```
In [1]: def install_reqs():
    #This defines a function named install_reqs.
    !pip install pandas
    #Installs the pandas library, used for reading and manipulating tabular data (li
    !pip install "matplotlib>=3.4"
    #Installs matplotlib version 3.4 or higher, used for plotting graphs and visuali
    !pip install numpy
    #Installs numpy, a fundamental package for numerical computations and arrays.
    !pip install statsmodels
    #Installs statsmodels, a library used for statistical tests (e.g., t-tests, Wilc
    !pip install scipy
    #Installs scipy, which contains scientific computing tools, including statistical
    install_reqs()
    # to install all the above dependencies uncomment the above line.
```

Defaulting to user installation because normal site-packages is not writeable Requirement already satisfied: pandas in c:\programdata\anaconda3\lib\site-packag es (2.3.0)

Requirement already satisfied: numpy>=1.22.4 in c:\users\hp\appdata\roaming\pytho n\python310\site-packages (from pandas) (1.26.4)

Requirement already satisfied: python-dateutil>=2.8.2 in c:\programdata\anaconda3 \lib\site-packages (from pandas) (2.9.0.post0)

Requirement already satisfied: pytz>=2020.1 in c:\programdata\anaconda3\lib\site-packages (from pandas) (2025.2)

Requirement already satisfied: tzdata>=2022.7 in c:\programdata\anaconda3\lib\sit e-packages (from pandas) (2025.2)

Requirement already satisfied: six>=1.5 in c:\programdata\anaconda3\lib\site-pack ages (from python-dateutil>=2.8.2->pandas) (1.17.0)

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Requirement already satisfied: contourpy>=1.0.1 in c:\programdata\anaconda3\lib\s ite-packages (from matplotlib>=3.4) (1.3.1)

Requirement already satisfied: cycler>=0.10 in c:\programdata\anaconda3\lib\site-packages (from matplotlib>=3.4) (0.11.0)

Requirement already satisfied: fonttools>=4.22.0 in c:\programdata\anaconda3\lib\site-packages (from matplotlib>=3.4) (4.55.3)

Requirement already satisfied: kiwisolver>=1.3.1 in c:\programdata\anaconda3\lib \site-packages (from matplotlib>=3.4) (1.4.8)

Requirement already satisfied: numpy>=1.23 in c:\users\hp\appdata\roaming\python \python310\site-packages (from matplotlib>=3.4) (1.26.4)

Requirement already satisfied: packaging>=20.0 in c:\programdata\anaconda3\lib\si te-packages (from matplotlib>=3.4) (25.0)

Requirement already satisfied: pillow>=8 in c:\programdata\anaconda3\lib\site-pac kages (from matplotlib>=3.4) (11.1.0)

Requirement already satisfied: pyparsing>=2.3.1 in c:\programdata\anaconda3\lib\s ite-packages (from matplotlib>=3.4) (3.2.0)

Requirement already satisfied: python-dateutil>=2.7 in c:\programdata\anaconda3\l ib\site-packages (from matplotlib>=3.4) (2.9.0.post0)

Requirement already satisfied: six>=1.5 in c:\programdata\anaconda3\lib\site-pack ages (from python-dateutil>=2.7->matplotlib>=3.4) (1.17.0)

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Defaulting to user installation because normal site-packages is not writeable Requirement already satisfied: numpy in c:\users\hp\appdata\roaming\python\python 310\site-packages (1.26.4)

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Defaulting to user installation because normal site-packages is not writeable
       Requirement already satisfied: statsmodels in c:\programdata\anaconda3\lib\site-p
       ackages (0.14.4)
       Requirement already satisfied: numpy<3,>=1.22.3 in c:\users\hp\appdata\roaming\py
       thon\python310\site-packages (from statsmodels) (1.26.4)
       Requirement already satisfied: scipy!=1.9.2,>=1.8 in c:\programdata\anaconda3\lib
       \site-packages (from statsmodels) (1.15.3)
       Requirement already satisfied: pandas!=2.1.0,>=1.4 in c:\programdata\anaconda3\li
       b\site-packages (from statsmodels) (2.3.0)
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       packages (from statsmodels) (1.0.1)
       Requirement already satisfied: packaging>=21.3 in c:\programdata\anaconda3\lib\si
       te-packages (from statsmodels) (25.0)
       Requirement already satisfied: python-dateutil>=2.8.2 in c:\programdata\anaconda3
       \lib\site-packages (from pandas!=2.1.0,>=1.4->statsmodels) (2.9.0.post0)
       Requirement already satisfied: pytz>=2020.1 in c:\programdata\anaconda3\lib\site-
       packages (from pandas!=2.1.0,>=1.4->statsmodels) (2025.2)
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       e-packages (from pandas!=2.1.0,>=1.4->statsmodels) (2025.2)
       Requirement already satisfied: six>=1.5 in c:\programdata\anaconda3\lib\site-pack
       ages (from python-dateutil>=2.8.2->pandas!=2.1.0,>=1.4->statsmodels) (1.17.0)
       WARNING: Ignoring invalid distribution -atplotlib (c:\programdata\anaconda3\lib\s
       ite-packages)
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       ackages)
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       ite-packages)
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       ackages)
       Defaulting to user installation because normal site-packages is not writeable
       Requirement already satisfied: scipy in c:\programdata\anaconda3\lib\site-package
       Requirement already satisfied: numpy<2.5,>=1.23.5 in c:\users\hp\appdata\roaming
       \python\python310\site-packages (from scipy) (1.26.4)
       WARNING: Ignoring invalid distribution -atplotlib (c:\programdata\anaconda3\lib\s
       ite-packages)
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       ackages)
       WARNING: Ignoring invalid distribution -atplotlib (c:\programdata\anaconda3\lib\s
       ite-packages)
       WARNING: Ignoring invalid distribution -orch (c:\programdata\anaconda3\lib\site-p
       ackages)
In [2]: import pandas as pd
        import numpy as np
        import matplotlib.pyplot as plt
In [3]: from compass analysis import cohens d, wilcoxon test, get reaction consistencies
In [4]: from matplotlib import __version__ as matplotlibversion
        if matplotlibversion < "3.4":</pre>
            print("Matplotlib versions older than 3.4 may not be able to generate figure
       Matplotlib versions older than 3.4 may not be able to generate figure 2E, as they
       do not support alpha arrays
In [5]: import os
        os.system("compass --data expression.tsv \
```

--model RECON2 mat --species mus musculus --media default-media --lamb

file:///C:/Users/HP/Downloads/Treg LN vs exTreg LN.html

```
--and-function mean --output-dir extdata/Th17 --penalty-diffusion knn
                   --isoform-summing legacy --num-processes 50")
Out[5]: 0
In [6]: import os
         print(os.getcwd())
        C:\Users\HP\Tcon vs Treg\extdata
In [7]: reaction_penalties = pd.read_csv("reactions.tsv", sep="\t", index_col=0)
In [8]: import pandas as pd
         # Load reactions.tsv
         reaction_penalties = pd.read_csv("reactions.tsv", sep="\t", index_col=0)
         # Extract cell IDs from the columns
         cell_ids = reaction_penalties.columns
         # Infer cell type from prefix
         def infer_cell_type(cell_id):
             if cell_id.startswith("Treg"):
                 return "Treg"
             elif cell_id.startswith("Tcon"):
                 return "Tcon"
             elif cell_id.startswith("exTreg"):
                 return "exTreg"
             else:
                 return "Unknown"
         cell_types = [infer_cell_type(cell) for cell in cell_ids]
         # Create metadata DataFrame
         cell metadata = pd.DataFrame({
             "cell_id": cell_ids,
             "cell_type": cell_types
         }).set_index("cell_id")
         # Save to CSV
         cell metadata.to csv("cell metadata.csv")
         print(" ✓ Created cell metadata.csv with the following cell types:")
         print(cell_metadata["cell_type"].value_counts())
        Created cell_metadata.csv with the following cell types:
        cell_type
        exTreg
                  20
                  20
        Tcon
        Treg
                  16
        Name: count, dtype: int64
In [9]: cell_metadata = pd.read_csv("cell_metadata.csv", sep=",")
         cell_metadata.set_index("cell_id", inplace=True)
In [10]: print(cell_metadata.columns.tolist())
         print(cell_metadata.head())
```

```
['cell_type']
                                 cell_type
        cell id
        exTreg_Lymph_exTreg2_L1
                                    exTreg
        Tcon_Lymph_Tcon9_L2
                                      Tcon
        exTreg Spleen exTreg8 L1
                                    exTreg
        Tcon_Spleen_Tcon5_L2
                                      Tcon
        Treg_Spleen_Treg5_L1
                                      Treg
In [11]: Treg_cells = cell_metadata.index[cell_metadata["cell_type"] == "Treg"]
         Tcon_cells = cell_metadata.index[cell_metadata["cell_type"] == "Tcon"]
         exTreg cells = cell metadata.index[cell metadata["cell type"] == "exTreg"]
In [12]: print(cell_metadata.columns.tolist())
        ['cell_type']
In [13]: print(cell_metadata.head())
         print(cell_metadata.columns.tolist())
                                 cell_type
        cell id
        exTreg_Lymph_exTreg2_L1
                                    exTreg
        Tcon_Lymph_Tcon9_L2
                                      Tcon
        exTreg_Spleen_exTreg8_L1
                                    exTreg
        Tcon_Spleen_Tcon5_L2
                                     Tcon
        Treg Spleen Treg5 L1
                                      Treg
        ['cell_type']
In [14]: cell_metadata = pd.read_csv("cell_metadata.csv", index_col=0)
         treg_ln_cells = cell_metadata[
             (cell_metadata['cell_type'] == 'Treg') &
             (cell_metadata.index.str.contains("Lymph"))
         ].index
         extreg_ln_cells = cell_metadata[
             (cell_metadata['cell_type'] == 'exTreg') &
             (cell_metadata.index.str.contains("Lymph"))
         ].index
In [15]: cell_metadata = pd.read_csv("cell_metadata.csv", index_col=0)
         treg_ln_cells = cell_metadata[
             (cell_metadata['cell_type'] == 'Treg') &
             (cell_metadata.index.str.contains("Lymph"))
         ].index
         extreg_ln_cells = cell_metadata[
             (cell_metadata['cell_type'] == 'exTreg') &
             (cell_metadata.index.str.contains("Lymph"))
         ].index
In [16]:
        reaction metadata = pd.read csv("reaction metadata.csv", index col = 0)
In [17]: reaction_metadata.loc[['r0281']]
```

1.00 *

Out[17]:

reaction_name formula associated_genes subsystem EC_num

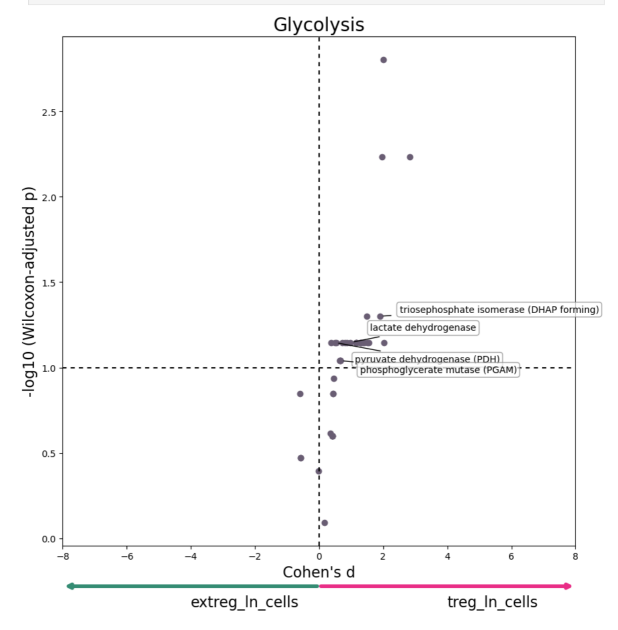
reaction_no_direction

```
Water
                                                                         Methionine
                              Putrescine:oxygen
                                                  [e] +
                                                                               and
                       r0281
                                 oxidoreductase
                                                 1.00 *
                                                                  AOC1
                                                                                         1.4
                                                                            cysteine
                                 (deaminating)... O2 [e] +
                                                                         metabolism
                                                  1.00 *
                                                  Putr...
In [18]: #This function is repeated here for clarity
         def get reaction consistencies(compass reaction penalties, min range=1e-3):
                  Converts the raw penalties outputs of compass into scores per reactions
             df = -np.log(compass_reaction_penalties + 1)
             df = df[df.max(axis=1) - df.min(axis=1) >= min_range]
             df = df - df.min().min()
             return df
        reaction_consistencies = get_reaction_consistencies(reaction_penalties)
In [19]:
In [20]: common_cells = list(set(reaction_consistencies.columns).intersection(set(cell_me
         reaction_consistencies = reaction_consistencies[common_cells]
In [21]: print(reaction_consistencies.columns.tolist()[:10])
        ['exTreg_Spleen_exTreg8_L1', 'exTreg_Lymph_exTreg3_L2', 'exTreg_Lymph_exTreg2_L
        2', 'exTreg_Spleen_exTreg9_L1', 'exTreg_Spleen_exTreg6_L2', 'Tcon_Spleen_Tcon1_L
        2', 'Tcon_Spleen_Tcon5_L1', 'Tcon_Spleen_Tcon3_L2', 'Tcon_Lymph_Tcon9_L2', 'exTre
        g_Spleen_exTreg9_L2']
In [22]: print(exTreg cells[:10])
        Index(['exTreg_Lymph_exTreg2_L1', 'exTreg_Spleen_exTreg8_L1',
               'exTreg_Spleen_exTreg7_L2', 'exTreg_Lymph_exTreg3_L2',
               'exTreg_Spleen_exTreg8_L2', 'exTreg_Lymph_exTreg2_L2',
               'exTreg_Lymph_exTreg3_L1', 'exTreg_Spleen_exTreg7_L1',
               'exTreg_Spleen_exTreg6_L2', 'exTreg_Lymph_exTreg4_L2'],
              dtype='object', name='cell_id')
In [23]: wilcox results = wilcoxon test(reaction consistencies, treg ln cells, extreg ln
         wilcox_results['metadata_r_id'] = ""
         for r in wilcox_results.index:
             if r in reaction metadata.index:
                  wilcox_results.loc[r, 'metadata_r_id'] = r
             elif r[:-4] in reaction metadata.index:
                  wilcox_results.loc[r, 'metadata_r_id'] = r[:-4]
             else:
                  print(f"Should not occur \rightarrow {r}")
In [24]: W = wilcox results.merge(reaction metadata, how='left',
                                   left_on='metadata_r_id', right_index=True, validate='m:
         W = W[W['confidence'].isin([0,4])]
```

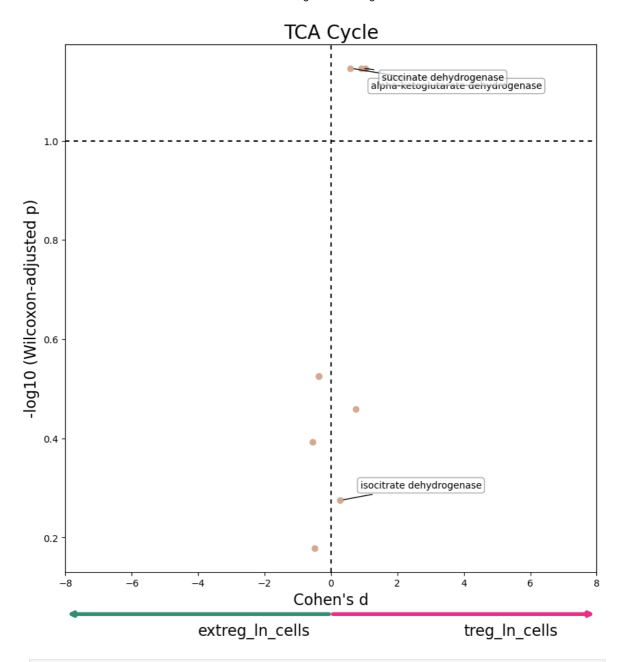
```
W = W[~W['EC_number'].isna()]
          W.loc[(W['formula'].map(lambda x: '[m]' not in x)) & (W['subsystem'] == "Citric
         wilcox results.loc[['r0281 pos']]
In [25]:
Out[25]:
                     wilcox_stat wilcox_pval cohens_d adjusted_pval metadata_r_id
          r0281 pos
                           45.0
                                   0.117882
                                             0.903637
                                                           0.141679
                                                                            r0281
In [26]:
          reaction_metadata.loc['r0281']['formula']
Out[26]: '1.00 * Water [e] + 1.00 * O2 [e] + 1.00 * Putrescine [e] --> 1.00 * Ammonium
          [e] + 1.00 * Hydrogen peroxide [e] + 1.00 * 4-Aminobutanal [e]\nAOC1'
In [106...
          def plot differential scores(data, title, c):
              plt.figure(figsize=(10,10))
              axs = plt.gca()
              axs.scatter(data['cohens_d'], -np.log10(data['adjusted_pval']), c=c)
              axs.set_xlabel("Cohen's d", fontsize=16)
              axs.set_ylabel("-log10 (Wilcoxon-adjusted p)", fontsize=16)
              # Visual markers and title
              axs.set_xlim(-8, 8)
              axs.axvline(0, dashes=(3,3), c='black')
              axs.axhline(1, dashes=(3,3), c='black')
              axs.set_title(title, fontdict={'fontsize':20})
              # Arrows for sample identity
              axs.annotate('', xy=(0.5, -0.08), xycoords='axes fraction', xytext=(0, -0.08)
                           arrowprops=dict(arrowstyle="<-", color='#348C73', linewidth=4))</pre>
              axs.annotate('treg_ln_cells', xy=(0.75, -0.12), xycoords='axes fraction', fo
              axs.annotate('', xy=(0.5, -0.08), xycoords='axes fraction', xytext=(1, -0.08
                            arrowprops=dict(arrowstyle="<-", color='#E92E87', linewidth=4))</pre>
              axs.annotate('extreg_ln_cells', xy=(0.25, -0.12), xycoords='axes fraction',
              # Dynamic annotation loop with variable offsets and labels
              for i, r in enumerate(data.index):
                  if r in labeled_reactions:
                      x = data.loc[r, 'cohens d']
                      y = -np.log10(data.loc[r, 'adjusted_pval'])
                      dx = 30 if x >= 0 else -120 # horizontal offset
                      dy = (i \% 6) * 12 - 30
                                                  # vertical offset varies to reduce stac
                      axs.annotate(
                           labeled reactions[r],
                           (x, y),
                          xytext=(dx, dy),
                           textcoords='offset pixels',
                          arrowprops=dict(arrowstyle="-", shrinkA=6, shrinkB=4),
                          fontsize=10,
                           zorder=5, # make sure text is on top
                          bbox=dict(boxstyle="round,pad=0.3", fc="white", ec="gray", alpha
                       )
          filtered data = pd.concat([W[W['subsystem'] == "Glycolysis/gluconeogenesis"],
In [108...
                       W[W['subsystem'] == "Citric acid cycle"],
```

```
W[W['subsystem'].isin(amino_acid_metab)],
W[W['subsystem'] == "Fatty acid oxidation"]])
```

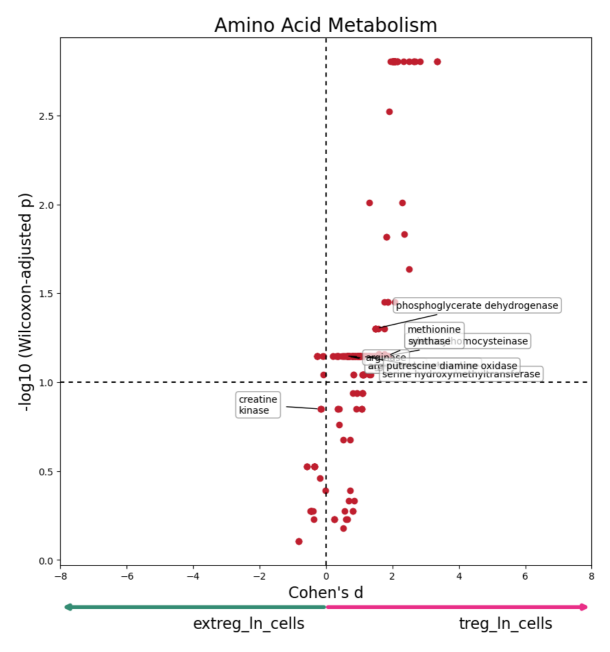
```
In [110... data = W[W['subsystem'] == "Glycolysis/gluconeogenesis"]
    plot_differential_scores(data, title='Glycolysis', c="#695D73")
```



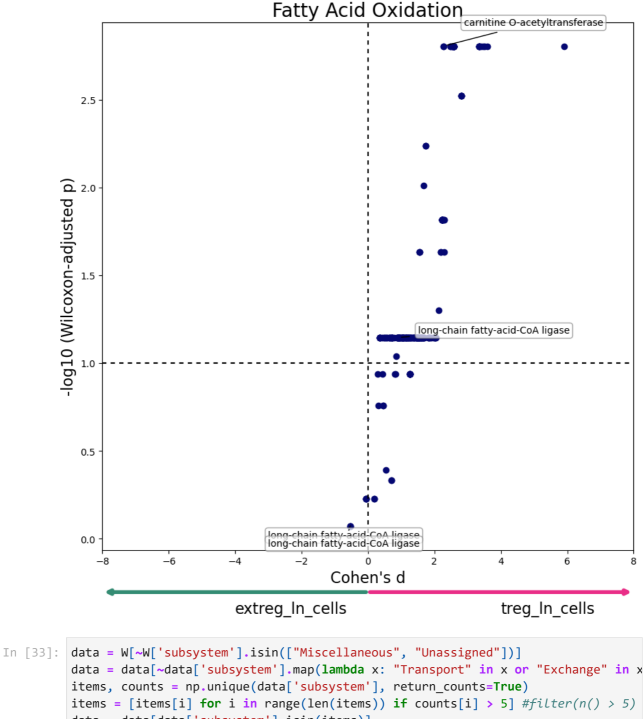
```
In [112... data = W[W['subsystem'] == "Citric acid cycle"]
    plot_differential_scores(data, title="TCA Cycle", c="#D3A991")
```



```
In [114... data = W[W['subsystem'].isin(amino_acid_metab)].copy()
  data['adjusted_pval'] = data['adjusted_pval'].clip(1e-12)
  plot_differential_scores(data, "Amino Acid Metabolism", c="#BF1E2E")
```



In [116... data = W[W['subsystem'] == "Fatty acid oxidation"]
 plot_differential_scores(data, "Fatty Acid Oxidation", c="#040772")



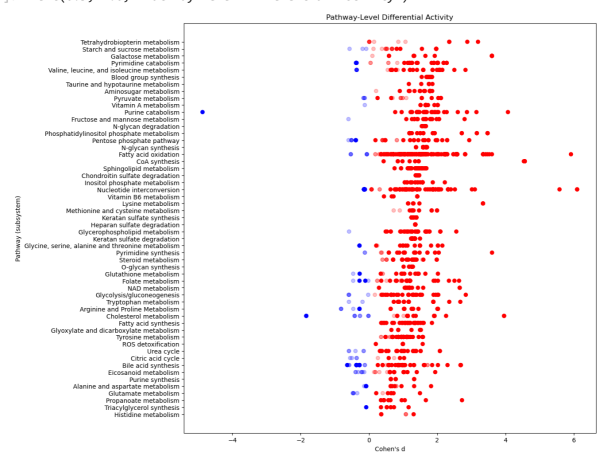
```
items, counts = np.unique(data[ subsystem ], return_counts=irue)
  items = [items[i] for i in range(len(items)) if counts[i] > 5] #filter(n() > 5)
  data = data[data['subsystem'].isin(items)]

In [34]: import matplotlib.pyplot as plt
  plt.figure(figsize=(12, 12))
  axs = plt.gca()

d = data[data['adjusted_pval'] < 0.1].groupby('subsystem')['cohens_d'].median().decounted = d.sort_values()
  axs.scatter(d_sorted, d_sorted.index, alpha=0)
  color = data['cohens_d'].map(lambda x: 'r' if x >= 0 else 'b')
  alpha = data['adjusted_pval'].map(lambda x: 1.0 if x < 0.1 else 0.25)
  axs.scatter(data['cohens_d'], data['subsystem'], c=color, alpha=alpha)</pre>
```

```
axs.set_xlabel("Cohen's d")
axs.set_ylabel("Pathway (subsystem)")
axs.set_title("Pathway-Level Differential Activity")
```

Out[34]: Text(0.5, 1.0, 'Pathway-Level Differential Activity')



```
reaction_penalties = pd.read_csv("reactions.tsv", sep="\t", index_col = 0)
In [35]:
          reaction_penalties[reaction_penalties <= 1e-4] = 0</pre>
          reaction_penalties = reaction_penalties[np.all(reaction_penalties != 0, axis=1)]
          reaction_penalties = reaction_penalties[reaction_penalties.max(axis=1) - reaction_penalties.max(axis=1) - reaction_penalties.
In [36]:
          meta rxns map = get metareactions(reaction penalties)
In [37]:
          meta rxns = reaction penalties.join(pd.DataFrame(meta rxns map, columns=["meta r
         meta rxn consistencies = get reaction consistencies(meta rxns)
          treg_cells = cell_metadata.index[cell_metadata["cell_type"] == "treg_ln_cells"]
In [39]:
          tcon_cells = cell_metadata.index[cell_metadata["cell_type"] == "extreg_ln_cells"
          wilcox meta rxn results = wilcoxon test(meta rxn consistencies, treg ln cells, e
In [40]:
          wilcox_meta_rxn_results.iloc[0:1]
Out[40]:
                       wilcox_stat wilcox_pval cohens_d adjusted_pval
          meta_rxn_id
```

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1

48.0

0.055944

1.135176

0.078347

```
In [41]: wilcox meta rxn expanded = pd.DataFrame(index=reaction penalties.index, columns=
         for i in range(len(wilcox_meta_rxn_expanded.index)):
             if (meta_rxns_map[i] in wilcox_meta_rxn_results.index):
                  wilcox_meta_rxn_expanded.loc[wilcox_meta_rxn_expanded.index[i]] = wilcox
         wilcox_meta_rxn_expanded = wilcox_meta_rxn_expanded.dropna().astype('float64')
         wilcox_meta_rxn_expanded['metadata_r_id'] = ""
In [42]:
         for r in wilcox_meta_rxn_expanded.index:
             if r in reaction_metadata.index:
                 wilcox_meta_rxn_expanded.loc[r, 'metadata_r_id'] = r
             elif r[:-4] in reaction metadata.index:
                 wilcox_meta_rxn_expanded.loc[r, 'metadata_r_id'] = r[:-4]
             else:
                  print("Should not occur")
In [43]: wilcox meta rxn expanded.iloc[0:1]
Out[43]:
                            wilcox_stat wilcox_pval cohens_d adjusted_pval metadata_r_id
          10FTHF5GLUtl_pos
                                  48.0
                                          0.055944
                                                    1.198422
                                                                 0.078347
                                                                           10FTHF5GLUtl
In [44]: outputs = {
             "wilcox_results.csv": wilcox_results,
             "reaction_consistencies.csv": reaction_consistencies,
             "reaction_metadata.csv": reaction_metadata,
             "wilcox_meta_rxn_results.csv": wilcox_meta_rxn_results,
             "wilcox_meta_rxn_expanded.csv": wilcox_meta_rxn_expanded,
             "final_stats_with_metadata.csv": W,
         }
         for name, df in outputs.items():
             df.to_csv(name)
In [45]: import zipfile
         with zipfile.ZipFile("Treg LN vs exTreg LN.zip", "w") as zipf:
             for filename in outputs:
                 zipf.write(filename)
In [46]: from IPython.display import FileLink
         FileLink("Treg LN vs exTreg LN.zip")
Out[46]: Treg LN vs exTreg LN.zip
 In [ ]:
```