#### main

August 21, 2021

# 1 Generate supplementary data for DEGs

```
[1]: import functools
import numpy as np
import pandas as pd
from scipy.stats import fisher_exact
from statsmodels.stats.multitest import multipletests
```

#### 1.1 Function

# 1.1.1 Cached functions

#### 1.1.2 Simple functions

```
return ds
def print_overlap(feature, tissue, ds):
    dx = get_de()[(get_de()["adj.P.Val"] < 0.05) &
                  (get_de()["Type"] == feature_map(feature)) &
                  (get de()["Tissue"] == tissue)].copy()
    dft = ds[(ds["Tissue"] == tissue)].copy()
    overlap = len(set(dx.Symbol) & set(dft.gene))
    print("There are \{\} (\{:.1\%\}) DS overlapping DE \{\}."
          .format(overlap, overlap/len(dft.gene.unique()), feature))
def subset_de(feature, tissue):
    dx = get_de()[(get_de()["Type"] == feature_map(feature)) &
                  (get_de()["Tissue"] == tissue)].copy()
    return dx.loc[:, ["Symbol", "adj.P.Val"]].dropna()
def merge_data(feature, tissue, ds):
    df = ds[(ds["Tissue"] == tissue)].copy()
    return df.loc[:, ["gene", "FDR", "annotation"]].

drop_duplicates(subset="gene")
\

             .merge(subset_de(feature, tissue), left_on="gene",_
→right_on="Symbol", how="outer")\
             .fillna(1)
def cal_fishers(feature, tissue, ds):
    dt = merge data(feature, tissue, ds)
    table = [[np.sum((dt['adj.P.Val']<0.05) & ((dt['FDR']<0.05))),
              np.sum((dt['adj.P.Val']<0.05) & ((dt['FDR']>=0.05)))],
             [np.sum((dt['adj.P.Val']>=0.05) & ((dt['FDR']<0.05))),
              np.sum((dt['adj.P.Val']>=0.05) & ((dt['FDR']>=0.05)))]]
    #print(table)
    return fisher_exact(table)
```

## 1.2 Summary

```
[4]: caudate = get_tissues_DS("Caudate")
    dlpfc = get_tissues_DS("DLPFC")
    gyrus = get_tissues_DS("Dentate Gyrus")
    hippo = get_tissues_DS("Hippocampus")
    ds = pd.concat([caudate, dlpfc, gyrus, hippo], axis=0)
    print(ds.shape)
    ds.head(2)
```

```
(5233, 9)
[4]:
          clusterID N
                                              coord gene annotation
                                                                               FDR \
     0 clu_128031_- 14 chr12:124911899-124913724 UBC
                                                                      0.000000e+00
                                                            cryptic
     1 clu 105375 ? 13 chr12:124911899-124913724 UBC
                                                            cryptic 2.510000e-170
          chr Type
                    Tissue
     0 chr12 DTU Caudate
     1 chr12 DTU Caudate
[5]: ds.groupby(["Tissue"]).size()
[5]: Tissue
     Caudate
                      1901
    DLPFC
                      1345
    Dentate Gyrus
                      655
                      1332
    Hippocampus
    dtype: int64
    1.2.1 Save files
[6]: ds.to_csv("diffSplicing_ancestry_FDR05_4regions.tsv", sep='\t', index=False)
    1.3 Overlap with DE
[7]: for tissue in ["Caudate", "Dentate Gyrus", "DLPFC", "Hippocampus"]:
        print(tissue)
        for feature in ["genes", "transcripts", "exons", "junctions"]:
            print_overlap(feature, tissue, ds)
        print("")
    Caudate
    There are 357 (23.2%) DS overlapping DE genes.
    There are 594 (38.5%) DS overlapping DE transcripts.
    There are 684 (44.4%) DS overlapping DE exons.
    There are 654 (42.4%) DS overlapping DE junctions.
    Dentate Gyrus
    There are 48 (9.0%) DS overlapping DE genes.
    There are 62 (11.7%) DS overlapping DE transcripts.
    There are 89 (16.7%) DS overlapping DE exons.
    There are 92 (17.3%) DS overlapping DE junctions.
    DLPFC
    There are 239 (21.6%) DS overlapping DE genes.
    There are 388 (35.1%) DS overlapping DE transcripts.
    There are 479 (43.3%) DS overlapping DE exons.
    There are 411 (37.2%) DS overlapping DE junctions.
```

```
Hippocampus
There are 216 (19.2%) DS overlapping DE genes.
There are 399 (35.5%) DS overlapping DE transcripts.
There are 513 (45.6%) DS overlapping DE exons.
There are 410 (36.5%) DS overlapping DE junctions.
```

## 1.4 Enrichment analysis

```
[8]: | feature_lt = []; pval_lt = []; oddratio_lt = []; tissue_lt = []
     for tissue in ["Caudate", "Dentate Gyrus", "DLPFC", "Hippocampus"]:
         print(tissue)
         for feature in ["genes", "transcripts", "exons", "junctions"]:
             oddratio, pval = cal_fishers(feature, tissue, ds)
             feature_lt.append(feature_map(feature))
             pval_lt.append(pval)
             oddratio_lt.append(oddratio)
             tissue_lt.append(tissue)
             print("Enrichment of DS within DE for {}:\nOdd Ratio: {:.2f}; P-value:
      \hookrightarrow{:.1e}\n"\
                   .format(feature_map(feature), oddratio, pval))
         print("")
     pval_lt = [1e-323 if x == 0 else x for x in pval_lt]
     _, fdr, _, _ = multipletests(pval_lt, method='fdr_bh')
     df = pd.DataFrame({"Tissue": tissue_lt, "Feature": feature_lt,
                        "OR": oddratio_lt, "PValue": pval_lt, "FDR": fdr})
     df.to_csv('diffSplice_enrichment_analysis.txt', sep='\t', index=False)
    Caudate
    Enrichment of DS within DE for Gene:
    Odd Ratio: 2.11; P-value: 1.1e-27
    Enrichment of DS within DE for Transcript:
    Odd Ratio: 2.84; P-value: 1.5e-139
    Enrichment of DS within DE for Exon:
    Odd Ratio: 2.77; P-value: 0.0e+00
    Enrichment of DS within DE for Junction:
    Odd Ratio: 3.38: P-value: 0.0e+00
    Dentate Gyrus
    Enrichment of DS within DE for Gene:
    Odd Ratio: 2.66; P-value: 2.1e-08
    Enrichment of DS within DE for Transcript:
```

```
Odd Ratio: 4.17; P-value: 1.0e-23
Enrichment of DS within DE for Exon:
Odd Ratio: 3.90; P-value: 7.2e-106
Enrichment of DS within DE for Junction:
Odd Ratio: 5.08; P-value: 2.7e-66
DLPFC
Enrichment of DS within DE for Gene:
Odd Ratio: 1.98; P-value: 2.9e-17
Enrichment of DS within DE for Transcript:
Odd Ratio: 3.32; P-value: 4.8e-113
Enrichment of DS within DE for Exon:
Odd Ratio: 3.11; P-value: 0.0e+00
Enrichment of DS within DE for Junction:
Odd Ratio: 4.06; P-value: 1.2e-252
Hippocampus
Enrichment of DS within DE for Gene:
Odd Ratio: 1.57; P-value: 3.4e-08
Enrichment of DS within DE for Transcript:
Odd Ratio: 2.65; P-value: 1.6e-83
Enrichment of DS within DE for Exon:
Odd Ratio: 2.52; P-value: 0.0e+00
Enrichment of DS within DE for Junction:
Odd Ratio: 3.50; P-value: 8.8e-221
```

[]: