main

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1 Extract unique male specific SZ-associated genes

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[1]: import functools
     import numpy as np
     import pandas as pd
     from os import environ
     from gtfparse import read_gtf
     from scipy.stats import mannwhitneyu
     from statsmodels.stats.multitest import fdrcorrection
[2]: environ['NUMEXPR MAX THREADS'] = '16'
[3]: @functools.lru_cache()
     def get_res_df(feature):
        return pd.read_csv('../../../interaction_model/hippocampus/_m/%s/
     →residualized_expression.tsv' %
                            feature, sep='\t').T
     @functools.lru_cache()
     def get_pheno_df():
        return pd.read_csv('/ceph/projects/v3_phase3_paper/inputs/phenotypes/_m/
     ⇔hippo_phenotypes.csv',
                            index_col=0)
     @functools.lru_cache()
     def get_res_pheno_df(feature):
        return pd.merge(get_pheno_df(), get_res_df(feature), left_index=True,_u
     →right index=True)
     @functools.lru_cache()
     def get_gtf(gtf_file):
        return read_gtf(gtf_file)
```

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[4]: def map_features(feature):
         return {"genes": "gene", 'transcripts': 'tx',
                 'exons': 'exon', 'junctions': 'jxn'}[feature]
     def gene_annot(feature):
         gtf_file = '/ceph/genome/human/gencode25/gtf.CHR/_m/gencode.v25.annotation.
     ⇔gtf'
         gtf0 = get_gtf(gtf_file)
         gtf = gtf0[(gtf0["feature"] == feature)]
         return gtf[["gene_id", "gene_name", "transcript_id", "exon_id", "gene_type",
                     "seqname", "start", "end", "strand"]]
     def get_de(feature):
         f = pd.read_csv('../../female_analysis/_m/%s/diffExpr_szVctl_full.txt' %__

→feature,
                         sep='\t', index_col=0)\
               .rename(columns={'gencodeGeneID': 'gencodeID'})
         m = pd.read_csv('../../male_analysis/_m/%s/diffExpr_szVctl_full.txt' %__

→feature,
                         sep='\t', index_col=0)\
               .rename(columns={'gencodeGeneID': 'gencodeID'})
         return f, m
     def get unique(x, y, thres=0.05):
         return x.merge(pd.DataFrame(index = list(set(x[(x['adj.P.Val'] <= thres)].</pre>
      ⇒index) -
                                                  set(y[(y['adj.P.Val'] <= thres)].</pre>
      →index))),
                        left_index=True, right_index=True)
     def subset_sz_female(feature):
         df = get_res_pheno_df(feature)
         ctl = df[(df['Dx'] == 'Control') & (df['Sex'] == 'F')].copy()
         sz = df[(df['Dx'] == 'Schizo') & (df['Sex'] == 'F')].copy()
         return ctl, sz
     def add_pvals_adjustPval(feature, df):
         ctl, sz = subset_sz_female(feature)
         f pval = []
         for gene_id in df.Feature:
             stat, pval = mannwhitneyu(ctl[gene_id], sz[gene_id])
             f_pval.append(pval)
```

1.1 Genes

```
[5]: gtf_annot = gene_annot('gene')
    INFO:root:Extracted GTF attributes: ['gene_id', 'gene_type', 'gene_status',
    'gene_name', 'level', 'havana_gene', 'transcript_id', 'transcript_type',
    'transcript_status', 'transcript_name', 'transcript_support_level', 'tag',
    'havana transcript', 'exon number', 'exon id', 'ont', 'protein id', 'ccdsid']
[6]: f, m = get_de('genes')
     m['Feature'] = m.index
     #qenes = qet_unique(qet_unique(m, f), a)
     genes = get_unique(m, f)
     genes = pd.merge(gtf_annot[['gene_id', 'seqname']], genes, left_on='gene_id',
                      right_on='Feature', how='right').rename(columns={'seqname':_
     genes = genes[['Feature', 'gencodeID', 'Symbol', 'ensemblID',
                    'Chrom', 'logFC', 't', 'adj.P.Val']].sort_values('adj.P.Val')
     genes = add_pvals_adjustPval('genes', genes)
     genes = genes[~(genes['Female Pval'] <= 0.05)].sort_values('adj.P.Val') ##__
     \hookrightarrowStringents
     genes['Type'] = 'gene'
     genes.shape
[6]: (104, 10)
```

1.2 Transcripts

[8]: (17, 10)

1.2.1 Exons

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[9]: gtf_annot = gene_annot('exon')
gtf_annot['ensemblID'] = gtf_annot.gene_id.str.replace('\\.\d+', '', regex=True)
```

[10]: (113, 10)

1.2.2 Junctions

```
juncs.shape
[11]: (3, 10)
     1.3 DE summary
     1.3.1 DE (feature)
[12]: gg = len(set(genes.index))
      tt = len(set(trans.index))
      ee = len(set(exons.index))
      jj = len(set(juncs.index))
      print("\nGene:\t\t%d\nTranscript:\t%d\nExon:\t\t%d\nJunction:\t%d" %
            (gg, tt, ee, jj))
     Gene:
                     104
     Transcript:
                     17
     Exon:
                     113
     Junction:
                     3
     DE (EnsemblID)
[13]: gg = len(set(genes['ensemblID']))
      tt = len(set(trans['ensemblID']))
      ee = len(set(exons['ensemblID']))
      jj = len(set(juncs['ensemblID']))
      print("\nGene:\t\t%d\nTranscript:\t%d\nExon:\t\t%d\nJunction:\t%d" %
            (gg, tt, ee, jj))
     Gene:
                     104
     Transcript:
                     15
     Exon:
                     50
     Junction:
                     3
     DE (Gene Symbol)
[14]: gg = len(set(genes['Symbol']))
      tt = len(set(trans['Symbol']))
      ee = len(set(exons['Symbol']))
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jj = len(set(juncs['Symbol']))

(gg, tt, ee, jj))

 $print("\nGene:\t\t\%d\nTranscript:\t\%d\nExon:\t\t\%d\nJunction:\t\%d" \ \%$

Transcript: 15 Exon: 47 Junction: 3

1.3.2 Feature effect size summary

```
[15]: feature_list = ['Genes', 'Transcript', 'Exons', 'Junctions']
      feature_df = [genes, trans, exons, juncs]
      for ii in range(4):
          ff = feature df[ii]
          half = len(set(ff[(np.abs(ff['logFC']) >= 0.5)].index))
          one = len(set(ff[(np.abs(ff['logFC']) >= 1)].index))
          print("\nThere are %d unique %s with abs(log2FC) >= 0.5" % (half,
       →feature_list[ii]))
          print("There are %d unique %s with abs(log2FC) >= 1" % (one, __
       →feature_list[ii]))
     There are 31 unique Genes with abs(log2FC) >= 0.5
     There are 1 unique Genes with abs(log2FC) >= 1
     There are 13 unique Transcript with abs(log2FC) >= 0.5
     There are 3 unique Transcript with abs(log2FC) >= 1
     There are 61 unique Exons with abs(log2FC) >= 0.5
     There are 3 unique Exons with abs(log2FC) >= 1
     There are 3 unique Junctions with abs(log2FC) >= 0.5
     There are 1 unique Junctions with abs(log2FC) >= 1
[16]: feature_list = ['Genes', 'Transcripts', 'Exons', 'Junctions']
      feature_df = [genes, trans, exons, juncs]
      for ii in range(4):
          ff = feature_df[ii]
          half = len(set(ff[(np.abs(ff['logFC']) >= 0.5)].ensemblID))
          one = len(set(ff[(np.abs(ff['logFC']) >= 1)].ensemblID))
          print("\nThere are %d unique %s with abs(log2FC) >= 0.5" % (half,
       →feature_list[ii]))
          print("There are %d unique %s with abs(log2FC) >= 1" % (one, __
       →feature_list[ii]))
     There are 31 unique Genes with abs(log2FC) >= 0.5
     There are 1 unique Genes with abs(log2FC) >= 1
     There are 11 unique Transcripts with abs(log2FC) >= 0.5
     There are 3 unique Transcripts with abs(log2FC) >= 1
```