
gasperini_data.obs
```

```
##
               batch
## cell_1
                  b1
## cell 2
                  b1
## cell_3
                  b1
## cell_4
                  b1
## cell_5
                  b1
## ...
## cell_44304
                  b2
## cell_44305
                  b2
## cell_44306
                  b2
                  b2
## cell_44307
## cell_44308
                  b2
##
## [44308 rows x 1 columns]
```

I propose we have a required variable called batch that specifies the batch for each cell. Even if the data only has one batch, we can have a variable with just one value. The other field is uns, which contains unstructured metadata. I have included two fields in uns: moi and inference\_results. The moi field specifies the MOI of the experiment (high or low):

```
gasperini_data.uns['moi'][0]
```

```
## 'high'
```

I propose for this to be a mandatory field. The inference\_results field is where the results of the inference will be stored:

```
pd.set_option('display.max_columns', None)
pd.DataFrame(gasperini_data.uns['inference_results'])
```

```
##
                                           log_2_FC p_value
                gene_id
                              grna_target
                                                                      pair_type
## 0
        ENSG00000069482
                         ENSG00000069482
                                                -9.0
                                                         -9.0
                                                               positive_control
## 1
        ENSG00000100316
                         ENSG00000100316
                                                -9.0
                                                               positive_control
```

```
## 2
        ENSG00000104131
                         ENSG00000104131
                                               -9.0
                                                         -9.0 positive_control
## 3
        ENSG00000122026 ENSG00000122026
                                               -9.0
                                                         -9.0
                                                              positive_control
## 4
        ENSG00000135821
                         ENSG00000135821
                                               -9.0
                                                         -9.0
                                                              positive_control
##
                                                                      discovery
## 615
        ENSG00000131094
                         candidate_enh_8
                                               -9.0
                                                         -9.0
## 616
        ENSG00000136448
                         candidate enh 8
                                               -9.0
                                                         -9.0
                                                                      discovery
        ENSG00000172992
                         candidate enh 8
                                               -9.0
                                                         -9.0
                                                                      discovery
## 617
                         candidate_enh_8
## 618
        ENSG00000181513
                                               -9.0
                                                         -9.0
                                                                      discovery
                         candidate_enh_8
## 619
        ENSG00000161714
                                               -9.0
                                                         -9.0
                                                                      discovery
##
## [620 rows x 5 columns]
```

The columns <code>gene\_id</code> and <code>grna\_target</code> specify what pairs of gene and targeted element to test for association. The columns <code>log\_2\_FC</code> and <code>p\_value</code> will be filled in by the inference method, and should be initialized with placeholders such as <code>-9</code>. Finally, the optional column <code>pair\_type</code> specifies the type of pair being tested, e.g. positive control or discovery pair.

### 3 Items for discussion

- 1. We need to work together to settle on the precise specification of the MuData format, not just for the inference task but also for upstream tasks like gRNA assignment.
- 2. Do we agree that we should test for association between *targeted elements* and gene expression rather than between *individual gRNAs* and gene expression? Should we also have an option to test the latter? I think the latter is adding unnecessary complication at this stage.