PPMI Merge

2022-03-30

Here, we utilize PPMI subject clinical data files (PPMI Study Data) to create a single .csv file that contains longitudinal clinical information of PPMI subjects (included in Analytic Cohort).

Getting the data

- 1.) Get access to PPMI database and login.
- 2.) Click download from the navigation bar and select study data.
- 3.) Select ALL documents and zip files and click download.
- 4.) Unzip and save the folder on your computer.

Import required modules

```
import pandas as pd
import numpy as np
import boto3
from datetime import datetime
from dateutil import relativedelta
## PPMI Study Data path
userdir = '/Users/areardon/Desktop/ppmi_merge/'
ppmi_download_path = userdir + 'PPMI_Study_Data_Download/'
invicro_data_path = userdir
genetics_path = userdir + 'genetic_data/'
#### CLINICAL INFO ####
xlsx = pd.ExcelFile(ppmi_download_path +
→ 'Consensus Committee Analytic Datasets 280CT21.xlsx') # Read in main xlsx file
def create_cohort_df(xlsx, sheet) :
    cohort_df = pd.read_excel(xlsx, sheet)
    cols_delete = ['Unnamed', 'CONDATE'] # Columns to remove from sheet/ df
    cohort df = cohort df.loc[:, ~cohort df.columns.str.startswith(tuple(cols delete))] #
  Remove columns in cols_delete
    # Create new columns for Enrollment.Subtype and Consensus.Subtype in pd_df
    cohort_df["Enrollment.Subtype"] = '' # Create new column called 'Enrollment.Subtype'
    cohort_df["Consensus.Subtype"] = '' # Create new column called 'Consensus.Subtype' in
\rightarrow pd_df
    if sheet == 'HC' :
        cohort_df['Subgroup'] = 'Healthy Control'
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if sheet == 'SWEDD' :
       cohort_df['Comments'] = ''
   return cohort_df
def merge_columns(df : pd.DataFrame , old_df_columns : list, new_df_column_name : str,

    separator = str) :

   Takes entries in each of old_df_columns and joins them together with a sepator of
empty/nan column entries.
   df = df.replace(r'^\s*\s', np.NaN, regex=True) # Fill in empty cells with nan
   df[new_df_column_name] = df[old_df_columns].agg(lambda x: x.dropna().str.cat(sep=
df.drop(old_df_columns, axis = 1, inplace = True)
   return df
def getNamesFromDataframe(df, str) :
   col_names = [col for col in df.columns if str in col]
   return col_names
def merge_new_csv(df, csv_filename, list_cols) :
   demo df = pd.read csv(ppmi download path + csv filename)
   demo df = demo df[list cols]
   ppmi_merge = pd.merge(df, demo_df, on = ['PATNO', 'EVENT_ID'], how = "outer") # Merge
   return ppmi_merge
def isNaN(num):
   return num != num
def decode_0_1_no_yes(df, list) :
   for i in list :
       df[i] = df[i].astype(int, errors = "ignore")
       df[i].replace({0 : 'No' , 1 : 'Yes', 2 : 'Uncertain'}, inplace = True)
   return df
def decode_none2severe(df, list):
   for i in list :
       df[i] = df[i].astype(int, errors = "ignore")
       df[i].replace({0 : 'None', 1 : 'Slight', 2: 'Mild', 3 : 'Moderate', 4 :
return df
# Create cohort df
pd_df = create_cohort_df(xlsx, 'PD') # Create Parkinson's Disease data frame from 'PD'
→ sheet in 'Consensus_Committee_Analytic_Datasets_280CT21.xlsx'
prodromal_df = create_cohort_df(xlsx, 'Prodromal')
hc_df = create_cohort_df(xlsx, 'HC')
swedd_df = create_cohort_df(xlsx, 'SWEDD') # SWEDD subjects
# Decode Enrollment.Subtype ('Summary' sheet) and Consensus.Subtype ('Summary Analytic'
→ sheet) of 'Consensus_Committee_Analytic_Datasets_280CT21.xlsx'
```

```
pd_df.loc[(pd_df['ENRLPD'] == 1) & (pd_df['ENRLLRRK2'] == 0) & (pd_df['ENRLGBA'] == 0) &
→ (pd_df['ENRLSNCA'] == 0), 'Enrollment.Subtype'] = '' # Sporadic - don't need to
→ define here bc already covered in 'Subgroup' column in PD sheet
pd_df.loc[(pd_df['ENRLPD'] == 1) & (pd_df['ENRLLRRK2'] == 1), 'Enrollment.Subtype'] = '

    ∴ LRRK2¹

pd_df.loc[(pd_df['ENRLPD'] == 1) & (pd_df['ENRLGBA'] == 1), 'Enrollment.Subtype'] = ' :
pd_df.loc[(pd_df['ENRLPD'] == 1) & (pd_df['ENRLSNCA'] == 1), 'Enrollment.Subtype'] = ':
→ SNCA'
pd_df.loc[(pd_df['CONLRRK2'] == 1) & (pd_df['CONLRRK2'] == 0) | (pd_df['CONLRRK2'] == '.') &
pd_df.loc[(pd_df['CONPD'] == 1) & (pd_df['CONLRRK2'] == 1) & (pd_df['CONGBA'] == 0) &
pd_df.loc[(pd_df['CONPD'] == 1) & (pd_df['CONLRRK2'] == 0) & (pd_df['CONGBA'] == 1) &
\rightarrow (pd_df['CONSNCA'] == 0) , 'Consensus.Subtype'] = 'Genetic : GBA'
pd_df.loc[(pd_df['CONPD'] == 1) & (pd_df['CONLRRK2'] == 1) & (pd_df['CONGBA'] == 1) &
pd_df.loc[(pd_df['CONPD'] == 1) & (pd_df['CONLRRK2'] == 0) & (pd_df['CONGBA'] == 0) &
pd_df.loc[(pd_df['CONPD'] == 0) & (pd_df['CONPROD'] == 0) , 'Consensus.Subtype'] =
pd_df.loc[(pd_df['CONPD'] == 0) & (pd_df['CONPROD'] == 1) & (pd_df['CONLRRK2'] == 1) &
pd_df.loc[(pd_df['CONPD'] == 0) & (pd_df['CONPROD'] == 1) & (pd_df['CONLRRK2'] == 0) &

→ GBA Prodromal'

# Decode Enrollment.Subtype and Consensus.Subtype to be categorical variables for
\rightarrow prodromal df
prodromal_df.loc[(prodromal_df['ENRLPROD'] == 1) & (prodromal_df['ENRLLRRK2'] == 1),
prodromal_df.loc[(prodromal_df['ENRLPROD'] == 1) & (prodromal_df['ENRLGBA'] == 1),
→ 'Enrollment.Subtype'] = ' : GBA Prodromal'
prodromal_df.loc[(prodromal_df['ENRLPROD'] == 1) & (prodromal_df['ENRLSNCA'] == 1),
→ 'Enrollment.Subtype'] = ' : SNCA Prodromal'
prodromal_df.loc[(prodromal_df['ENRLPROD'] == 1) & (prodromal_df['ENRLHPSM'] == 1)
→ ,'Enrollment.Subtype'] = '' # Hyposmia already covered in 'Subgroup' column
prodromal_df.loc[(prodromal_df['ENRLPROD'] == 1) & (prodromal_df['ENRLRBD'] == 1),
→ 'Enrollment.Subtype'] = '' # RBD already covered in 'Subgroup' column
prodromal_df.loc[(prodromal_df['CONPROD'] == 1) & (prodromal_df['PHENOCNV'] == 0) &
   (prodromal_df['CONLRRK2'] == 1) & (prodromal_df['CONGBA'] == 0) &
   (prodromal_df['CONSNCA'] == 0) , 'Consensus.Subtype'] = 'Genetic : LRRK2 Prodromal'
prodromal_df.loc[(prodromal_df['CONPROD'] == 1) & (prodromal_df['PHENOCNV'] == 1) &
prodromal_df.loc[(prodromal_df['CONPROD'] == 0) & (prodromal_df['CONLRRK2'] == 1) &
→ = 'Genetic : LRRK2 not Prodromal'
prodromal_df.loc[(prodromal_df['CONPROD'] == 1) & (prodromal_df['PHENOCNV'] == 0) &
→ (prodromal_df['CONSNCA'] == 0), 'Consensus.Subtype'] = 'Genetic : GBA Prodromal'
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prodromal_df.loc[(prodromal_df['CONPROD'] == 1) & (prodromal_df['PHENOCNV'] == 1) &
(prodromal_df['CONSNCA'] == 0) & (prodromal_df['CONRBD'] == 0), 'Consensus.Subtype']
  = 'Genetic : GBA Phenoconverted'
prodromal_df.loc[(prodromal_df['CONPROD'] == 0) & (prodromal_df['CONLRRK2'] == 0) &
→ ] = 'Genetic : GBA not Prodromal'
prodromal df.loc[(prodromal df['CONPROD'] == 1) & (prodromal df['PHENOCNV'] == 0) &
→ (prodromal_df['CONSNCA'] == 0) , 'Consensus.Subtype'] = 'Genetic : LRRK2 + GBA
prodromal df.loc[(prodromal df['CONPROD'] == 1) & (prodromal df['PHENOCNV'] == 1) &
  (prodromal_df['CONLRRK2'] == 1) & (prodromal_df['CONGBA'] == 1) &
 (prodromal_df['CONSNCA'] == 0) , 'Consensus.Subtype'] = 'Genetic : LRRK2 + GBA
→ Phenoconverted'
prodromal_df.loc[(prodromal_df['CONPROD'] == 0) & (prodromal_df['CONLRRK2'] == 1) &
→ ] = 'Genetic : LRRK2 + GBA not Prodromal'
prodromal_df.loc[(prodromal_df['CONPROD'] == 1) & (prodromal_df['PHENOCNV'] == 0) &
  (prodromal_df['CONLRRK2'] == 0) & (prodromal_df['CONGBA'] == 0) &
  (prodromal_df['CONSNCA'] == 1), 'Consensus.Subtype'] = 'Genetic : SNCA Prodromal'
prodromal_df.loc[(prodromal_df['CONPROD'] == 1) & (prodromal_df['PHENOCNV'] == 1) &
→ (prodromal_df['CONSNCA'] == 1), 'Consensus.Subtype'] = 'Genetic : SNCA
→ Phenoconverted'
prodromal_df.loc[(prodromal_df['CONPROD'] == 0) & (prodromal_df['CONLRRK2'] == 0) &
→ ] = 'Genetic : SNCA not Prodromal'
prodromal df.loc[(prodromal df['CONPROD'] == 0) & (prodromal df['CONLRRK2'] == 0) &
→ ] = 'No Mutation not Prodromal'
prodromal_df.loc[(prodromal_df['CONPROD'] == 1) & (prodromal_df['PHENOCNV'] == 0) &
  (prodromal_df['CONLRRK2'] == 0) & (prodromal_df['CONGBA'] == 0) &
  (prodromal_df['CONSNCA'] == 0) & (prodromal_df['CONHPSM'] == 1) &
 (prodromal_df['CONRBD'] == '.'), 'Consensus.Subtype'] = 'Hyposmia'
prodromal_df.loc[(prodromal_df['CONPROD'] == 1) & (prodromal_df['PHENOCNV'] == 1) &
→ (prodromal_df['CONRBD'] == '.'), 'Consensus.Subtype'] = 'Hyposmia: Phenoconverted'
prodromal df.loc[(prodromal df['CONPROD'] == 0) & (prodromal df['CONLRRK2'] == 0) &
→ ] = 'Hyposmia : not Prodromal'
prodromal_df.loc[(prodromal_df['CONPROD'] == 1) & (prodromal_df['CONRBD'] == 1) &
prodromal_df.loc[(prodromal_df['CONPROD'] == 1) & (prodromal_df['CONRBD'] == 1) &
→ = 'RBD : Phenoconverted'
prodromal_df.loc[(prodromal_df['CONPROD'] == 1) & (prodromal_df['CONRBD'] == 1) &
→ ] = 'RBD : Phenoconverted with GBA'
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```
# Decode Enrollment. Subtype and Consensus. Subtype to be categorical variables for Healthy
\hookrightarrow Control df
hc_df.loc[(hc_df['ENRLHC'] == 1), 'Enrollment.Subtype'] = ''
hc_df.loc[(hc_df['CONHC'] == 1) & (hc_df['CONLRRK2'] == 0) & (hc_df['CONGBA'] == 0) &
hc_df.loc[(hc_df['CONHC'] == 1) & (hc_df['CONLRRK2'] == 1) & (hc_df['CONGBA'] == 0) &
hc_df.loc[(hc_df['CONHC'] == 1) & (hc_df['CONLRRK2'] == 0) & (hc_df['CONGBA'] == 1) &
hc_df.loc[(hc_df['CONHC'] == 1) & (hc_df['CONLRRK2'] == 1) & (hc_df['CONGBA'] == 1) &
hc df.loc[(hc df['CONHC'] == 1) & (hc df['CONLRRK2'] == 0) & (hc df['CONGBA'] == 0) &
hc_df.loc[(hc_df['CONHC'] == 0), 'Consensus.Subtype'] = 'non-HC' # FIXME
# Decode Enrollment.Subtype and Consensus.Subtype to be categorical variables for swedd
\hookrightarrow df
swedd_df.loc[swedd_df['ENRLSWEDD'] == 1 , 'Enrollment.Subtype'] = 'SWEDD Legacy'
swedd_df.loc[swedd_df['CONSWEDD'] == 0 , 'Consensus.Subtype'] = 'SWEDD/PD Active'
swedd_df.loc[swedd_df['CONSWEDD'] == 1 ,'Consensus.Subtype'] = 'SWEDD/non-PD Active'
# Merge the four dataframes (HC, Prodromal, PD, SWEDD)
full_df = pd_df.append([prodromal_df, hc_df, swedd_df]) # Concat all 4 cohort dfs
# Decode and re-organize full df
full_df['CONPD'].replace({1 : 'Parkinson\'s Disease', 0 : ''}, inplace = True)
full_df['CONPROD'].replace({1 : 'Prodromal', 0 : ''}, inplace = True)
full_df['CONHC'].replace({1 : 'Healthy Control', 0 : ''}, inplace = True)
full df['CONSWEDD'].replace({1 : 'SWEDD', 0 : 'SWEDD/PD'}, inplace = True)
full_df['Comments'].replace({'MSA' : 'Multiple System Atrophy'}, inplace = True)
full_df = decode_0_1_no_yes(full_df, ['PHENOCNV'])
full_df = merge_columns(full_df, ['CONPD', 'CONPROD', 'CONHC', 'CONSWEDD'],
→ 'Consensus.Diagnosis_temp', ', ') # Get one column for Consensus Diagnosis called
→ Consensus.Diagnosis_temp
full_df = merge_columns(full_df, ['Consensus.Diagnosis_temp', 'Comments'],
→ 'Consensus.Diagnosis', ': ') # Include the 'Comments' column in with
→ Consensus.Diagnosis temp into a new column called 'Consensus.Diagnosis'
full_df = merge_columns(full_df, ['Subgroup', 'Enrollment.Subtype'], 'Enroll.Subtype',
→ '') # Get one column for Enroll.Subtype
full_df = full_df.loc[:, ~full_df.columns.str.startswith(tuple(['CON', 'ENRL']))] # Remove
→ columns that begin with CON and ENRL
full_df.rename(columns = {'Cohort' : 'Enroll.Diagnosis' , 'PHENOCNV' :
→ 'Subject.Phenoconverted'}, inplace = True)
full_df = full_df[['PATNO', 'Enroll.Diagnosis', 'Enroll.Subtype', 'Consensus.Diagnosis',
→ 'Consensus.Subtype', 'Subject.Phenoconverted', 'DIAG1', 'DIAG1VIS', 'DIAG2', 'DIAG2VIS']]
→ # Reorganize column order
analytic_cohort_subids = full_df['PATNO'].unique() # subids for analytic cohort
## Add in age info at each visit
age_df = pd.read_csv(ppmi_download_path + 'Age_at_visit.csv', skipinitialspace = True) #
\rightarrow Age info
age_df.rename(columns = {'AGE_AT_VISIT' : 'Age'}, inplace = True) # Rename column
ppmi_merge = pd.merge(full_df, age_df, on = 'PATNO', how = "outer") # Merge full_df with
\hookrightarrow age_df
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ppmi_merge['EVENT_ID'].fillna('LOG', inplace = True) # Replace nan cells with 'LOG' in
→ EVENT_ID column -- FIXME (not sure if I should fill this with LOG)-subject 41358
## Add deographics, vital signs, and pd diagnosis history info
ppmi_merge = merge_new_csv(ppmi_merge, 'Demographics.csv', ['PATNO', 'EVENT_ID', 'SEX',
→ 'HANDED', 'BIRTHDT']) # Bday, sex, handedness
ppmi_merge = merge_new_csv(ppmi_merge, 'Vital_Signs.csv', ['PATNO', 'EVENT_ID', 'INFODT',
\hookrightarrow 'WGTKG', 'HTCM']) # Visit date, weight and height
ppmi_merge = merge_new_csv(ppmi_merge, 'PD_Diagnosis_History.csv', ['PATNO', 'EVENT_ID',
→ 'SXDT', 'PDDXDT']) # First symptom date, PD diagnosis date
ppmi_merge['SEX'].replace({0 : 'Female', 1 : 'Male' }, inplace = True) # Decode sex
ppmi_merge['HANDED'].replace({1 : 'Right', 2 : 'Left', 3 : 'Mixed' }, inplace = True) #
→ Decode handedness
ppmi_merge.rename(columns = {'SEX' : 'Sex', 'HANDED' : 'Handed', 'BIRTHDT' : 'BirthDate',
→ 'WGTKG' : 'Weight(kg)', 'HTCM' : 'Height(cm)', 'SXDT' : 'First.Symptom.Date',
→ 'PDDXDT': 'PD.Diagnosis.Date'}, inplace = True) # Rename columns
## Add a PD.Disease.Duration variable (in years)
ppmi_merge['PD.Diagnosis.Duration'] = '' # Initialize PD.Diagnosis.Duration variable
for row_num in range(len(ppmi_merge['PD.Diagnosis.Date'])) :
    if isinstance(ppmi_merge['PD.Diagnosis.Date'].loc[row_num], str) and
    → isinstance(ppmi_merge['INFODT'].loc[row_num], str): # If we have both a PD
    → Diagnosis date and an event id date
       diag_year = int(ppmi_merge['PD.Diagnosis.Date'].loc[row_num].split('/')[1]) #
  Diagnosis year
       diag_month = int(ppmi_merge['PD.Diagnosis.Date'].loc[row_num].split('/')[0]) #
  Diagnosis month
       event_year = int(ppmi_merge['INFODT'].loc[row_num].split('/')[1]) # Visit date
       event_month = int(ppmi_merge['INFODT'].loc[row_num].split('/')[0]) # Visit date
   month
       diff = relativedelta.relativedelta(datetime(event_year, event_month, 1),
→ datetime(diag_year, diag_month, 1)) # FIXME ASSUMPTION visit date was the first of
   the month, diagnosis date was the first of the month
       ppmi_merge['PD.Diagnosis.Duration'].iloc[row_num] = ((diff.years)*12 +
   diff.months)/12 # PD.Diagnosis.Duration in years
# Add Final (FNL) Event ID and info
ppmi_merge = merge_new_csv(ppmi_merge, 'Conclusion_of_Study_Participation.csv', ['PATNO',
→ 'EVENT_ID', 'COMPLT', 'WDRSN', 'WDDT']) # completed study, whithdrawal reason,
\hookrightarrow withdrawal date
ppmi_merge = decode_0_1_no_yes(ppmi_merge, ['COMPLT'])
ppmi_merge['WDRSN'].replace({1 : 'Adverse Event', 2 : 'Completed study per protocol',3 :
→ 'Death', 4: 'Family, care-partner, or social issues', 5: 'Lost to follow up', 6:
_{\rightarrow} 'Non-compliance with study procedures' , 7 : 'Transportation/Travel issues' , 8 :
\hookrightarrow 'Institutionalized' , 9 : 'Subject transitioning to a new cohort' , 10 : 'Subject
→ withdrew consent', 11 : 'Investigator decision', 12 : 'Sponsor decision', 13 :
ppmi_merge.rename(columns = {'COMPLT' : 'Completed.Study' , 'WDRSN':
→ 'Reason.for.Withdrawal','WDDT' : 'Withdrawal.Date'}, inplace = True)
## Add diagnosis change info on event id
def condensed_df(df, keep_col_list, rename_col_dict, drop_col_list) :
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new_df = df[keep_col_list]
   new_df.rename(columns = rename_col_dict, inplace = True)
   new_df.dropna(subset = drop_col_list, inplace = True)
   return new_df
diag_vis1 = condensed_df(full_df,['PATNO', 'DIAG1', 'DIAG1VIS'], {'DIAG1VIS' :

    'EVENT ID'}, ['EVENT ID'])

diag vis2 = condensed df(full df, ['PATNO', 'DIAG2', 'DIAG2VIS'], {'DIAG2VIS' :

    'EVENT ID'},['EVENT ID'])

ppmi_merge.drop(['DIAG1', 'DIAG1VIS', 'DIAG2', 'DIAG2VIS'], axis = 1, inplace = True) #
→ Drop these from ppmi_merge so there aren't duplicates when we merge the diag_vis dfs
ppmi merge = pd.merge(diag vis1, ppmi merge, on = ['EVENT ID', 'PATNO'], how = "outer" )
→ # Merge in first diagnosis change
ppmi_merge = pd.merge(diag_vis2, ppmi_merge, on = ['EVENT_ID', 'PATNO'], how = "outer" )
→ # Merge in second diagnosis change
ppmi_merge['DIAG1'].replace({ 'PD' : 'Parkinson\'s Disease', 'DLB': 'Dimentia with Lewy
→ Bodies'}, inplace = True) # Decode
ppmi_merge['DIAG2'].replace({ 'MSA' : 'Multiple System Atrophy', 'DLB': 'Dimentia with
→ Lewy Bodies'}, inplace = True) # Decode
ppmi_merge.rename(columns = {'DIAG1' : 'First.Diagnosis.Change', 'DIAG2' :
→ 'Second.Diagnosis.Change'}, inplace = True) # Rename columns
## Dominant side of disease
ppmi merge = merge new csv(ppmi merge,
- 'PPMI_Original_Cohort_BL_to_Year_5_Dataset_Apr2020.csv',['PATNO', 'EVENT_ID',
→ 'DOMSIDE']) # Domside
ppmi_merge['DOMSIDE'].replace({1 : 'Left', 2 : 'Right', 3 : 'Symmetric'}, inplace = True)

→ # Decode

## Participant Motor Function Questionnaire
ppmi_merge = merge_new_csv(ppmi_merge, 'Participant_Motor_Function_Questionnaire.csv',
- ['PATNO', 'EVENT_ID', 'PAG_NAME', 'CMPLBY2', 'TRBUPCHR', 'WRTSMLR', 'VOICSFTR', 'POORBAL', 'FTSTUCK', 'LSSXP
ppmi_merge['CMPLBY2'].replace({1: 'Participant', 2: 'Caregiver', 3: 'Participant and
ppmi_merge['PAG_NAME'].replace({'PQUEST' : 'Participant Motor Function Questionnaire'},
→ inplace = True)
ppmi_merge = decode_0_1_no_yes(ppmi_merge, ['TRBUPCHR', 'WRTSMLR', 'VOICSFTR', 'POORBAL',
- 'FTSTUCK', 'LSSXPRSS', 'ARMLGSHK', 'TRBBUTTN', 'SHUFFLE', 'MVSLOW', 'TOLDPD'])
ppmi_merge = ppmi_merge.rename(columns = {'PAG_NAME' : 'Motor.Function.Page.Name',
→ 'CMPLBY2': 'Motor.Function.Source', 'TRBUPCHR': 'Trouble.Rising.Chair', 'WRTSMLR':
→ 'Writing.Smaller', 'VOICSFTR': 'Voice.Softer', 'POORBAL': 'Poor.Balance',
→ 'FTSTUCK' : 'Feet.Stuck', 'LSSXPRSS' : 'Less.Expressive',
   'ARMLGSHK':'Arms/Legs.Shake', 'TRBBUTTN': 'Trouble.Buttons', 'SHUFFLE':
→ 'Shuffle.Feet' , 'MVSLOW' : 'Slow.Movements' , 'TOLDPD' : 'Been.Told.PD' })
## Cognitive symptoms - Cognitive Categorization
ppmi_merge = merge_new_csv(ppmi_merge, 'Cognitive_Categorization.csv',['PATNO' ,
- 'EVENT_ID', 'PAG_NAME', 'COGDECLN', 'FNCDTCOG', 'COGDXCL', 'PTCGBOTH', 'COGSTATE',
→ 'COGCAT_TEXT']) # Visit date, weight and height
ppmi_merge = decode_0_1_no_yes(ppmi_merge, ['COGDECLN', 'FNCDTCOG'])
ppmi_merge['COGDXCL'].replace({1 : '90 - 100%', 2 : '50 - 89%', 3 : '10 - 49%', 4 : '0 -
ppmi_merge['PTCGBOTH'].replace({1 : 'Participant', 2 : 'Caregiver', 3 : 'Participant and
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ppmi_merge['COGSTATE'].replace({1 : 'Normal Condition', 2 : 'Mild Cognitive Impairment',
ppmi_merge = ppmi_merge.rename(columns = {'PAG_NAME' : 'Cognitive.Page.Name', 'COGDECLN'
→ : 'Cognitive.Decline', 'FNCDTCOG' : 'Functional.Cognitive.Impairment', 'COGDXCL' :
   'Confidence.Level.Cognitive.Diagnosis', 'PTCGBOTH' : 'Cognitive.Source', 'COGSTATE' :
ppmi merge['Cognitive.Page.Name'].replace({'COGCATG' : 'Cognitive Categorization'},

    inplace = True) # Rename

## Cognitive symptoms - MOCA
ppmi_merge = merge_new_csv(ppmi_merge,
→ 'Montreal Cognitive Assessment MoCA .csv',['PATNO', 'EVENT ID', 'MCATOT'])
ppmi_merge.rename(columns = {'MCATOT' : 'MOCA.Total'}, inplace = True) # Rename
## Medication Status - Concomitant Med Log
med_df = pd.read_csv(ppmi_download_path + 'Concomitant_Medication_Log.csv',
med_df.replace({';' : ','}, regex = True, inplace = True) # Replace ';' with ','
med_df['CMTRT'] = med_df['CMTRT'].str.title() # Capitalize all medication names
med_df['STARTDT'].fillna('NA', inplace = True) # Fillna
med_df['STOPDT'].fillna('NA', inplace = True) # Fillna
med_df = med_df.astype({"STARTDT" : 'str', "STOPDT" : 'str'}) # Change start and stop date
med_df = merge_columns(med_df, ['STARTDT', 'STOPDT'], 'Start_Stop', '-') # Merge columns
→ Start and stop date together
med_df['Start_Stop'] = '(' + med_df['Start_Stop'].astype(str) + ')'# Put parenthesis
→ around dates so when you merge it with LED_med_and dose it is more organized
med_df = merge_columns(med_df, ['CMTRT', 'Start_Stop'], 'Medication', ' ')
med df = med df.groupby(['PATNO', 'EVENT ID'])['Medication'].apply('; '.join)
ppmi_merge = pd.merge(ppmi_merge, med_df, on = ['PATNO', 'EVENT_ID'], how = "outer") #

    Merge med_df in

ppmi_merge = ppmi_merge.sort_values(by = ['PATNO', 'Age']).reset_index(drop = True) # Sort
→ values by subject and age (similar to event id bc age in order of event id)
## LEDD Medication Status
LEDD_med_df = pd.read_csv(ppmi_download_path + 'LEDD_Concomitant_Medication_Log.csv',
LEDD_med_df = LEDD_med_df[['PATNO', 'LEDD', 'STARTDT', 'STOPDT']]
ppmi_merge['INFODT'] = pd.to_datetime(ppmi_merge['INFODT'], format = '%m%Y', errors =
→ 'ignore') # change to INFODT to type datetime so we can sort according to date
LEDD med df['STARTDT'] = pd.to datetime(LEDD med df['STARTDT'], format = '%m%Y', errors =
→ 'ignore') # change to INFODT to type datetime so we can sort according to date
LEDD_med_df['STOPDT'] = pd.to_datetime(LEDD_med_df['STOPDT'], format = '%m%Y', errors =
→ 'ignore') # change to INFODT to type datetime so we can sort according to date
LEDD_med_df['STOPDT2'] = LEDD_med_df['STOPDT'] # Initialize second stop date variable
LEDD_med_df['STOPDT2'].fillna(LEDD_med_df['STARTDT'], inplace = True)# Fill in NaN stop
\hookrightarrow dates with start dates
LEDD_med_df = LEDD_med_df.merge(LEDD_med_df.apply(lambda s: pd.date_range(s.STARTDT,

    s.STOPDT2, freq='MS', inclusive = 'both'),
→ 1).explode().rename('INFODT').dt.strftime('%m/%Y'), left_index=True,
→ right_index=True) # Create a row for every month/year for LEDD
LEDD_med_df.drop(['STOPDT2'], axis = 1, inplace = True)
LEDD_med_df.rename(columns = {'STARTDT' : 'LEDD.STARTDT', 'STOPDT': 'LEDD.STOPDT'},

    inplace = True) # Rename
```

```
# If one event id has > 1 LEDD - take max LEDD only
ppmi_merge_temp = ppmi_merge[['PATNO', 'EVENT_ID', 'INFODT']]
ppmi_merge_temp = pd.merge(ppmi_merge_temp, LEDD_med_df, on = ['PATNO', 'INFODT'], how =

    "left")

ppmi_merge_temp = ppmi_merge_temp[['PATNO', 'EVENT_ID', 'LEDD', 'LEDD.STARTDT',
→ 'LEDD.STOPDT']] # keep only
ppmi_merge_temp = ppmi_merge_temp[ppmi_merge_temp["LEDD"].str.contains("LD") == False] #
\rightarrow FIXME not sure what to do with LD x .33 etc rows - for now remove
uids = ppmi_merge_temp['PATNO'].unique() # Unique subject ids
myevs = ppmi_merge_temp['EVENT_ID'].unique() # Unique event ids
ppmi merge temp['LEDD2'] = False
for myev in myevs :
      for u in uids :
              selu = ppmi_merge_temp.loc[(ppmi_merge_temp['PATNO'] == u) &
composition of the composit
              if len(selu) == 1:
                     idx = selu.index
                     ppmi_merge_temp.loc[idx, 'LEDD2'] = True
              if len(selu) > 1:
                     selu['LEDD'] = selu['LEDD'].astype(float)
                     maxidx = selu.LEDD.idxmax()
                     ppmi_merge_temp.loc[maxidx, 'LEDD2'] = True
ledd_keep = ppmi_merge_temp['LEDD2'] == True
LEDD_new_df = ppmi_merge_temp[ledd_keep]
LEDD_new_df.drop(['LEDD2'], axis = 1, inplace = True)
ppmi_merge = pd.merge(ppmi_merge, LEDD_new_df, on = ['PATNO', 'EVENT_ID'], how = "left") #
→ Merge into ppmi merge
## Comorbidities
comorbid_df = pd.read_csv(ppmi_download_path + 'Medical_Conditions_Log.csv',

    skipinitialspace=True) # Medication history

comorbid_df.replace({';' : ','}, regex = True, inplace = True) # Replace ';' with ','
comorbid_df = comorbid_df[['PATNO', 'EVENT_ID', 'MHDIAGDT', 'MHTERM']] # keep only
comorbid_df['MHTERM'] = comorbid_df['MHTERM'].str.capitalize() # Capitalize all MHTERM
comorbid_df['MHDIAGDT'].fillna('NA', inplace = True) # If no diagnosis date - fill in
\hookrightarrow with NA
comorbid_df['MHDIAGDT'] = '(' + comorbid_df['MHDIAGDT'].astype(str) + ')'# Put
→ parentheses around diagnosis date
comorbid_df = comorbid_df.astype({"MHDIAGDT" : 'str'}) # Change date to string
comorbid_df = merge_columns(comorbid_df, ['MHTERM', 'MHDIAGDT'],
→ 'Medical.History.Description(Diagnosis.Date)' , ' ')
comorbid_df =
comorbid_df.groupby(['PATNO', 'EVENT_ID'])['Medical.History.Description(Diagnosis.Date)'].apply(';
\hookrightarrow '.join)
ppmi_merge = pd.merge(ppmi_merge, comorbid_df, on = ['PATNO', 'EVENT_ID'], how = "outer")
## Education (in years) - FIXME make integer
education_df = pd.read_csv(ppmi_download_path + 'Socio-Economics.csv', skipinitialspace =
→ True) # Education info
education_df = education_df[['PATNO', 'EVENT_ID', 'EDUCYRS']] # Keep info
```

```
education_df.rename(columns = {'EDUCYRS' : 'Education.Years'}, inplace = True) # Rename
education_df = education_df.groupby('PATNO').mean().reset_index() # Take the mean of
- education years if there are 2 different number of years for one subject
ppmi_merge = pd.merge(ppmi_merge, education_df, on = ['PATNO'], how = "outer") # Merge
## Add in analytic cohort column
analytic cohort = ppmi merge['PATNO'].isin(analytic cohort subids)
ppmi_merge['Analytic.Cohort'] = '' # Initialize Analytic.Cohort col
ppmi_merge['Analytic.Cohort'].loc[analytic_cohort] = 'Analytic Cohort'
ppmi_merge['Analytic.Cohort'].fillna('Not Analytic Cohort', inplace = True)
## Reindex
ppmi_merge = ppmi_merge.reindex(columns = ['PATNO', 'EVENT_ID', 'INFODT' ,
→ 'Enroll.Diagnosis', 'Enroll.Subtype', 'Consensus.Diagnosis',
→ 'Consensus.Subtype', 'Analytic.Cohort', 'Subject.Phenoconverted', 'First.Diagnosis.Change',
→ 'Second.Diagnosis.Change', 'First.Symptom.Date', 'PD.Diagnosis.Date',
→ 'PD.Diagnosis.Duration', 'BirthDate', 'Age', 'Sex', 'Handed', 'Weight(kg)',
→ 'Height(cm)', 'Education.Years', 'DOMSIDE', 'Motor.Function.Page.Name',
→ 'Motor.Function.Source', 'Trouble.Rising.Chair', 'Writing.Smaller',
→ 'Voice.Softer', 'Poor.Balance', 'Feet.Stuck', 'Less.Expressive',
_{\rightarrow} 'Arms/Legs.Shake', 'Trouble.Buttons', 'Shuffle.Feet', 'Slow.Movements',
→ 'Been.Told.PD', 'Cognitive.Page.Name', 'Cognitive.Source', 'Cognitive.Decline',
→ 'Functional.Cognitive.Impairment', 'Confidence.Level.Cognitive.Diagnosis',
→ 'Cognitive.State', 'Cognitive.Tscore.Cat', 'MOCA.Total', 'Medication', 'LEDD',
→ 'LEDD.STARTDT', 'LEDD.STOPDT', 'Medical.History.Description(Diagnosis.Date)'])
## Add in other csvs
ppmi_merge = merge_new_csv(ppmi_merge, 'Modified_Boston_Naming_Test.csv', ['PATNO',
→ 'EVENT ID', 'MBSTNSCR', 'MBSTNCRC', 'MBSTNCRR', 'MBSTNVRS'])
ppmi_merge = merge_new_csv(ppmi_merge, 'Clock_Drawing.csv', ['PATNO', 'EVENT_ID',
ppmi_merge = merge_new_csv(ppmi_merge, 'Benton_Judgement_of_Line_Orientation.csv',
ppmi_merge = merge_new_csv(ppmi_merge, 'Letter_-_Number_Sequencing.csv', ['PATNO',

    'EVENT_ID', 'LNS_TOTRAW'])

ppmi_merge = merge_new_csv(ppmi_merge, 'Modified_Semantic_Fluency.csv', ['PATNO',

    'EVENT_ID', 'DVS_SFTANIM'])

ppmi_merge = merge_new_csv(ppmi_merge,
→ 'Hopkins_Verbal_Learning_Test_-_Revised.csv',['PATNO', 'EVENT_ID',

    'DVT_DELAYED_RECALL', 'DVT_TOTAL_RECALL'])
ppmi_merge = merge_new_csv(ppmi_merge, 'Symbol_Digit_Modalities_Test.csv', ['PATNO',

    'EVENT ID', 'SDMTOTAL'])

ppmi_merge.rename(columns = {'CLCKTOT' : 'Clock.Drawing.Total', 'JLOTOTCALC' :
→ 'JOLO.Total', 'LNS_TOTRAW' : 'Letter.Number.Sequencing.Total', 'DVS_SFTANIM' :
→ 'Semantic.Fluency.Total','DVT_TOTAL_RECALL' : 'DVT.Total.RECALL','SDMTOTAL' :
## REM Sleeep behavior disorder questionnaire
ppmi_merge = merge_new_csv(ppmi_merge, 'REM_Sleep_Behavior_Disorder_Questionnaire.csv',
_{\rightarrow} ['PATNO', 'EVENT_ID', 'PAG_NAME', 'PTCGBOTH', 'DRMVIVID', 'DRMAGRAC', 'DRMNOCTB',
→ 'SLPLMBMV', 'SLPINJUR', 'DRMVERBL', 'DRMFIGHT', 'DRMUMV', 'DRMOBJFL', 'MVAWAKEN',
→ 'DRMREMEM', 'SLPDSTRB', 'STROKE', 'HETRA',
                                                 'PARKISM', 'RLS', 'NARCLPSY',
→ 'DEPRS', 'EPILEPSY', 'BRNINFM', 'CNSOTH'])
```

```
ppmi_merge.rename(columns = {'PAG_NAME' : 'REM.Sleep.Behavior.Disorder.Page.Name'},

    inplace = True)

ppmi_merge['REM.Sleep.Behavior.Disorder.Page.Name'].replace({'REMSLEEP' : 'REM Sleep
⇔ Behavior Disorder Questionnaire'}, inplace = True)
ppmi_merge['PTCGBOTH'].replace({1 : 'Participant', 2 : 'Caregiver', 3: 'Participant and
rem list = ['DRMVIVID', 'DRMAGRAC', 'DRMNOCTB', 'SLPLMBMV', 'SLPINJUR', 'DRMVERBL',
-> 'DRMFIGHT', 'DRMUMV', 'DRMOBJFL', 'MVAWAKEN', 'DRMREMEM', 'SLPDSTRB', 'STROKE',
              'PARKISM', 'RLS', 'NARCLPSY', 'DEPRS', 'EPILEPSY', 'BRNINFM',
→ 'HETRA',
ppmi_merge['RBDTotal.REM'] = ppmi_merge[rem_list].sum(axis = 1) # Add an RBDTotal.REM
ppmi_merge = decode_0_1_no_yes(ppmi_merge, rem_list)
ppmi_merge.rename(columns = {'PTCGBOTH' : 'Sleep.Behavior.Source.REM', 'DRMVIVID' :
→ 'Vivid.Dreams.REM' , 'DRMAGRAC': 'Aggressive.or.Action-packed.Dreams.REM',
→ 'DRMNOCTB':'Nocturnal.Behaviour.REM', 'SLPLMBMV':'Move.Arms/legs.During.Sleep.REM',
→ 'SLPINJUR': 'Hurt.Bed.Partner.REM', 'DRMVERBL': 'Speaking.in.Sleep.REM', 'DRMFIGHT':
→ 'Sudden.Limb.Movements.REM', 'DRMUMV': 'Complex.Movements.REM',
→ 'EPILEPSY': 'Epilepsy.REM', 'BRNINFM': 'Inflammatory.Disease.of.the.Brain.REM',
#### IMAGING INFO ####
# FIXME - laterality issue? SUV ?
ppmi merge = merge new csv(ppmi merge, 'DaTScan Analysis.csv',
- ['PATNO','EVENT_ID','DATSCAN_DATE','DATSCAN_CAUDATE_R','DATSCAN_CAUDATE_L','DATSCAN_PUTAMEN_R','DAT
ppmi_merge = merge_new_csv(ppmi_merge, 'DaTScan_Imaging.csv',
- ['PATNO', 'EVENT_ID', 'PAG_NAME', 'INFODT', 'DATSCAN', 'PREVDATDT', 'SCNLOC', 'SCNINJCT', 'VSINTRPT', 'VSRPT
ppmi_merge['DATSCAN'].replace({0 : 'Not Completed', 1: 'Completed', 2 : 'Completed using
→ a previously acquired DaTscan (i.e., acquired prior to participant\'s consent to
→ PPMI)' }, inplace = True)
ppmi_merge['SCNLOC'].replace({ 1 : 'Site', 2 : 'IND'}, inplace = True )
ppmi_merge['SCNINJCT'].replace({ 1 : 'DaTSCAN', 2 : 'Beta-CIT'}, inplace = True)
ppmi_merge['VSINTRPT'].replace({ '1' : 'Consistent with evidence' , '2' : 'Not consistent
→ with evidence', '3' : 'No visual interpretation report provided'}, inplace = True)
ppmi_merge['VSRPTELG'].replace({1 : 'Eligible', 2 : 'Not eligible'}, inplace = True)
ppmi merge.rename(columns = {'INFODT x' : 'INFODT', 'INFODT y' : 'DaTScan.INFODT',
\hookrightarrow 'SCNLOC' : 'Location.Scan.Completed' , 'PREVDATDT' :
→ 'Date.DaTscan.Imaging.Completed.Previously', 'SCNINJCT': 'Scan.Injection',
→ 'VSINTRPT' : 'Visual.Interpretation.Report', 'VSRPTELG' :
→ 'Visual.Interpretation.Report(eligible/not)'}, inplace = True)
ppmi_merge['INFODT'].fillna(ppmi_merge['DaTScan.INFODT'], inplace = True)# Fill in NaN
→ infodts with datscan infodts if na
## MRI.csv
mri_df = pd.read_csv(ppmi_download_path + 'Magnetic_Resonance_Imaging_MRI_.csv',

    skipinitialspace=True)

mri_df = mri_df[[ 'PATNO', 'EVENT_ID', 'INFODT', 'MRICMPLT', 'MRIWDTI', 'MRIWRSS',
→ 'MRIRSLT', 'MRIRSSDF' ]] # Keep only
```

```
mri_df['MRICMPLT'].replace({0 : 'Not Completed', 1 : 'Completed'}, inplace = True) #
→ Decode
mri_df = decode_0_1_no_yes(mri_df, ['MRIWDTI', 'MRIWRSS', 'MRIRSSDF'])
mri_df['MRIRSLT'].replace({1 : 'Normal', 2 : 'Abnormal, not clinically significant', 3 :
→ 'Abnormal, clinically significant'}, inplace = True) # Decode
mri_df.rename(columns = { 'INFODT' : 'Image.Acquisition.Date', 'MRICMPLT' :
→ 'MRI.Completed', 'MRIWDTI': 'MRI.DTI', 'MRIWRSS': 'MRI.Resting.State', 'MRIRSLT'
- : 'MRI.Results', 'MRIRSSDF': 'Resting.State.Dif.Day.PDMed.Use'}, inplace = True) #
\rightarrow Rename
## FIXME - do we want to get rid of these?
# Some subjects had two baseline rows - 1 with incomplete MRI. Completed and 1 with
\rightarrow complete as MRI.Complete - I am only keeping the one that is complete bc the data we
→ have on s3 is complete
duplicate_mri = mri_df[['PATNO','EVENT_ID']].duplicated(keep = False) # Find locations of
→ True for duplicated subs w/ 2 MRI at baseline
duplicate_mri_index = mri_df[duplicate_mri == True].index.tolist() # Get index of
\rightarrow duplicates
dup_subid_list = [] # Initialize duplicate subid list variable
[dup_subid_list.append(index) for index in duplicate_mri_index if
→ mri_df['MRI.Completed'][index] == 'Not Completed'] # Get the indices of duplicate
\Rightarrow subids that were labeled as Not Completed
mri_df = mri_df.reset_index(drop = True)
[mri_df.drop(index = i, axis = 1, inplace = True) for i in reversed(dup_subid_list) if
→ mri_df['MRI.Completed'][i] == 'Not Completed'] # Get rid of the duplicate subids that
\rightarrow were labeled as Not Completed (that also have another labeled as completed)
ppmi_merge = pd.merge(ppmi_merge, mri_df, on = ['PATNO', 'EVENT_ID'], how = "outer")
## For all subjects - see if we have T1 images in ppmi-image-data bucket for given dates
def search_s3(bucket, prefix, search_string):
    client = boto3.client('s3', region_name="us-east-1")
   paginator = client.get_paginator('list_objects')
   pages = paginator.paginate(Bucket=bucket, Prefix=prefix)
   keys = []
   for page in pages:
        contents = page['Contents']
        for c in contents:
           keys.append(c['Key'])
    if search_string:
        keys = [key for key in keys if search_string in key]
    return keys
keys = search_s3('invicro-ia-object-repository', 'refined/ppmi/data/PPMI/', 'T1w/') #
→ PPMI1.0 and PPMI2.0
# Set a variable in ppmi_merge 'Subid.Date.TEMP' that is the subid and image acquisition
→ date to match s3 image to
ppmi_merge['Subid.Date.TEMP'] = ''
for row_num in range(len(ppmi_merge['Image.Acquisition.Date'])) :
    if isinstance(ppmi_merge['Image.Acquisition.Date'].loc[row_num], str) :
        ppmi_merge['Subid.Date.TEMP'].iloc[row_num] =
→ ppmi_merge["PATNO"].iloc[row_num].astype(str) + '/' +
→ ppmi_merge["Image.Acquisition.Date"].iloc[row_num].split('/')[1] +
→ ppmi_merge['Image.Acquisition.Date'].iloc[row_num].split('/')[0] # Combine subid and
\rightarrow date into 1 col in df
```

```
subid_date_ordered = ppmi_merge['Subid.Date.TEMP'].dropna().tolist() # Make column into
\hookrightarrow list
# Get keys of subjects in ppmi-image-data bucket with T1w/ image folder
woutppmi = [key.split('PPMI/')[1] for key in keys] # Remove PPMI/ from key
woutt1w = [key.split('/T1w/')[0] for key in woutppmi] # Remove 'T1w' from key
s3woutdate = [current_subid_date[:-2] for current_subid_date in woutt1w] # Remove the day
→ from date - want only yearmonth i.e. 202106
matches = [current_subid_date for current_subid_date in subid_date_ordered if
→ current_subid_date in s3woutdate] # PPMI images that are in S3 that have T1w
# Add in T1 s3 Info
s3_df = pd.DataFrame(columns = ['PATNO', 'EVENT_ID', 'T1.s3.Image.Name']) # create s3_df
\rightarrow dataframe
for current_subid_date_temp in matches :
    for image_id in woutppmi :
        if current_subid_date_temp in image_id and image_id.endswith('.nii.gz'):
            image id split = image id.split('/')
            s3_df = s3_df.append({'PATNO' : image_id_split[0], 'EVENT_ID' :
→ ppmi_merge.loc[ppmi_merge['Subid.Date.TEMP'] == current_subid_date_temp, 'EVENT_ID'].iloc[0],

    'T1.s3.Image.Name' : image_id_split[-1]}, ignore_index = True)

# Create a column in s3_df for just object name so later we can merge Taylor's T1 file
→ with object name
s3_df['PATNO'] = s3_df['PATNO'].astype(int) # needed for merge on PATNO
s3 df['Image ID merge'] = ''
for row_num in range(len(s3_df['T1.s3.Image.Name'])) :
    image0 = s3_df['T1.s3.Image.Name'].iloc[row_num].split('.')[0] # Get name of image

    before .nii.qz

   s3_df['Image_ID_merge'].iloc[row_num] = image0.split('-')[4] # Get ImageID from s3
→ filename and put in Image_ID_merge column
ppmi_merge = pd.merge(ppmi_merge, s3_df, on = ['PATNO', 'EVENT_ID'], how = "outer")
#### MDS-UPDRS Scores UPDRS 1-4 as Numeric Variables ####
def setup_updrs_df(updrs_filename, drop_cols, decode_dict, rename_col_dict) :
    updrs_df = pd.read_csv(ppmi_download_path + updrs_filename, skipinitialspace = True)
   updrs_df.drop(drop_cols, axis = 1, inplace = True)
   updrs_df['PAG_NAME'].replace(decode_dict, inplace = True)
   updrs_df.rename(columns = rename_col_dict, inplace = True)
   return updrs df
updrs_part1_df =

→ setup_updrs_df('MDS-UPDRS_Part_I.csv',['ORIG_ENTRY','LAST_UPDATE','REC_ID'],

→ {'NUPDRS1' : 'MDS-UPDRS Part I: Non-Motor Aspects of Experiences of Daily Living'},
→ {'PAG_NAME' : 'UPDRS.Part1.Page.Name', 'NUPSOURC' : 'UPDRS.Part1.Source'})
updrs_part1_pq_df = setup_updrs_df('MDS-UPDRS_Part_I_Patient_Questionnaire.csv',
→ ['ORIG_ENTRY', 'LAST_UPDATE', 'REC_ID', 'INFODT'], {'NUPDRS1P' : 'MDS-UPDRS Part I
-- Patient Questionnaire: Non-Motor Aspects of Experiences of Daily Living', 'NUPDRSP':
  'MDS-UPDRS Part IB and Part II'}, {'PAG_NAME' :
→ 'UPDRS.Part1.Patient.Questionnaire.Page.Name', 'NUPSOURC' : 'UPDRS.Part1.PQ.Source'})
updrs_part2_pq_df = setup_updrs_df('MDS_UPDRS_Part_II__Patient_Questionnaire.csv',
→ ['ORIG_ENTRY', 'LAST_UPDATE', 'REC_ID', 'INFODT'], {'NUPDRS2P' : 'MDS-UPDRS Part II
-- Patient Questionnaire: Motor Aspects of Experiences of Daily Living', 'NUPDRSP' :
→ 'MDS-UPDRS Part IB and Part II'}, {'PAG_NAME' : 'UPDRS.Part2.Page.Name', 'NUPSOURC' :

    'UPDRS.Part2.Source'})
                                            13
```

```
updrs_part3_dos_df =

→ setup_updrs_df('MDS-UPDRS_Part_III_ON_OFF_Determination___Dosing.csv',
→ ['ORIG_ENTRY','LAST_UPDATE','REC_ID','INFODT'], {'NUPDRDOSE' : 'MDS-UPDRS Part III
\hookrightarrow ON/OFF Determination & Dosing', 'NUPDRDOSER' : 'MDS-UPDRS Part III examination
→ administered at remote visit'}, {'PAG_NAME' : 'UPDRS.Part3.Dosage.Page.Name'})
updrs_part4_motor_df = setup_updrs_df('MDS-UPDRS_Part_IV__Motor_Complications.csv',
→ ['ORIG ENTRY', 'LAST UPDATE', 'REC ID', 'INFODT'], {'NUPDRS4' : 'MDS-UPDRS Part IV:
→ Motor Complications'}, {'PAG_NAME' : 'UPDRS.Part4.Page.Name'})
# UPDRS Part 3
updrs_part3_df = pd.read_csv(ppmi_download_path + 'MDS_UPDRS_Part_III.csv',

    skipinitialspace = True)

updrs_part3_df.drop(['ORIG_ENTRY','LAST_UPDATE','REC_ID','INFODT'], axis = 1, inplace =
→ True)
updrs_part3_df.rename(columns = {'PAG_NAME' : 'UPDRS.Part3.Page.Name'}, inplace = True)
# Split up UPDRS Part 3 into four parts
nupdrs3 = updrs_part3_df.loc[updrs_part3_df['UPDRS.Part3.Page.Name'] == 'NUPDRS3']
nupdrs3A = updrs_part3_df.loc[updrs_part3_df['UPDRS.Part3.Page.Name'] == 'NUPDRS3A']
nupdr3OF = updrs_part3_df.loc[updrs_part3_df['UPDRS.Part3.Page.Name'] == 'NUPDR3OF']
nupdr30N = updrs_part3_df.loc[updrs_part3_df['UPDRS.Part3.Page.Name'] == 'NUPDR30N']
# Change all UPDRS dataframe cols begin with 'N' to floats
updrs_list = [updrs_part1_df, updrs_part1_pq_df,updrs_part2_pq_df, nupdrs3, nupdrs3A,
-- nupdr30F, nupdr30N, updrs_part3_dos_df, updrs_part4_motor_df]
for df in updrs_list :
   for col_name in df :
        if col_name.startswith('N') :
            df[col name] = pd.to numeric(df[col name], errors = 'coerce', downcast =
→ 'float')
# Create a copy of each dataframe to use later to create categorical versions of
\hookrightarrow varibales
updrs_part1_df_copy = updrs_part1_df.copy()
updrs_part1_pq_df_copy = updrs_part1_pq_df.copy()
updrs_part2_pq_df_copy = updrs_part2_pq_df.copy()
nupdrs3_copy = nupdrs3.copy()
nupdrs3A_copy = nupdrs3A.copy()
nupdr30F_copy = nupdr30F.copy()
nupdr30N_copy = nupdr30N.copy()
updrs_part3_dos_df_copy = updrs_part3_dos_df.copy()
updrs_part4_motor_df_copy = updrs_part4_motor_df.copy()
# For all UPDRS of columns - add the respective extension for which UPDRS assessment it
def add_extension_to_column_names(df, skip_col_list, ext):
   for col_name in df :
        if col_name not in skip_col_list :
            df.rename(columns = {str(col_name) : str(col_name) + ext }, inplace = True)
    return df
# Add extensions to updrs dfs
updrs_list_str = ['.UPDRS1','.UPDRS1','.UPDRS2', '.UPDRS3','.UPDRS3A', '.UPDR30F',
→ '.UPDR3ON','.UPDRDOSE','.UPDRS4'] # extensions
```

```
for i in range(len(updrs_list)) :
    updrs_list[i] = add_extension_to_column_names(updrs_list[i],
# Merge the four UPDRS3 dfs together
updrs3 = pd.merge(nupdrs3, nupdrs3A, on = ['PATNO', 'EVENT_ID'], how = "outer")
updrs3 = pd.merge(updrs3, nupdr30F, on = ['PATNO', 'EVENT_ID'], how = "outer")
updrs3 = pd.merge(updrs3, nupdr30N, on = ['PATNO', 'EVENT_ID'], how = "outer")
# Create one df for updrs_numeric
updrs_numeric = pd.merge(updrs_part1_df, updrs_part1_pq_df , on = ['PATNO', 'EVENT_ID'],
→ how = "outer")
updrs_numeric = pd.merge(updrs_numeric, updrs_part2_pq_df , on = ['PATNO', 'EVENT_ID'],
→ how = "outer")
updrs_numeric = pd.merge(updrs_numeric, updrs3, on = ['PATNO', 'EVENT_ID'], how =
→ "outer")
updrs_numeric = pd.merge(updrs_numeric,updrs_part3_dos_df , on = ['PATNO', 'EVENT_ID'],
→ how = "outer")
updrs_numeric = pd.merge(updrs_numeric, updrs_part4_motor_df, on = ['PATNO', 'EVENT_ID'],
→ how = "outer")
## Keep only variables in .Num that are numeric variables
numeric_vars = []
for col_name in updrs_numeric :
    if col_name.startswith('NP') or col_name.startswith('PATNO') or

    col_name.startswith('EVENT') or col_name.startswith('NHY'):

       numeric_vars.append(col_name)
updrs_numeric = updrs_numeric[numeric_vars]
# Rename columns in updrs_numeric
updrs_numeric.rename(columns = {'NHY.UPDRS3' : 'Hoehn.and.Yahr.Stage.UPDRS3',
'NP3BRADY.UPDRS3' : 'Global.Spontaneity.of.Movement.UPDRS3', 'NP3PTRMR.UPDRS3' :
'Postural.Tremor.Right.Hand.UPDRS3' , 'NP3PTRML.UPDRS3' : 'Postural.Tremor.Left.Hand.UPDRS3'
, 'NP3KTRMR.UPDRS3' : 'Kinetic.Tremor.Right.Hand.UPDRS3', 'NP3KTRML.UPDRS3' :
'Kinetic.Tremor.Left.Hand.UPDRS3', 'NP3RTARU.UPDRS3': 'Rest.Tremor.Amplitude.RUE.UPDRS3',
'NP3RTALU.UPDRS3' : 'Rest.Tremor.Amplitude.LUE.UPDRS3', 'NP3RTARL.UPDRS3' :
'Rest.Tremor.Amplitude.RLE.UPDRS3' ,'NP3RTALL.UPDRS3' : 'Rest.Tremor.Amplitude.LLE.UPDRS3'
,'NP3RTALJ.UPDRS3' : 'Rest.Tremor.Amplitude.Lip.Jaw.UPDRS3', 'NP3RTCON.UPDRS3' :
'Constancy.of.Rest.Tremor.UPDRS3', 'NP3SPCH.UPDRS3' : 'UPDRS3.Speech.Difficulty.UPDRS3',
'NP3FACXP.UPDRS3': 'Facial.Expression.Difficulty.UPDRS3', 'NP3RIGN.UPDRS3':
'Rigidity.Neck.UPDRS3' , 'NP3RIGRU.UPDRS3' : 'Rigidity.RUE.UPDRS3', 'NP3RIGLU.UPDRS3'
: 'Rigidity.LUE.UPDRS3', 'NP3RIGRL.UPDRS3' : 'Rigidity.RLE.UPDRS3', 'NP3RIGLL.UPDRS3'
: 'Rigidity.LLE.UPDRS3', 'NP3FTAPR.UPDRS3' : 'Finger.Tapping.Right.Hand.UPDRS3'
,'NP3FTAPL.UPDRS3' : 'Finger.Tapping.Left.Hand.UPDRS3' ,'NP3HMOVR.UPDRS3' :
'Hand.Movements.Right.Hand.UPDRS3', 'NP3HMOVL.UPDRS3': 'Hand.Movements.Left.Hand.UPDRS3',
'NP3PRSPR.UPDRS3' : 'Pronation.Supination.Right.Hand.UPDRS3', 'NP3PRSPL.UPDRS3' :
'Pronation.Supination.Left.Hand.UPDRS3' , 'NP3TTAPR.UPDRS3' : 'Toe.Tapping.Right.Foot.UPDRS3'
, 'NP3TTAPL.UPDRS3' : 'Toe.Tapping.Left.Foot.UPDRS3', 'NP3LGAGR.UPDRS3' :
'Leg.Agility.Right.Leg.UPDRS3', 'NP3LGAGL.UPDRS3' : 'Leg.Agility.Left.Leg.UPDRS3',
'NP3RISNG.UPDRS3' : 'UPDRS3.Rising.from.Chair.UPDRS3', 'NP3GAIT.UPDRS3'
: 'Gait.Problems.UPDRS3' ,'NP3FRZGT.UPDRS3' : 'Freezing.of.Gait.UPDRS3'
,'NP3PSTBL.UPDRS3': 'Postural.Stability.Problems.UPDRS3', 'NP3POSTR.UPDRS3'
: 'Posture.Problems.UPDRS3', 'NP3TOT.UPDRS3':'UPDRS.Part3.Total.UPDRS3',
'NP3SPCH.UPDRS3A' : 'UPDRS3.Speech.Difficulty.UPDRS3A', 'NP3FACXP.UPDRS3A' :
'Facial.Expression.Difficulty.UPDRS3A' , 'NP3RIGN.UPDRS3A' : 'Rigidity.Neck.UPDRS3A'
, 'NP3RIGRU.UPDRS3A' : 'Rigidity.RUE.UPDRS3A' : 'NP3RIGLU.UPDRS3A' : 'Rigidity.LUE.UPDRS3A', 'NP3RIGLL.UPDRS3A' : 'Rigidity.RLE.UPDRS3A', 'NP3RIGLL.UPDRS3A'
: 'Rigidity.LLE.UPDRS3A', 'NP3FTAPR.UPDRS3A' : 'Finger.Tapping.Right.Hand.UPDRS3A'
,'NP3FTAPL.UPDRS3A' : 'Finger.Tapping.Left.Hand.UPDRS3A' ,'NP3HMOVR.UPDRS3A' :
'Hand Movements Right Hand UPDRS3A' 'NP3HMOVI UPDRS3A' · 'Hand Movements Left Hand UPDRS3A'
```

```
## Subscore info from :
# https://www.ncbi.nlm.nih.gov/pmc/articles/PMC7242837/
→ https://www.movementdisorders.org/MDS-Files1/PDFs/Rating-Scales/MDS-UPDRS_English_FINAL.pdf
# FIXME - if one measurement is missing in caclulating subscores - do not calculate
→ subscores for this subject?
\#\# Add Brady-Rigidity, Tremor and PIGD Subscores
def calculate_subscores(df, new_col_name, subscore_list) :
   subscore_only = df[subscore_list] # extract df of only cols we are interested in for
df[new_col_name] = 0 # initalize new col for subscore
   idx = subscore_only.loc[pd.isnull(subscore_only).any(1), :].index
   df[new_col_name] = subscore_only.sum(axis = 1) # Get sum of subscore
   df[new_col_name].iloc[idx] = np.nan # Replace rows that should be nan with nan
   return df
updrs_numeric = calculate_subscores(updrs_numeric, 'Brady.Rigidity.Subscore.UPDRS3',
→ ['Rigidity.Neck.UPDRS3', 'Rigidity.RUE.UPDRS3', 'Rigidity.LUE.UPDRS3',
   'Rigidity.RLE.UPDRS3', 'Rigidity.LLE.UPDRS3', 'Finger.Tapping.Right.Hand.UPDRS3',
→ 'Finger.Tapping.Left.Hand.UPDRS3' ,
→ 'Hand.Movements.Right.Hand.UPDRS3', 'Hand.Movements.Left.Hand.UPDRS3', 'Pronation.Supination.Right.Ha
→ 'Pronation.Supination.Left.Hand.UPDRS3',
  'Toe.Tapping.Right.Foot.UPDRS3','Toe.Tapping.Left.Foot.UPDRS3','Leg.Agility.Right.Leg.UPDRS3',
updrs_numeric = calculate_subscores(updrs_numeric, 'Brady.Rigidity.Subscore.UPDRS3A',
→ ['Rigidity.Neck.UPDRS3A', 'Rigidity.RUE.UPDRS3A', 'Rigidity.LUE.UPDRS3A',
→ 'Rigidity.RLE.UPDRS3A', 'Rigidity.LLE.UPDRS3A', 'Finger.Tapping.Right.Hand.UPDRS3A'
, 'Finger.Tapping.Left.Hand.UPDRS3A', 'Hand.Movements.Right.Hand.UPDRS3A',
   'Hand.Movements.Left.Hand.UPDRS3A', 'Pronation.Supination.Right.Hand.UPDRS3A',
→ 'Pronation.Supination.Left.Hand.UPDRS3A', 'Toe.Tapping.Right.Foot.UPDRS3A',
→ 'Toe.Tapping.Left.Foot.UPDRS3A', 'Leg.Agility.Right.Leg.UPDRS3A',
→ 'Leg.Agility.Left.Leg.UPDRS3A'])
updrs_numeric = calculate_subscores(updrs_numeric, 'Brady.Rigidity.Subscore.UPDR3ON',
→ ['Rigidity.Neck.UPDR30N', 'Rigidity.RUE.UPDR30N', 'Rigidity.LUE.UPDR30N',
- 'Rigidity.RLE.UPDR30N', 'Rigidity.LLE.UPDR30N', 'Finger.Tapping.Right.Hand.UPDR30N'
, 'Finger.Tapping.Left.Hand.UPDR3ON', 'Hand.Movements.Right.Hand.UPDR3ON', 'Hand.Movements.Left.Hand.U
   'Pronation.Supination.Right.Hand.UPDR3ON', 'Pronation.Supination.Left.Hand.UPDR3ON',
  'Toe.Tapping.Right.Foot.UPDR30N', 'Toe.Tapping.Left.Foot.UPDR30N',
updrs_numeric = calculate_subscores(updrs_numeric, 'Brady.Rigidity.Subscore.UPDR3OF',
→ ['Rigidity.Neck.UPDR30F', 'Rigidity.RUE.UPDR30F', 'Rigidity.LUE.UPDR30F',
→ 'Rigidity.RLE.UPDR30F', 'Rigidity.LLE.UPDR30F', 'Finger.Tapping.Right.Hand.UPDR30F'
→ ,'Finger.Tapping.Left.Hand.UPDR30F' ,
   'Hand.Movements.Right.Hand.UPDR30F', 'Hand.Movements.Left.Hand.UPDR30F',
→ 'Pronation.Supination.Right.Hand.UPDR3OF', 'Pronation.Supination.Left.Hand.UPDR3OF',
→ 'Toe.Tapping.Right.Foot.UPDR30F', 'Toe.Tapping.Left.Foot.UPDR30F',

→ 'Leg.Agility.Right.Leg.UPDR30F', 'Leg.Agility.Left.Leg.UPDR30F'])

updrs_numeric = calculate_subscores(updrs_numeric, 'Tremor.Subscore.UPDRS3',
→ ['Tremor.UPDRS2', 'Postural.Tremor.Right.Hand.UPDRS3'
  ,'Postural.Tremor.Left.Hand.UPDRS3'
    , 'Kinetic.Tremor.Right.Hand.UPDRS3', 'Kinetic.Tremor.Left.Hand.UPDRS3',
→ 'Rest.Tremor.Amplitude.RUE.UPDRS3', 'Rest.Tremor.Amplitude.LUE.UPDRS3',
→ 'Rest.Tremor.Amplitude.RLE.UPDRS3' , 'Rest.Tremor.Amplitude.LLE.UPDRS3'
    'Rest.Tremor.Amplitude.Lip.Jaw.UPDRS3', 'Constancy.of.Rest.Tremor.UPDRS3'])
```

```
updrs numeric = calculate_subscores(updrs_numeric, 'Tremor.Subscore.UPDRS3A',
→ ['Tremor.UPDRS2', 'Postural.Tremor.Right.Hand.UPDRS3A'
    , 'Postural.Tremor.Left.Hand.UPDRS3A', 'Kinetic.Tremor.Right.Hand.UPDRS3A',
    'Kinetic.Tremor.Left.Hand.UPDRS3A', 'Rest.Tremor.Amplitude.RUE.UPDRS3A',
   'Rest.Tremor.Amplitude.LUE.UPDRS3A', 'Rest.Tremor.Amplitude.RLE.UPDRS3A'
  'Rest.Tremor.Amplitude.LLE.UPDRS3A', 'Rest.Tremor.Amplitude.Lip.Jaw.UPDRS3A',
  'Constancy.of.Rest.Tremor.UPDRS3A'])
updrs_numeric = calculate_subscores(updrs_numeric, 'Tremor.Subscore.UPDR30N',
→ ['Tremor.UPDRS2', 'Postural.Tremor.Right.Hand.UPDR30N'
  , 'Postural.Tremor.Left.Hand.UPDR3ON',
   'Kinetic.Tremor.Right.Hand.UPDR30N', 'Kinetic.Tremor.Left.Hand.UPDR30N',
→ 'Rest.Tremor.Amplitude.RUE.UPDR3ON', 'Rest.Tremor.Amplitude.LUE.UPDR3ON',
   'Rest.Tremor.Amplitude.RLE.UPDR3ON' , 'Rest.Tremor.Amplitude.LLE.UPDR3ON'
  'Rest.Tremor.Amplitude.Lip.Jaw.UPDR30N', 'Constancy.of.Rest.Tremor.UPDR30N'])
updrs_numeric = calculate_subscores(updrs_numeric, 'Tremor.Subscore.UPDR30F',
→ ['Tremor.UPDRS2', 'Postural.Tremor.Right.Hand.UPDR30F'
    , 'Postural.Tremor.Left.Hand.UPDR30F' ,
   'Kinetic.Tremor.Right.Hand.UPDR30F', 'Kinetic.Tremor.Left.Hand.UPDR30F',
'Rest.Tremor.Amplitude.Lip.Jaw.UPDR3OF', 'Constancy.of.Rest.Tremor.UPDR30F'])
updrs_numeric = calculate_subscores(updrs_numeric, 'PIGD.Subscore.UPDRS3',
→ ['Walking.Difficulty.UPDRS2', 'Freezing.while.Walking.UPDRS2'
→ ,'Gait.Problems.UPDRS3' , 'Freezing.of.Gait.UPDRS3' ,
→ 'Postural.Stability.Problems.UPDRS3'])
updrs_numeric = calculate_subscores(updrs_numeric, 'PIGD.Subscore.UPDRS3A',
\rightarrow ['Walking.Difficulty.UPDRS2',
→ 'Freezing.while.Walking.UPDRS2','Gait.Problems.UPDRS3A','Freezing.of.Gait.UPDRS3A',
→ 'Postural.Stability.Problems.UPDRS3A'])
updrs_numeric = calculate_subscores(updrs_numeric, 'PIGD.Subscore.UPDR3ON',
_{\hookrightarrow} ['Walking.Difficulty.UPDRS2' , 'Freezing.while.Walking.UPDRS2'
  , 'Gait.Problems.UPDR30N' , 'Freezing.of.Gait.UPDR30N' ,
  'Postural.Stability.Problems.UPDR30N'])
updrs_numeric = calculate_subscores(updrs_numeric, 'PIGD.Subscore.UPDR3OF',
\hookrightarrow ['Walking.Difficulty.UPDRS2', 'Freezing.while.Walking.UPDRS2',
→ 'Gait.Problems.UPDR30F' , 'Freezing.of.Gait.UPDR30F' ,
→ 'Postural.Stability.Problems.UPDR30F'])
updrs_numeric = add_extension_to_column_names(updrs_numeric, ['PATNO', 'EVENT_ID',
→ 'PIGD.Subscore.UPDRS3', 'Tremor.Subscore.UPDRS3', 'Brady.Rigidity.Subscore.UPDRS3',
  'PIGD.Subscore.UPDRS3A', 'Tremor.Subscore.UPDRS3A',
   'Brady.Rigidity.Subscore.UPDRS3A', 'PIGD.Subscore.UPDR30N',
→ 'Tremor.Subscore.UPDR30N', 'Brady.Rigidity.Subscore.UPDR30N',
→ 'PIGD.Subscore.UPDR30F', 'Tremor.Subscore.UPDR30F',
→ 'Brady.Rigidity.Subscore.UPDR30F'], '.Num') # Add a .Num extension to column names w

    updrs numeric vars

### UPDRS CATEGORICAL ####
# UPDRS3 (four dataframes) decode and rename in loop
updrs3_df_list = [nupdrs3_copy, nupdrs3A_copy, nupdr30N_copy, nupdr30F_copy]
ext_list = ['.UPDRS3', '.UPDRS3A', '.UPDR3ON', '.UPDR3OF']
```

```
for df in [nupdrs3_copy, nupdrs3A_copy, nupdr30N_copy, nupdr30F_copy] :
        df = decode_none2severe(df, ['NP3SPCH', 'NP3FACXP', 'NP3RIGN', 'NP3RIGRU',
        'NP3RIGLU', 'NP3RIGRL', 'NP3RIGLL', 'NP3FTAPR', 'NP3FTAPL', 'NP3HMOVR', 'NP3HMOVL',
        'NP3PRSPR', 'NP3PRSPL', 'NP3TTAPR', 'NP3TTAPL', 'NP3LGAGR', 'NP3LGAGL', 'NP3RISNG',
       'NP3GAIT', 'NP3FRZGT', 'NP3PSTBL', 'NP3POSTR', 'NP3BRADY', 'NP3PTRMR', 'NP3PTRML',
→ 'NP3KTRMR', 'NP3KTRML', 'NP3RTARU', 'NP3RTALU', 'NP3RTALL', 'NP
→ 'NP3RTCON'])
        df['DBS_STATUS'].replace({0 : 'OFF', 1 : 'ON'}, inplace = True)
        df = decode_0_1_no_yes(df, ['DYSKPRES','DYSKIRAT'])
        df['NHY'].replace({0 : 'Asymptomatic', 1 : 'Unilateral Movement Only', 2: 'Bilateral
       involvement without impairment of balance', 3 : 'Mild to moderate involvement',4:
       'Severe disability', 5 : 'Wheelchair bound or bedridden'}, inplace = True)
        df['PDTRTMNT'].replace({0 : False , 1: True}, inplace = True)
        df.rename(columns = {'DBS_STATUS' : 'Deep.Brain.Stimulation.Treatment' , 'NP3SPCH' :
        'UPDRS3.Speech.Difficulty', 'NP3FACXP' : 'Facial.Expression.Difficulty', 'NP3RIGN' :
       'Rigidity.Neck', 'NP3RIGRU': 'Rigidity.RUE', 'NP3RIGLU': 'Rigidity.LUE',
        'NP3RIGRL' : 'Rigidity.RLE', 'NP3RIGLL' : 'Rigidity.LLE', 'NP3FTAPR' :
        'Finger.Tapping.Right.Hand' ,'NP3FTAPL' : 'Finger.Tapping.Left.Hand' ,'NP3HMOVR' :
        'Hand.Movements.Right.Hand', 'NP3HMOVL' : 'Hand.Movements.Left.Hand', 'NP3PRSPR' :
       'Pronation.Supination.Right.Hand', 'NP3PRSPL' : 'Pronation.Supination.Left.Hand',
       'NP3TTAPR' : 'Toe.Tapping.Right.Foot', 'NP3TTAPL' : 'Toe.Tapping.Left.Foot',
        'NP3LGAGR' : 'Leg.Agility.Right.Leg', 'NP3LGAGL' : 'Leg.Agility.Left.Leg', 'NP3RISNG'
       : 'UPDRS3.Rising.from.Chair', 'NP3GAIT' : 'Gait.Problems', 'NP3FRZGT' :
       'Freezing.of.Gait', 'NP3PSTBL': 'Postural.Stability.Problems', 'NP3POSTR':
        'Posture.Problems' , 'NP3TOT':'UPDRS.Part3.Total', 'RMONOFF' :
        'MDS-UPDRS.Part3.Remote.Visit.ON.or.OFF',
        'Most.Recent.PD.Med.Dose.Date.Time.Before.OFF.Exam' :
       'Most.Recent.PD.Med.Dose.Date.Time.Before.OFF.Exam', 'RMTOFFRSN' :
        'Reason.OFF.Exam.at.Remote.Visit', 'ONEXAM': 'ON.Exam.Performed', 'ONEXAMTM':
        'ON.Exam.Time', 'ONNORSN': 'Reason.ON.Exam.Not.Performed', 'ONOFFORDER':
        'ON.or.OFF.Exam.Performed.First', 'Most.Recent.PD.Med.Dose.Date.Time.Before.ON.Exam'
       :'Most.Recent.PD.Med.Dose.Date.Time.Before.ON.Exam', 'PDMEDYN' : 'On.PD.Medication',
        'DBSONTM' : 'Time.DBS.Turned.on.before.ON.Exam', 'DBSOFFTM' :
        'Time.DBS.Turned.off.before.OFF.Exam', 'OFFNORSN': 'Reason.OFF.Exam.Not.Performed',
       'OFFEXAM' : 'OFF.Exam.Performed', 'OFFEXAMTM' : 'OFF.Exam.Time', 'DBSYN' : 'Has.DBS',
     'HRPOSTMED' : 'Hours.btwn.PD.Med.and.UPDRS3.Exam', 'HRDBSOFF' :
→ 'Hours.btwn.DBS.Device.Off.and.UPDRS3.Exam', 'HRDBSON' :
        'Hours.btwn.DBS.Device.On.and.UPDRS3.Exam' , 'RMEXAM' :
       'Remote.UPDRS3.Exam', 'DYSKPRES' : 'Dyskinesias.Present', 'DYSKIRAT' :
       'Movements.Interefered.with.Ratings', 'NHY': 'Hoehn.and.Yahr.Stage', 'PDTRTMNT':
\hookrightarrow
        'On.PD.Treatment', 'NP3BRADY' : 'Global.Spontaneity.of.Movement', 'NP3PTRMR' :
        'Postural.Tremor.Right.Hand' , 'NP3PTRML' : 'Postural.Tremor.Left.Hand' , 'NP3KTRMR'
     : 'Kinetic.Tremor.Right.Hand', 'NP3KTRML' : 'Kinetic.Tremor.Left.Hand', 'NP3RTARU' :
\label{eq:continuous} \mbox{$\hookrightarrow$} \quad \mbox{'Rest.Tremor.Amplitude.LUE', 'NP3RTARL' : 'Rest.Tremor.Amplitude.LUE', 'Rest.Tremor.Amplitud
     'Rest.Tremor.Amplitude.RLE' ,'NP3RTALL' : 'Rest.Tremor.Amplitude.LLE' ,'NP3RTALJ' :
       'Rest.Tremor.Amplitude.Lip.Jaw', 'NP3RTCON' : 'Constancy.of.Rest.Tremor', 'PDSTATE' :
      'Functional.State', 'EXAMTM' : 'UPDRS.Part3.Exam.Time' }, inplace = True)
        df = add_extension_to_column_names(df, ['PATNO', 'EVENT_ID', 'INFODT'],
       ext_list[count])
        count += 1
# Combine pd med dose date and time into one column
```

```
nupdrs3 copy = merge columns(nupdrs3 copy, ['PDMEDDT.UPDRS3', 'PDMEDTM.UPDRS3'],
→ 'Most.Recent.PD.Med.Dose.Date.Time.UPDRS3',' ')
nupdrs3A_copy = merge_columns(nupdrs3A_copy, ['PDMEDDT.UPDRS3A', 'PDMEDTM.UPDRS3A'],
→ 'Most.Recent.PD.Med.Dose.Date.Time.UPDRS3A', '')
nupdr30N_copy = merge_columns(nupdr30N_copy, ['PDMEDDT.UPDR30N', 'PDMEDTM.UPDR30N'],
→ 'Most.Recent.PD.Med.Dose.Date.Time.UPDR30N', ' ')
nupdr30F copy = merge columns(nupdr30F copy, ['PDMEDDT.UPDR30F', 'PDMEDTM.UPDR30F'],
→ 'Most.Recent.PD.Med.Dose.Date.Time.UPDR30F', '')
# Merge NUPDRS3, NUPDRS3A, NUPDR30N, NUPDR30F
updrs3_merge = pd.merge(nupdrs3_copy, nupdrs3A_copy, on = ['PATNO', 'EVENT_ID'], how =
→ "outer")
updrs3_merge = pd.merge(updrs3_merge, nupdr30F_copy, on = ['PATNO', 'EVENT_ID'], how =
→ "outer")
updrs3_merge = pd.merge(updrs3_merge, nupdr30N_copy, on = ['PATNO', 'EVENT_ID'], how =
→ "outer")
# Rename page name variables
updrs3_merge['UPDRS.Part3.Page.Name.UPDRS3'].replace({'NUPDRS3' : 'MDS-UPDRS Part III (No

¬ Treatment)'}, inplace = True)

updrs3_merge['UPDRS.Part3.Page.Name.UPDRS3A'].replace({'NUPDRS3A' : 'MDS-UPDRS Part III
updrs3_merge['UPDRS.Part3.Page.Name.UPDR30F'].replace({'NUPDR30F' : 'MDS-UPDRS Part III
updrs3 merge['UPDRS.Part3.Page.Name.UPDR30N'].replace({'NUPDR30N' : 'MDS-UPDRS Part III
# Merge all UPDRS dfs together
updrs cat = pd.merge(updrs part1 df copy, updrs part1 pq df copy, on = ['PATNO',

    'EVENT_ID'], how = "outer")

updrs_cat = pd.merge(updrs_cat, updrs_part2_pq_df_copy, on = ['PATNO', 'EVENT_ID'], how =
→ "outer")
updrs_cat = pd.merge(updrs_cat, updrs3_merge, on = ['PATNO', 'EVENT_ID'], how = "outer")
updrs_cat = pd.merge(updrs_cat, updrs_part3_dos_df_copy, on = ['PATNO', 'EVENT_ID'], how
updrs_cat = pd.merge(updrs_cat, updrs_part4_motor_df_copy, on = ['PATNO', 'EVENT_ID'],
→ how = "outer")
## Decode UPDRS df variables
# MNDS_UPDRS Part 1
updrs cat['UPDRS.Part1.Source'].replace({1 : 'Patient' , 2 : 'Caregiver' , 3 : 'Patient

    and Caregiver'}, inplace = True)

updrs_cat = decode_none2severe(updrs_cat, ['NP1COG', 'NP1HALL', 'NP1DPRS', 'NP1ANXS',
→ 'NP1APAT', 'NP1DDS'])
updrs_cat.rename(columns = {'UPDRS.Part1.Source' : 'UPDRS.Part1.Source.UPDRS1', 'NP1COG'
→ : 'Cognitive.Impairment.UPDRS1', 'NP1HALL' : 'Hallucinations.and.Psychosis.UPDRS1',
→ 'NP1DPRS': 'Depressed.Moods.UPDRS1', 'NP1ANXS': 'Anxious.Moods.UPDRS1', 'NP1APAT':
→ 'Apathy.UPDRS1', 'NP1DDS': 'Features.of.Dopamine.Dysregulation.Syndrome.UPDRS1',
→ 'NP1RTOT' : 'UPDRS.Part1.Rater.Completed.Total.UPDRS1'}, inplace = True)
# MDS_UPDRS_Part 1 Patient Questionnaire
updrs_cat['UPDRS.Part1.PQ.Source'].replace({1 : 'Patient', 2 : 'Caregiver', 3: 'Patient

    and Caregiver'}, inplace = True)
```

```
updsr_cat = decode_none2severe(updrs_cat, ['NP1SLPN', 'NP1SLPD', 'NP1PAIN', 'NP1URIN',
→ 'NP1CNST', 'NP1LTHD', 'NP1FATG'])
updrs_cat.rename(columns = {'UPDRS.Part1.PQ.Source' :
→ 'UPDRS.Part1.Patient.Questionnaire.Source.UPDRS1','NP1SLPN' :
    'Sleep.Problems.Night.UPDRS1', 'NP1SLPD' : 'Daytime.Sleepiness.UPDRS1', 'NP1PAIN' :
   'Pain.UPDRS1', 'NP1URIN': 'Urinary.Problems.UPDRS1', 'NP1CNST':
→ 'Constipation.Problems.UPDRS1', 'NP1LTHD': 'Lightheadedness.on.Standing.UPDRS1',
  'NP1FATG' : 'Fatigue.UPDRS1', 'NP1PTOT' :
  'UPDRS.Part1.Patient.Completed.Total.UPDRS1'}, inplace = True)
# MDS_UPDRS Part 2
updrs_cat['UPDRS.Part2.Source'].replace({1 : 'Patient', 2 : 'Caregiver', 3 : 'Patient and
updrs_cat =
decode_none2severe(updrs_cat,['NP2SPCH','NP2SALV','NP2SWAL','NP2EAT','NP2DRES','NP2HYGN','NP2HWRT',
updrs_cat.rename(columns = {'UPDRS.Part2.Source' : 'UPDRS.Part2.Source.UPDRS2', 'NP2SPCH'
- : 'UPDRS2.Speech.Difficulty.UPDRS2' , 'NP2SALV' : 'Saliva.Drooling.UPDRS2'
  ,'NP2SWAL': 'Chewing.Swallowing.Difficulty.UPDRS2', 'NP2EAT':
   'Eating.Difficulty.UPDRS2', 'NP2DRES': 'Dressing.Difficulty.UPDRS2', 'NP2HYGN':
   'Hygiene.Difficulty.UPDRS2' , 'NP2HWRT' : 'Handwriting.Difficulty.UPDRS2' ,'NP2HOBB'
  : 'Hobbies.Difficulty.UPDRS2', 'NP2TURN': 'Turning.in.Bed.Difficulty.UPDRS2',
→ 'NP2TRMR' : 'Tremor.UPDRS2', 'NP2RISE' :
→ 'UPDRS2.Rising.from.Chair.Difficulty.UPDRS2', 'NP2WALK' : 'Walking.Difficulty.UPDRS2'
    , \mbox{'NP2FREZ'} : 'Freezing.while.Walking.UPDRS2' , 'NP2PTOT':
  'UPDRS.Part2.Total.UPDRS2'}, inplace = True)
# MDS_UPDRS Part 3 On OFF determination Dosing
updrs_cat = decode_0_1_no_yes(updrs_cat, ['RMEXAM', 'DBSYN', 'OFFEXAM', 'OFFEXAM',
→ 'ONEXAM'])
updrs_cat['RMTOFFRSN'].replace({1 : 'ON state not reached', 2 : 'Scheduling issues', 3 :
updrs_cat['ONOFFORDER'].replace({1 : 'OFF', 2 : 'ON'}, inplace = True)
updrs_cat['RMONOFF'].replace({1 : 'OFF', 2 : 'ON'}, inplace = True)
updrs_cat['ONNORSN'].replace({1 : 'ON state not reached', 2 : 'Scheduling issues', 3 :
→ 'Other reason'}, inplace = True)
updrs_cat['PDMEDYN' ].replace({0 : False , 1 : True}, inplace = True)
updrs_cat = merge_columns(updrs_cat, ['ONPDMEDDT', 'ONPDMEDTM'],
→ 'Most.Recent.PD.Med.Dose.Date.Time.Before.ON.Exam', '')
updrs_cat['OFFNORSN'].replace({1 : 'Disease severity preventing turning off of DBS', 2 :
\hookrightarrow 'Did not bring medication to turn on', 3: 'Forgot to refrain from taking
→ medication', 4: 'Does not want to turn off DBS', 5: 'Forgot to turn off DBS', 6:
   'Forgot to bring DBS', 7: 'Unsure if participant was full off', 8: 'Site scheduling

    issues', 9 : 'Other reason'}, inplace = True)

updrs_cat = merge_columns(updrs_cat, ['OFFPDMEDDT','OFFPDMEDTM'],
→ 'Most.Recent.PD.Med.Dose.Date.Time.Before.OFF.Exam', ' ')
updrs_cat.rename(columns = {'RMONOFF' :
   'MDS-UPDRS.Part3.Remote.Visit.ON.or.OFF.UPDRDOSE',
   'Most.Recent.PD.Med.Dose.Date.Time.Before.OFF.Exam':
   'Most.Recent.PD.Med.Dose.Date.Time.Before.OFF.Exam.UPDRDOSE', 'RMTOFFRSN':
    'Reason.OFF.Exam.at.Remote.Visit.UPDRDOSE', 'ONEXAM' : 'ON.Exam.Performed.UPDRDOSE'
  ,'ONEXAMTM' : 'ON.Exam.Time.UPDRDOSE', 'ONNORSN' :
   'Reason.ON.Exam.Not.Performed.UPDRDOSE', 'ONOFFORDER':
  'ON.or.OFF.Exam.Performed.First.UPDRDOSE',
  'Most.Recent.PD.Med.Dose.Date.Time.Before.ON.Exam'

→ :'Most.Recent.PD.Med.Dose.Date.Time.Before.ON.Exam.UPDRDOSE', 'PDMEDYN':

\rightarrow 'On.PD.Medication.UPDRDOSE', 'DBSONTM' :_{20}
→ 'UPDRS3.Time.DBS.Turned.on.before.ON.Exam.UPDRDOSE', 'DBSOFFTM' :
   'UPDRS3.Time.DBS.Turned.off.before.OFF.Exam.UPDRDOSE', 'DBSONTM y':
   'UPDRS3.Dos.Time.DBS.Turned.on.before.ON.Exam.UPDRDOSE', 'DBSOFFTM y':
```

'UPDRS3 Dos Time DRS Turned off before OFF Exam UPDRDOSE' 'OFFNORSN' .

```
# MDS_UPDRS Part 4
updrs_cat['NP4WDYSK'].replace({0 : 'No dyskinesias', 1: 'Slight: <= 25% of waking day',
\rightarrow 2: 'Mild: 26-50% of waking day', 3: 'Moderate: 51-75% of waking day', 4: 'Severe: >
→ 75% of waking day'}, inplace = True)
updrs_cat['NP4DYSKI'].replace({0 : 'No dyskinesias', 1 : 'Slight', 2: 'Mild', 3 :
→ 'Moderate', 4: 'Severe'}, inplace = True)
updrs_cat['NP40FF'].replace({0 : 'Normal: No OFF time', 1 : 'Slight: <= 25% of waking day
\rightarrow ', 2: 'Mild : 26-50% of waking day', 3 : 'Moderate: 51-75% of waking day', 4 :
updrs_cat['NP4FLCTI'].replace({0 : 'Normal', 1: 'Slight', 2: 'Mild', 3: 'Moderate', 4:
updrs_cat['NP4DYSTN'].replace({0 : 'No dystonia', 1: 'Slight: <= 25% of time in OFF
→ state', 2: 'Mild: 26-50% of time in OFF state', 3: 'Moderate: 51-75% of time in OFF

    state', 4: 'Severe: > 75% of time in OFF state'}, inplace = True)

updrs_cat['NP4FLCTX'].replace({0: 'Normal', 1: 'Slight', 2: 'Mild', 3: 'Moderate', 4:
updrs_cat['RMNOPRT3'].replace({1 : 'Visit was not conducted with video', 2 : 'Scheduling

    issues', 3: 'Other reason'}, inplace = True)

updrs_cat.rename(columns = {'RMNOPRT3':
→ 'Reason.UPDRSPart3.Not.Administered.Remote.Visit.UPDRDOSE','NP4WDYSKDEN':'4.1.Time.with.Dyskinesias
→ 'NP4WDYSKNUM' : '4.1.Total.Hours.Awake.UPDRS4', 'NP4WDYSKPCT' :
→ 'Percent.Dyskinesia.UPDRS4', 'NP40FFDEN': '4.3.Total.Hours.OFF.UPDRS4', 'NP40FFNUM':
   '4.3.Total.Hours.Awake.UPDRS4','NP4OFFPCT' : 'Percent.OFF.UPDRS4', 'NP4DYSTNDEN'
→ :'4.6.Total.Hours.OFF.with.Dystonia.UPDRS4',
→ 'NP4DYSTNNUM':'4.6.Total.Hours.OFF.UPDRS4',
   'NP4DYSTNPCT':'4.6.Percent.OFF.Dystonia.UPDRS4', 'NP4WDYSK':
   'Time.Spent.with.Dyskinesias.UPDRS4',
→ 'NP4DYSKI': 'Functional.Impact.of.Dyskinesias.UPDRS4', 'NP4OFF':
→ 'Time.Spent.in.OFF.State.UPDRS4',
→ 'NP4FLCTI': 'Functional.Impact.Fluctuations.UPDRS4',
   'NP4FLCTX':'Complexity.of.Motor.Fluctuations.UPDRS4'
→ 'NP4DYSTN': 'Painful.OFF-state.Dystonia.UPDRS4' , 'NP4TOT':
→ 'UPDRS.Part4.Total.UPDRS4', 'NP4WDYSKDEN' : '4.1.Total.Hours.with.Dyskinesia',
    'NP4WDYSKNUM' : '4.1. Total. Hours. Awake', 'NP4WDYSKPCT' : '4.1. %. Dyskinesia',
→ 'NP40FFDEN': '4.3.Total.Hours.OFF', 'NP40FFNUM': '4.3.Total.Hours.Awake'
→ 'NP4OFFPCT': '4.3.%.OFF', 'NP4DYSTNDEN' : '4.6.Total.Hours.OFF.with.Dystonia',
→ 'NP4DYSTNNUM' :'4.6.Total.Hours.OFF', 'NP4DYSTNPCT' : '4.6.%.OFF.Dystonia'
→ ,'NP4DYSTNPCT' :'4.6.%.OFF.Dystonia'}, inplace = True)
updrs_cat = add_extension_to_column_names(updrs_cat, ['PATNO', 'EVENT_ID', 'INFODT'],
- '.Cat') # Add a .Cat extension to column names w updrs cat vars
# Create one df with UPDRS scores .Num (numeric) and all UPDRS scores .Cat (categorical)
updrs_cat.drop(['INFODT'], axis = 1, inplace = True)
updrs_merged = pd.merge(updrs_cat, updrs_numeric, on = ['PATNO', 'EVENT_ID'], how =
updrs_merged.replace({'UR' : np.nan}, inplace = True) # Unable to Rate --> nan
ppmi_merge = pd.merge(ppmi_merge, updrs_merged, on = ['PATNO', 'EVENT_ID'], how =
→ "outer")
#### CLEAN UP DF ####
ppmi_merge.drop(['Subid.Date.TEMP'], axis = 1, inplace = True)
ppmi_merge.rename(columns = {'EVENT_ID': 'Event.ID', 'INFODT' :
'Event.ID.Date','Medication': 'Medication(Dates)', 'DOMSIDE':
→ 'Dominant.Side.Disease'}, inplace = True)
```

```
# Change names of event ids to be indicative of months
ppmi_merge['Event.ID'].replace({'FNL' : 'Final Visit', 'BL' : 'Baseline', 'SC' :
→ 'Screening', 'LOG' : 'Logs', 'PW' : 'Premature Withdrawal', 'R04' : 'Remote Visit
_{\hookrightarrow} Month 18', 'R06' : 'Remote Visit Month 18', 'R08' : 'Remote Visit Month 42', 'R10' :
\hookrightarrow 'Remote Visit Month 54', 'R12' : 'Remote Visit Month 66', 'R13' : 'Remote Visit Month
_{\rightarrow} 78' , 'R14' : 'Remote Visit Month 90' , 'R15' : 'Remote Visit Month 102' , 'R16' :
→ 'Remote Visit Month 114', 'R17' : 'Remote Visit Month 126', 'RS1' : 'Re-screen', 'ST'
\hookrightarrow : 'Symptomatic Therapy', 'U01' : 'Unscheduled Visit 1', 'U02' : 'Unscheduled Visit
→ 2', 'V01' : 'Visit Month 3', 'V02' : 'Visit Month 6', 'V03' : 'Visit Month 9', 'V04'
\hookrightarrow : 'Visit Month 12', 'V05' : 'Visit Month 18', 'V06' : 'Visit Month 24', 'V07' :
\hookrightarrow 'Visit Month 30', 'V08' : 'Visit Month 36', 'V09' : 'Visit Month 42', 'V10' : 'Visit
\hookrightarrow Month 48', 'V11' : 'Visit Month 54', 'V12' : 'Visit Month 60', 'V13' : 'Visit Month
→ 72', 'V14': 'Visit Month 84', 'V15': 'Visit Month 96', 'V16': 'Visit Month 108',
_{\rm \hookrightarrow} 'V17' : 'Visit Month 120', 'V18' : 'Visit Month 132', 'P78' : 'Phone Visit (Month
\rightarrow 78)'}, inplace = True)
# Fill in cells that are NA with subject information that we know from other event
→ ids/rows for fixed variables
fixed_var_list = ['Enroll.Diagnosis', 'Enroll.Subtype','Consensus.Diagnosis',
→ 'Consensus.Subtype', 'Subject.Phenoconverted', 'BirthDate', 'Sex', 'Handed',
→ 'Analytic.Cohort'] # fixed variables
for col_name in fixed_var_list :
    ppmi_merge[col_name].fillna('NA', inplace = True)
    for row_num in range(len(ppmi_merge[col_name])):
        if ppmi_merge[col_name].iloc[row_num] == 'NA' : # if any entry is NA
            current_sub = ppmi_merge['PATNO'].iloc[row_num] # get current subid
            fixed_var_value = ppmi_merge.loc[(ppmi_merge['PATNO'] == current_sub ) &
    (ppmi merge[col name] != 'NA'), col name].values # get value from another event id
            if fixed_var_value.any() :
                ppmi_merge.loc[row_num, col_name] = fixed_var_value[0] # fill in baseline
\hookrightarrow value at NA
ppmi_merge = ppmi_merge.rename(columns = {'PATNO' : 'Subject.ID'})
ppmi_merge.replace({'NA' : np.nan}, inplace = True)
#### GENETICS INFO ####
lrrk2_genetics_df = pd.read_csv(genetics_path + 'lrrk2_geno_012_mac5_missing_geno.csv')
scna_genetics_df = pd.read_csv(genetics_path + 'scna_geno_012_mac5_missing_geno.csv')
apoe_genetics_df = pd.read_csv(genetics_path + 'apoe_geno_012_mac5_missing_geno.csv')
tmem_genetics_df = pd.read_csv(genetics_path + 'tmem175_geno_012_mac5_missing_geno.csv')
gba genetics df = pd.read csv(genetics path + 'gba geno 012 mac5 missing geno.csv')
def format_genetics_df(genetics_df : pd.DataFrame ) :
    Format genetics_df to make merge-able with ppmi_merge
    genetics_df.drop(['COUNTED', 'ALT', 'SNP', '(C)M'], axis = 1, inplace = True) #
→ Remove unnecessary columns
    # Change column names to be just subid
    for col in genetics_df:
        if '_' in col :
            subid = int(col.split('_')[-1])
```

```
genetics_df.rename(columns = {col : subid}, inplace = True)
   # Combine CHR and POS columns
   genetics_df['CHR'] = 'CHR' + genetics_df['CHR'].astype(str) # Need to be strings
→ before you use merge_columns function
   genetics_df['POS'] = 'POS' + genetics_df['POS'].astype(str) # Need to be strings
→ before you use merge_columns function
   genetics_df = merge_columns(genetics_df, ['CHR', 'POS'], 'Chromosome.Position', '.')
   # Pivot df so position is column name and subid is row
   genetics_df = genetics_df.T # Transpose df so that rows are subid
   genetics df.rename(columns = genetics df.iloc[-1], inplace = True) # Move Chr.Pos to
genetics_df.index.names = ['Subject.ID'] # Rename index to 'Subject.ID'
   genetics_df = genetics_df.drop(['Chromosome.Position'], axis = 0) # Drop last row
genetics_df = genetics_df.reset_index(drop = False)
   return genetics_df
lrrk2_genetics_df_formatted = format_genetics_df(lrrk2_genetics_df)
scna_genetics_df_formatted = format_genetics_df(scna_genetics_df)
apoe_genetics_df_formatted = format_genetics_df(apoe_genetics_df)
tmem_genetics_df_formatted = format_genetics_df(tmem_genetics_df)
gba_genetics_df_formatted = format_genetics_df(gba_genetics_df)
# Merge genetics dataframes together to create one genetics_df
genetics_df = pd.merge(lrrk2_genetics_df_formatted, scna_genetics_df_formatted, on =
genetics_df = pd.merge(genetics_df, apoe_genetics_df_formatted, on = ['Subject.ID'], how
genetics_df = pd.merge(genetics_df, tmem_genetics_df_formatted, on = ['Subject.ID'], how
genetics_df = pd.merge(genetics_df, gba_genetics_df_formatted, on = ['Subject.ID'], how =
→ "outer")
# Change genetics col names int to float (remove .0 in all 'CHR.POS' columns)
for col_name in genetics_df :
   if col name.startswith('CHR'):
       genetics_df[col_name] = genetics_df[col_name].fillna(-9999.0)
       genetics_df[col_name] = genetics_df[col_name].astype(int)
genetics_df.replace({-9999.0 : 'NA'}, inplace = True)
# Merge ppmi_merge with genetics df
ppmi_merge_genetics = pd.merge(ppmi_merge, genetics_df, on = 'Subject.ID', how = "outer")
#### T1 Info - Taylor's File ####
ppmi_t1_df = pd.read_csv(invicro_data_path + 'ppmi_mergewide_t1.csv') # Read in Taylor's
→ T1 results file
ppmi_t1_df.rename(columns = {'u_hier_id_OR': 'Subject.ID'}, inplace = True) # Rename
→ subject id column in Taylors of to match ppmi_merge
# Create a column for object name to merge on with ppmi_merge
```

```
ppmi_t1_df['Image_ID_merge'] = ''
for row_num in range(len(ppmi_t1_df['ImageID'])) :
   image_id = ppmi_t1_df['ImageID'].iloc[row_num].split('-')[2]
   ppmi_t1_df['Image_ID_merge'].iloc[row_num] = image_id
# Merge ppmi_merge_genetics with t1 info
ppmi merge = pd.merge(ppmi merge genetics, ppmi t1 df, on =
ppmi_merge.drop(['Image_ID_merge'], axis = 1, inplace = True) # Drop
# Put full date in Image.Acquisition.Date column
for row num in range(len(ppmi merge['T1.s3.Image.Name'])) :
   if isinstance(ppmi_merge['T1.s3.Image.Name'].iloc[row_num],str) :
       date = ppmi_merge['T1.s3.Image.Name'].iloc[row_num].split('-')[2]
       ppmi_merge['Image.Acquisition.Date'].iloc[row_num] = date[4:6] + '/' + date[6:8]
  +'/' + date[0:4]
## Get Enrollment Diagnosis for subjects in Not Analytic Cohort - do this using the

→ participants_status.csv

analytic = ppmi_merge[ppmi_merge['Analytic.Cohort'] == 'Analytic Cohort'] # Split up
→ Analytic cohort df and not Analytic Cohort df
not_analytic = ppmi_merge[ppmi_merge['Analytic.Cohort'] == 'Not Analytic Cohort'] # Split
→ up Analytic cohort df and not Analytic Cohort df
participant status = pd.read csv(ppmi download path + 'Participant Status.csv') # Read in
→ participant status.csv
participant status = participant status[['PATNO', 'COHORT DEFINITION']] # Keep only
participant_status.rename(columns = {'PATNO' : 'Subject.ID', 'COHORT_DEFINITION' :
not analytic participant status = pd.merge(not analytic, participant status, on =
→ ['Subject.ID'], how = "left") # Merge not Atlantic subids with enrollment diagnosis
not_analytic_participant_status.drop(['Enroll.Diagnosis_x'], axis = 1, inplace = True) #
→ Remove the extra Enroll.Diagnosis created at merge
not_analytic_participant_status.rename(columns = {'Enroll.Diagnosis_y' :
not_analytic_participant_status.loc[not_analytic_participant_status['Enroll.Diagnosis']
→ == 'Healthy Control', 'Enroll.Subtype'] = 'Healthy Control' # For Healthy Control
→ subjects in the Not Analytic Cohort - make 'Enroll.Subtype' = Healthy Control
# Merge df of Not Analytic and Analytic subjects and sort by SubID and Event.ID.Date
ppmi merge = pd.concat([analytic, not analytic participant status])
# Change Event. ID. Date to date time and corrected format
ppmi_merge['Event.ID.Date'] = ppmi_merge['Event.ID.Date'].astype(str)
ppmi_merge['Event.ID.Date'] = pd.to_datetime(ppmi_merge['Event.ID.Date'], errors =
→ "ignore") # Change event. ID. Date column to date time so we can sort according to this
ppmi_merge = ppmi_merge.sort_values(by = ['Subject.ID', 'Event.ID.Date']) # Sort values by
→ subject and event id date
ppmi_merge['Event.ID.Date'] = ppmi_merge['Event.ID.Date'].astype(str) # Change
→ Event.ID.Date back to string so we can reformat
# Reformat Event.ID.Date from pd.to_datetime to month/year
```

```
for row_num in range(len(ppmi_merge['Event.ID.Date'])):
    if ppmi_merge['Event.ID.Date'].iloc[row_num] != 'NaT':
       split = ppmi_merge['Event.ID.Date'].iloc[row_num].split('-')
       new_date = split[1] +'/' + split[0] # month/year format
       ppmi_merge['Event.ID.Date'].iloc[row_num] = new_date
ppmi_merge['Event.ID.Date'] = ppmi_merge['Event.ID.Date'].replace('NaT','NA')
#### Add in ppmi_qc_BA.csv - Brian sent on slack 1/31/22 ####
ppmi_qc_BA = pd.read_csv(invicro_data_path + 'ppmi_qc_BA.csv')
ppmi_qc_BA = ppmi_qc_BA.reset_index(drop = False) # Move index to first column so we can
ppmi_qc_BA.rename(columns = {'ID' : 'ImageID'}, inplace = True) # Rename ImageID
# Update ImageID column bc info from T1 file (where ImageID was created from) does not
→ contain all the files from s3 (need this to merge in subs from QC csv)
for row_num in range(len(ppmi_merge['ImageID'])) :
    if isinstance(ppmi_merge['T1.s3.Image.Name'].iloc[row_num], str):
       imageID = ppmi_merge['T1.s3.Image.Name'].iloc[row_num].split('.')[0] # take info
  before .nii.gz
       ppmi_merge['ImageID'].iloc[row_num] = imageID.split('-')[1] + '-' +
  imageID.split('-')[2] + '-' + imageID.split('-')[4]
ppmi_merge = pd.merge(ppmi_merge, ppmi_qc_BA, on = ['ImageID'], how = "left") # Merge -
→ keep only from ImageIDs we already have
#### BILATERAL SUBTYPE SCORES (Tremor and Brady) ####
brady_right3 = ['Dominant.Side.Disease', 'Rigidity.Neck.UPDRS3.Num',
→ 'Rigidity.RUE.UPDRS3.Num', 'Rigidity.RLE.UPDRS3.Num',
→ 'Finger.Tapping.Right.Hand.UPDRS3.Num', 'Hand.Movements.Right.Hand.UPDRS3.Num',
→ 'Pronation.Supination.Right.Hand.UPDRS3.Num', 'Toe.Tapping.Right.Foot.UPDRS3.Num',
→ 'Leg.Agility.Right.Leg.UPDRS3.Num']
brady_right3a = ['Dominant.Side.Disease', 'Rigidity.Neck.UPDRS3A.Num',
→ 'Rigidity.RUE.UPDRS3A.Num', 'Rigidity.RLE.UPDRS3A.Num',
→ 'Finger.Tapping.Right.Hand.UPDRS3A.Num', 'Hand.Movements.Right.Hand.UPDRS3A.Num',
  'Pronation.Supination.Right.Hand.UPDRS3A.Num', 'Toe.Tapping.Right.Foot.UPDRS3A.Num',
→ 'Leg.Agility.Right.Leg.UPDRS3A.Num']
brady_right30N = ['Dominant.Side.Disease', 'Rigidity.Neck.UPDR30N.Num',
→ 'Rigidity.RUE.UPDR3ON.Num', 'Rigidity.RLE.UPDR3ON.Num',
→ 'Finger.Tapping.Right.Hand.UPDR3ON.Num', 'Hand.Movements.Right.Hand.UPDR3ON.Num',
- 'Pronation.Supination.Right.Hand.UPDR3ON.Num', 'Toe.Tapping.Right.Foot.UPDR3ON.Num',
→ 'Leg.Agility.Right.Leg.UPDR30N.Num']
brady_right30F = ['Dominant.Side.Disease', 'Rigidity.Neck.UPDR30F.Num',
→ 'Rigidity.RUE.UPDR30F.Num', 'Rigidity.RLE.UPDR30F.Num',
→ 'Finger.Tapping.Right.Hand.UPDR3OF.Num', 'Hand.Movements.Right.Hand.UPDR3OF.Num',
→ 'Pronation.Supination.Right.Hand.UPDR3OF.Num', 'Toe.Tapping.Right.Foot.UPDR3OF.Num',
→ 'Leg.Agility.Right.Leg.UPDR30F.Num']
brady_left3 = ['Dominant.Side.Disease', 'Rigidity.Neck.UPDRS3.Num',
→ 'Rigidity.LUE.UPDRS3.Num', 'Rigidity.LLE.UPDRS3.Num',
→ 'Pronation.Supination.Left.Hand.UPDRS3.Num', 'Toe.Tapping.Left.Foot.UPDRS3.Num',
→ 'Leg.Agility.Left.Leg.UPDRS3.Num']
brady_left3a = ['Dominant.Side.Disease', 'Rigidity.Neck.UPDRS3A.Num',
→ 'Rigidity.LUE.UPDRS3A.Num', 'Rigidity.LLE.UPDRS3A.Num',
→ 'Finger.Tapping.Left.Hand.UPDRS3A.Num', 'Hand.Movements.Left.Hand.UPDRS3A.Num',
{\scriptstyle \leftarrow} \quad \text{'Pronation.Supination.Left.Hand.UPDRS3A.Num', 'Toe.Tapping.Left.Foot.UPDRS3A.Num', }
```

```
brady_left30N = ['Dominant.Side.Disease', 'Rigidity.Neck.UPDR30N.Num',
→ 'Rigidity.LUE.UPDR30N.Num', 'Rigidity.LLE.UPDR30N.Num',
\hookrightarrow
      'Finger.Tapping.Left.Hand.UPDR3ON.Num', 'Hand.Movements.Left.Hand.UPDR3ON.Num',
      'Pronation.Supination.Left.Hand.UPDR3ON.Num', 'Toe.Tapping.Left.Foot.UPDR3ON.Num',
    'Leg.Agility.Left.Leg.UPDR30N.Num']
brady_left30F = ['Dominant.Side.Disease', 'Rigidity.Neck.UPDR30F.Num',
→ 'Rigidity.LUE.UPDR30F.Num', 'Rigidity.LLE.UPDR30F.Num',
→ 'Finger.Tapping.Left.Hand.UPDR30F.Num', 'Hand.Movements.Left.Hand.UPDR30F.Num',
→ 'Pronation.Supination.Left.Hand.UPDR30F.Num', 'Toe.Tapping.Left.Foot.UPDR30F.Num',
→ 'Leg.Agility.Left.Leg.UPDR30F.Num']
brady sym3 = ['Dominant.Side.Disease', 'Rigidity.Neck.UPDRS3.Num',
→ 'Rigidity.RUE.UPDRS3.Num', 'Rigidity.LUE.UPDRS3.Num', 'Rigidity.RLE.UPDRS3.Num',
    'Rigidity.LLE.UPDRS3.Num', 'Finger.Tapping.Right.Hand.UPDRS3.Num',
→ 'Finger.Tapping.Left.Hand.UPDRS3.Num', 'Hand.Movements.Right.Hand.UPDRS3.Num',
→ 'Hand.Movements.Left.Hand.UPDRS3.Num', 'Pronation.Supination.Right.Hand.UPDRS3.Num',
→ 'Pronation.Supination.Left.Hand.UPDRS3.Num', 'Toe.Tapping.Right.Foot.UPDRS3.Num',
→ 'Toe.Tapping.Left.Foot.UPDRS3.Num', 'Leg.Agility.Right.Leg.UPDRS3.Num',

    'Leg.Agility.Left.Leg.UPDRS3.Num']

brady_sym3a = ['Dominant.Side.Disease', 'Rigidity.Neck.UPDRS3A.Num',
\label{eq:continuous} \mbox{$\overset{\cdot}{\to}$} \mbox{$\ '$Rigidity.RUE.UPDRS3A.Num', 'Rigidity.RLE.UPDRS3A.Num', 'Rigidi
→ 'Rigidity.LLE.UPDRS3A.Num', 'Finger.Tapping.Right.Hand.UPDRS3A.Num',
→ 'Finger.Tapping.Left.Hand.UPDRS3A.Num', 'Hand.Movements.Right.Hand.UPDRS3A.Num',
→ 'Hand.Movements.Left.Hand.UPDRS3A.Num'.
→ 'Pronation.Supination.Right.Hand.UPDRS3A.Num',
→ 'Pronation.Supination.Left.Hand.UPDRS3A.Num', 'Toe.Tapping.Right.Foot.UPDRS3A.Num',
→ 'Toe.Tapping.Left.Foot.UPDRS3A.Num', 'Leg.Agility.Right.Leg.UPDRS3A.Num',
    'Leg.Agility.Left.Leg.UPDRS3A.Num']
brady sym30N = ['Dominant.Side.Disease', 'Rigidity.Neck.UPDR30N.Num',
→ 'Rigidity.RUE.UPDR3ON.Num', 'Rigidity.LUE.UPDR3ON.Num', 'Rigidity.RLE.UPDR3ON.Num',
→ 'Rigidity.LLE.UPDR30N.Num', 'Finger.Tapping.Right.Hand.UPDR30N.Num',
→ 'Finger.Tapping.Left.Hand.UPDR3ON.Num', 'Hand.Movements.Right.Hand.UPDR3ON.Num',
→ 'Hand.Movements.Left.Hand.UPDR30N.Num',
→ 'Pronation.Supination.Right.Hand.UPDR3ON.Num',
→ 'Pronation.Supination.Left.Hand.UPDR3ON.Num', 'Toe.Tapping.Right.Foot.UPDR3ON.Num',
→ 'Toe.Tapping.Left.Foot.UPDR3ON.Num', 'Leg.Agility.Right.Leg.UPDR3ON.Num',

    'Leg.Agility.Left.Leg.UPDR30N.Num']

brady_sym30F = ['Dominant.Side.Disease', 'Rigidity.Neck.UPDR30F.Num',
→ 'Rigidity.RUE.UPDR30F.Num', 'Rigidity.LUE.UPDR30F.Num', 'Rigidity.RLE.UPDR30F.Num',
→ 'Rigidity.LLE.UPDR30F.Num', 'Finger.Tapping.Right.Hand.UPDR30F.Num',
→ 'Finger.Tapping.Left.Hand.UPDR30F.Num', 'Hand.Movements.Right.Hand.UPDR30F.Num',
→ 'Hand.Movements.Left.Hand.UPDR30F.Num',
→ 'Pronation.Supination.Right.Hand.UPDR30F.Num',
→ 'Pronation.Supination.Left.Hand.UPDR30F.Num', 'Toe.Tapping.Right.Foot.UPDR30F.Num',
→ 'Toe.Tapping.Left.Foot.UPDR30F.Num', 'Leg.Agility.Right.Leg.UPDR30F.Num',
→ 'Leg.Agility.Left.Leg.UPDR30F.Num']
# Make any dominant side of disease that are NA into 'Symmetric'
domsideisna = ppmi_merge['Dominant.Side.Disease'].isna()
ppmi_merge['Dominant.Side.Disease'].loc[domsideisna] = 'Symmetric'
def add_lateralized_subscores(df : pd.DataFrame, subscore_side_list : list, side : str,
→ new_col_name : str) :
```

```
Include lateralized subscroes (i.e. Brady Rigidity and Tremor subscores) into
\hookrightarrow dataframe.
    Arguments
    df: pd.DataFrame containing scores that make up subscore
    subscore_side_list : list containing column names of the scores that make up the
→ subscore
    side : 'Left' or 'Right'
    new_col_name : name of new column name with lateralized subscore
    11 11 11
    subscore_side = df[subscore_side_list] # Get dataframe of only columns in brady_left
    df[new_col_name] = 0 # Initialize lateralized variable
    subscore_side.loc[subscore_side['Dominant.Side.Disease'] != side, :] = np.nan # Make
\hookrightarrow all rows nan if dominant side of disease is not left
    subscore_side_temp = subscore_side.drop('Dominant.Side.Disease',1) # Drop dominant
→ side of disease - necessary because this cannot be summed in next line
    idx = subscore_side_temp.loc[pd.isnull(subscore_side_temp).any(1), :].index
   df[new_col_name] = subscore_side_temp.sum(axis = 1) # Sum of all columns in each row
\rightarrow where dom side is left
   df[new_col_name].iloc[idx] = np.nan # Fill in subscores that should be nans as nan
   return df
def add_symmetric_subscores(df : pd.DataFrame, subscore_side_list : list, side : str,
→ new_col_name : str, ext : str) :
    subscore_side = df[subscore_side_list] # Get dataframe of only columns in brady_left
    df[new_col_name] = 0 # Initialize lateralized variable
    subscore_side.loc[subscore_side['Dominant.Side.Disease'] != side, :] = np.nan # Make
\hookrightarrow all rows nan if dominant side of disease is not side
   subscore_side_temp = subscore_side.drop('Dominant.Side.Disease',1) # Drop dominant
→ side of disease - necessary because this cannot be summed in next line
   idx = subscore_side_temp.loc[pd.isnull(subscore_side_temp).any(1), :].index
   x = subscore_side_temp.fillna(0) # because adding 1 plus nan equals nan
    if "Brady" in new col name :
        df[new_col_name] = x['Rigidity.Neck' + ext] + (x['Rigidity.RUE' + ext] +
→ x['Rigidity.LUE'+ ext])/2 + (x['Rigidity.RLE'+ ext] + x['Rigidity.LLE'+ ext])/2 +
→ (x['Finger.Tapping.Right.Hand'+ ext] +x['Finger.Tapping.Left.Hand'+ ext])/2 +
→ (x['Hand.Movements.Right.Hand'+ ext] + x['Hand.Movements.Left.Hand'+ ext])/2 +
→ (x['Pronation.Supination.Right.Hand'+ ext]+x['Pronation.Supination.Left.Hand'+
ext])/2 + (x['Toe.Tapping.Right.Foot'+ ext]+ x['Toe.Tapping.Left.Foot'+ ext])/2 +

→ (x['Leg.Agility.Right.Leg'+ ext] + x['Leg.Agility.Left.Leg'+ ext])/2

        df[new_col_name].iloc[idx] = np.nan # Fill in subscores that should be nans as
    elif "Tremor" in new_col_name :
        df[new_col_name] = x['Tremor.UPDRS2.Num'] + (x['Postural.Tremor.Right.Hand' +
→ ext ] + x['Postural.Tremor.Left.Hand' + ext ])/2 + (x['Kinetic.Tremor.Right.Hand' +
→ ext ] + x['Kinetic.Tremor.Left.Hand'+ ext ])/2 + (x['Rest.Tremor.Amplitude.RUE' + ext
→ ]+ x['Rest.Tremor.Amplitude.LUE' + ext])/2 + (x['Rest.Tremor.Amplitude.RLE' + ext] +
\rightarrow x['Rest.Tremor.Amplitude.LLE' + ext])/2 \frac{1}{2}x['Rest.Tremor.Amplitude.Lip.Jaw' + ext] +

    x['Constancy.of.Rest.Tremor' + ext]
```

```
df[new_col_name].iloc[idx] = np.nan
    return df
ppmi_merge = add_lateralized_subscores(ppmi_merge, brady_left3, 'Left',
→ 'Brady.Rigidity.Subscore-left.UPDRS3')
ppmi_merge = add_lateralized_subscores(ppmi_merge, brady_left3a, 'Left',
→ 'Brady.Rigidity.Subscore-left.UPDRS3A')
ppmi_merge = add_lateralized_subscores(ppmi_merge, brady_left30N, 'Left',
→ 'Brady.Rigidity.Subscore-left.UPDR30N')
ppmi_merge = add_lateralized_subscores(ppmi_merge, brady_left30F, 'Left',
→ 'Brady.Rigidity.Subscore-left.UPDR30F')
ppmi_merge = add_lateralized_subscores(ppmi_merge, brady_right3, 'Right',
→ 'Brady.Rigidity.Subscore-right.UPDRS3')
ppmi_merge = add_lateralized_subscores(ppmi_merge, brady_right3a, 'Right',
→ 'Brady.Rigidity.Subscore-right.UPDRS3A')
ppmi_merge = add_lateralized_subscores(ppmi_merge, brady_right30N, 'Right',
→ 'Brady.Rigidity.Subscore-right.UPDR30N')
ppmi_merge = add_lateralized_subscores(ppmi_merge, brady_right30F, 'Right',
→ 'Brady.Rigidity.Subscore-right.UPDR30F')
ppmi_merge = add_symmetric_subscores(ppmi_merge, brady_sym3, 'Symmetric',
→ 'Brady.Rigidity.Subscore-sym.UPDRS3', '.UPDRS3.Num')
ppmi_merge = add_symmetric_subscores(ppmi_merge, brady_sym3a, 'Symmetric',
→ 'Brady.Rigidity.Subscore-sym.UPDRS3A', '.UPDRS3A.Num')
ppmi_merge = add_symmetric_subscores(ppmi_merge, brady_sym30N, 'Symmetric',
→ 'Brady.Rigidity.Subscore-sym.UPDR30N' , '.UPDR30N.Num')
ppmi_merge = add_symmetric_subscores(ppmi_merge, brady_sym30F, 'Symmetric',
→ 'Brady.Rigidity.Subscore-sym.UPDR30F', '.UPDR30F.Num')
# Combine left and right and sym subscores into same column
ppmi_merge["Brady.Rigidity.Subscore.lateralized.UPDRS3"] =
ppmi_merge.pop("Brady.Rigidity.Subscore-right.UPDRS3").fillna(ppmi_merge.pop("Brady.Rigidity.Subscore-right.UPDRS3").
ppmi_merge["Brady.Rigidity.Subscore.lateralized.UPDRS3"] =
→ ppmi_merge.pop("Brady.Rigidity.Subscore.lateralized.UPDRS3").fillna(ppmi_merge.pop("Brady.Rigidity.
ppmi_merge["Brady.Rigidity.Subscore.lateralized.UPDRS3A"] =
ppmi_merge.pop("Brady.Rigidity.Subscore-right.UPDRS3A").fillna(ppmi_merge.pop("Brady.Rigidity.Subscore-right.UPDRS3A").
ppmi_merge["Brady.Rigidity.Subscore.lateralized.UPDRS3A"] =
→ ppmi_merge.pop("Brady.Rigidity.Subscore.lateralized.UPDRS3A").fillna(ppmi_merge.pop("Brady.Rigidity
ppmi_merge["Brady.Rigidity.Subscore.lateralized.UPDR30N"] =
→ ppmi_merge.pop("Brady.Rigidity.Subscore-right.UPDR30N").fillna(ppmi_merge.pop("Brady.Rigidity.Subsc
ppmi_merge["Brady.Rigidity.Subscore.lateralized.UPDR30N"] =
ppmi_merge.pop("Brady.Rigidity.Subscore.lateralized.UPDR30N").fillna(ppmi_merge.pop("Brady.Rigidity
ppmi_merge["Brady.Rigidity.Subscore.lateralized.UPDR30F"] =
ppmi_merge.pop("Brady.Rigidity.Subscore-right.UPDR30F").fillna(ppmi_merge.pop("Brady.Rigidity.Subscore-right.UPDR30F").
ppmi_merge["Brady.Rigidity.Subscore.lateralized.UPDR30F"] =
ppmi_merge.pop("Brady.Rigidity.Subscore.lateralized.UPDR30F").fillna(ppmi_merge.pop("Brady.Rigidity
#### TREMOR ####
```

```
tremor_right3 = ['Dominant.Side.Disease', 'Tremor.UPDRS2.Num',
          'Postural.Tremor.Right.Hand.UPDRS3.Num', 'Kinetic.Tremor.Right.Hand.UPDRS3.Num',
          'Rest.Tremor.Amplitude.RUE.UPDRS3.Num', 'Rest.Tremor.Amplitude.RLE.UPDRS3.Num',
          'Rest.Tremor.Amplitude.Lip.Jaw.UPDRS3.Num', 'Constancy.of.Rest.Tremor.UPDRS3.Num']
tremor_right3a = ['Dominant.Side.Disease', 'Tremor.UPDRS2.Num',
          'Postural.Tremor.Right.Hand.UPDRS3A.Num', 'Kinetic.Tremor.Right.Hand.UPDRS3A.Num',
          'Rest.Tremor.Amplitude.RUE.UPDRS3A.Num', 'Rest.Tremor.Amplitude.RLE.UPDRS3A.Num',
          'Rest.Tremor.Amplitude.Lip.Jaw.UPDRS3A.Num', 'Constancy.of.Rest.Tremor.UPDRS3A.Num']
tremor_right30N = ['Dominant.Side.Disease', 'Tremor.UPDRS2.Num',
          'Postural.Tremor.Right.Hand.UPDR3ON.Num', 'Kinetic.Tremor.Right.Hand.UPDR3ON.Num',
          'Rest.Tremor.Amplitude.RUE.UPDR3ON.Num', 'Rest.Tremor.Amplitude.RLE.UPDR3ON.Num',
          'Rest.Tremor.Amplitude.Lip.Jaw.UPDR3ON.Num', 'Constancy.of.Rest.Tremor.UPDR3ON.Num']
tremor_right30F = ['Dominant.Side.Disease', 'Tremor.UPDRS2.Num',
          'Postural.Tremor.Right.Hand.UPDR3OF.Num', 'Kinetic.Tremor.Right.Hand.UPDR3OF.Num',
          'Rest.Tremor.Amplitude.RUE.UPDR30F.Num', 'Rest.Tremor.Amplitude.RLE.UPDR30F.Num',
          'Rest.Tremor.Amplitude.Lip.Jaw.UPDR30F.Num', 'Constancy.of.Rest.Tremor.UPDR30F.Num']
tremor_left3 = ['Dominant.Side.Disease', 'Tremor.UPDRS2.Num',
          'Postural.Tremor.Left.Hand.UPDRS3.Num', 'Kinetic.Tremor.Left.Hand.UPDRS3.Num',
          'Rest.Tremor.Amplitude.LUE.UPDRS3.Num', 'Rest.Tremor.Amplitude.LLE.UPDRS3.Num'
          'Rest.Tremor.Amplitude.Lip.Jaw.UPDRS3.Num', 'Constancy.of.Rest.Tremor.UPDRS3.Num']
tremor_left3a = ['Dominant.Side.Disease', 'Tremor.UPDRS2.Num',
         'Postural.Tremor.Left.Hand.UPDRS3A.Num', 'Kinetic.Tremor.Left.Hand.UPDRS3A.Num',
          'Rest.Tremor.Amplitude.LUE.UPDRS3A.Num', 'Rest.Tremor.Amplitude.LLE.UPDRS3A.Num',
          'Rest.Tremor.Amplitude.Lip.Jaw.UPDRS3A.Num', 'Constancy.of.Rest.Tremor.UPDRS3A.Num']
tremor_left30N = ['Dominant.Side.Disease', 'Tremor.UPDRS2.Num',
          'Postural.Tremor.Left.Hand.UPDR3ON.Num', 'Kinetic.Tremor.Left.Hand.UPDR3ON.Num',
          'Rest.Tremor.Amplitude.LUE.UPDR30N.Num', 'Rest.Tremor.Amplitude.LLE.UPDR30N.Num',
          'Rest.Tremor.Amplitude.Lip.Jaw.UPDR3ON.Num', 'Constancy.of.Rest.Tremor.UPDR3ON.Num']
tremor_left30F = ['Dominant.Side.Disease', 'Tremor.UPDRS2.Num',
          \verb|'Postural.Tremor.Left.Hand.UPDR30F.Num', |'Kinetic.Tremor.Left.Hand.UPDR30F.Num', |'Kinetic.Tremor.Left.Hand.UPDR30
          'Rest.Tremor.Amplitude.LUE.UPDR30F.Num', 'Rest.Tremor.Amplitude.LLE.UPDR30F.Num',
          'Rest.Tremor.Amplitude.Lip.Jaw.UPDR3OF.Num', 'Constancy.of.Rest.Tremor.UPDR3OF.Num']
tremor_sym3 = ['Dominant.Side.Disease', 'Tremor.UPDRS2.Num',
          'Postural.Tremor.Right.Hand.UPDRS3.Num', 'Postural.Tremor.Left.Hand.UPDRS3.Num',
          'Kinetic.Tremor.Right.Hand.UPDRS3.Num', 'Kinetic.Tremor.Left.Hand.UPDRS3.Num',
          'Rest.Tremor.Amplitude.RUE.UPDRS3.Num', 'Rest.Tremor.Amplitude.LUE.UPDRS3.Num',
          'Rest.Tremor.Amplitude.RLE.UPDRS3.Num', 'Rest.Tremor.Amplitude.LLE.UPDRS3.Num',
          'Rest.Tremor.Amplitude.Lip.Jaw.UPDRS3.Num', 'Constancy.of.Rest.Tremor.UPDRS3.Num']
tremor sym3a = ['Dominant.Side.Disease', 'Tremor.UPDRS2.Num',
          'Postural.Tremor.Right.Hand.UPDRS3A.Num', 'Postural.Tremor.Left.Hand.UPDRS3A.Num',
          'Kinetic.Tremor.Right.Hand.UPDRS3A.Num', 'Kinetic.Tremor.Left.Hand.UPDRS3A.Num',
          \verb|'Rest.Tremor.Amplitude.RUE.UPDRS3A.Num', \verb|'Rest.Tremor.Amplitude.LUE.UPDRS3A.Num'|, amplitude.LUE.UPDRS3A.Num'|, amplitude.LUE.UPDRS3A.Num'|, amplitu
          'Rest.Tremor.Amplitude.RLE.UPDRS3A.Num', 'Rest.Tremor.Amplitude.LLE.UPDRS3A.Num',
          'Rest.Tremor.Amplitude.Lip.Jaw.UPDRS3A.Num', 'Constancy.of.Rest.Tremor.UPDRS3A.Num']
tremor_sym30N = ['Dominant.Side.Disease', 'Tremor.UPDRS2.Num',
          'Postural.Tremor.Right.Hand.UPDR3ON.Num', 'Postural.Tremor.Left.Hand.UPDR3ON.Num',
          'Kinetic.Tremor.Right.Hand.UPDR3ON.Num', 'Kinetic.Tremor.Left.Hand.UPDR3ON.Num',
          'Rest.Tremor.Amplitude.RUE.UPDR30N.Num', 'Rest.Tremor.Amplitude.LUE.UPDR30N.Num'
          'Rest.Tremor.Amplitude.RLE.UPDR30N.Num', 'Rest.Tremor.Amplitude.LLE.UPDR30N.Num',
          'Rest.Tremor.Amplitude.Lip.Jaw.UPDR3ON.Num', 'Constancy.of.Rest.Tremor.UPDR3ON.Num']
tremor_sym30F = ['Dominant.Side.Disease', 'Tremor.UPDRS2.Num',
          'Postural.Tremor.Right.Hand.UPDR30F.Num', 'Postural.Tremor.Left.Hand.UPDR30F.Num',
          'Kinetic.Tremor.Right.Hand.UPDR3OF.Num', 'Kinetic.Tremor.Left.Hand.UPDR3OF.Num',
          \label{lem:continuous} \begin{tabular}{ll} \tt 'Rest.Tremor.Amplitude.LUE.UPDR30F.Num', 29' \tt Rest.Tremor.Amplitude.LUE.UPDR30F.Num', \tt 'Rest.Tremor.Amplitude.LLE.UPDR30F.Num', \tt 'Rest.Tremor.Ampli
          'Rest.Tremor.Amplitude.Lip.Jaw.UPDR3OF.Num', 'Constancy.of.Rest.Tremor.UPDR3OF.Num']
```

```
ppmi_merge = add_lateralized_subscores(ppmi_merge, tremor_left3, 'Left',
ppmi_merge = add_lateralized_subscores(ppmi_merge, tremor_left3a, 'Left',
ppmi_merge = add_lateralized_subscores(ppmi_merge, tremor_left30N, 'Left',
→ 'Tremor.Subscore-left.UPDR30N')
ppmi_merge = add_lateralized_subscores(ppmi_merge, tremor_left30F, 'Left',
→ 'Tremor.Subscore-left.UPDR30F')
ppmi_merge = add_lateralized_subscores(ppmi_merge, tremor_right3, 'Right',
ppmi_merge = add_lateralized_subscores(ppmi_merge, tremor_right3a, 'Right',
→ 'Tremor.Subscore-right.UPDRS3A')
ppmi_merge = add_lateralized_subscores(ppmi_merge, tremor_right30N, 'Right',
→ 'Tremor.Subscore-right.UPDR3ON')
ppmi_merge = add_lateralized_subscores(ppmi_merge, tremor_right30F, 'Right',
→ 'Tremor.Subscore-right.UPDR30F')
ppmi_merge = add_symmetric_subscores(ppmi_merge, tremor_sym3, 'Symmetric',
→ 'Tremor.Subscore-sym.UPDRS3', '.UPDRS3.Num')
ppmi_merge = add_symmetric_subscores(ppmi_merge, tremor_sym3a, 'Symmetric',
→ 'Tremor.Subscore-sym.UPDRS3A', '.UPDRS3A.Num')
ppmi_merge = add_symmetric_subscores(ppmi_merge, tremor_sym30N, 'Symmetric',
→ 'Tremor.Subscore-sym.UPDR30N', '.UPDR30N.Num')
ppmi_merge = add_symmetric_subscores(ppmi_merge, tremor_sym30F, 'Symmetric',
→ 'Tremor.Subscore-sym.UPDR30F', '.UPDR30F.Num')
# Combine left and right and sym scores into same column (.lateralized)
ppmi_merge["Tremor.Subscore.lateralized.UPDRS3"] =
ppmi_merge.pop("Tremor.Subscore-right.UPDRS3").fillna(ppmi_merge.pop("Tremor.Subscore-left.UPDRS3")
ppmi_merge["Tremor.Subscore.lateralized.UPDRS3"] =
→ ppmi_merge.pop("Tremor.Subscore.lateralized.UPDRS3").fillna(ppmi_merge.pop("Tremor.Subscore-sym.UPD
ppmi_merge["Tremor.Subscore.lateralized.UPDRS3A"] =
ppmi_merge.pop("Tremor.Subscore-right.UPDRS3A").fillna(ppmi_merge.pop("Tremor.Subscore-left.UPDRS3A")
ppmi_merge["Tremor.Subscore.lateralized.UPDRS3A"] =
ppmi_merge.pop("Tremor.Subscore.lateralized.UPDRS3A").fillna(ppmi_merge.pop("Tremor.Subscore-sym.UPDRS3A").
ppmi_merge["Tremor.Subscore.lateralized.UPDR30N"] =
ppmi_merge.pop("Tremor.Subscore-right.UPDR30N").fillna(ppmi_merge.pop("Tremor.Subscore-left.UPDR30N").
ppmi_merge["Tremor.Subscore.lateralized.UPDR30N"] =
ppmi_merge.pop("Tremor.Subscore.lateralized.UPDR30N").fillna(ppmi_merge.pop("Tremor.Subscore-sym.UPDR30N").
ppmi_merge["Tremor.Subscore.lateralized.UPDR30F"] =
ppmi_merge.pop("Tremor.Subscore-right.UPDR30F").fillna(ppmi_merge.pop("Tremor.Subscore-left.UPDR30F")
ppmi_merge["Tremor.Subscore.lateralized.UPDR30F"] =
ppmi_merge.pop("Tremor.Subscore.lateralized.UPDR30F").fillna(ppmi_merge.pop("Tremor.Subscore-sym.UPDR30F").
## If lateralized subscore is nan - input the non-lateralized score for Tremor and
\rightarrow Brady.Rigidity
def fill_non_lat_subscore(df, subscore_lateralized_name, subscore_name) :
```

```
latisna = df[subscore lateralized name].isna() & df[subscore name].notna()
   df[subscore_lateralized_name].loc[latisna] = df[subscore_name].loc[latisna]
   return df
ppmi_merge = fill_non_lat_subscore(ppmi_merge, 'Tremor.Subscore.lateralized.UPDRS3',
ppmi merge = fill non lat subscore(ppmi merge, 'Tremor.Subscore.lateralized.UPDRS3A',
ppmi_merge = fill_non_lat_subscore(ppmi_merge, 'Tremor.Subscore.lateralized.UPDR3ON',
ppmi_merge = fill_non_lat_subscore(ppmi_merge, 'Tremor.Subscore.lateralized.UPDR3OF',
ppmi_merge = fill_non_lat_subscore(ppmi_merge,
→ 'Brady.Rigidity.Subscore.lateralized.UPDRS3', 'Brady.Rigidity.Subscore.UPDRS3')
ppmi_merge = fill_non_lat_subscore(ppmi_merge,
→ 'Brady.Rigidity.Subscore.lateralized.UPDRS3A', 'Brady.Rigidity.Subscore.UPDRS3A')
ppmi_merge = fill_non_lat_subscore(ppmi_merge,
→ 'Brady.Rigidity.Subscore.lateralized.UPDR3ON', 'Brady.Rigidity.Subscore.UPDR3ON')
ppmi_merge = fill_non_lat_subscore(ppmi_merge,
→ 'Brady.Rigidity.Subscore.lateralized.UPDR30F', 'Brady.Rigidity.Subscore.UPDR30F')
## Inlcude columns for bestEventID (bestScreening, bestBaseline, etc) and denote the
→ highest resnetGrade with True (else = False)
myevs = ppmi_merge['Event.ID'].unique() # Unique event ids
uids = ppmi_merge['Subject.ID'].unique() # Unique subject ids
for myev in myevs :
   mybe = "best" + myev # Create best Visit column
   ppmi_merge[mybe] = False # Set all best visit to be False
   for u in uids :
       selu = ppmi_merge.loc[(ppmi_merge['Subject.ID'] == u) & (ppmi_merge['Event.ID']
--- myev) & (ppmi_merge['resnetGrade'].notna())] # For one subject at one event id if
\rightarrow resnetGrade not na
       if len(selu) == 1 : # If there is one event id for that subject
           idx = selu.index # Ge the index
           ppmi_merge.loc[idx, mybe] = True
       if len(selu) > 1 : # IF there is more than one event id for that subject and
        → resnet grade is not na
           maxidx = selu[['resnetGrade']].idxmax() # Get the higher resnetGrade for each
  visit if there are more than one
           ppmi_merge.loc[maxidx, mybe] = True
## Include a column for bestAtImage.Acquisition.Date - denote the one or highest
→ resnetGrade with True (else = False)
ppmi_merge['bestAtImage.Acquisition.Date'] = False # Initialize
→ bestAtImage.Acuqisition.Date col
for myev in myevs :
   for u in uids :
       selu = ppmi_merge.loc[(ppmi_merge['Subject.ID'] == u) & (ppmi_merge['Event.ID']
  == myev) & (ppmi_merge['resnetGrade'].notna())]
       if len(selu) == 1 : # If there is one event id for that subject
           idx = selu.index # Get the index
```

```
ppmi_merge.loc[idx, 'bestAtImage.Acquisition.Date'] = True
       if len(selu) > 1 : # IF there is more than one event id for that subject and
        → resnet grade is not na
           maxidx = selu[['resnetGrade']].idxmax() # Get the higher resnetGrade for each
→ visit if there are more than one
           ppmi_merge.loc[maxidx, 'bestAtImage.Acquisition.Date'] = True
# DX simplified column
ppmi_merge['DXsimplified'] = '' # Initialize DXsimplfied
ppmi_merge.loc[ppmi_merge['Consensus.Subtype'] == 'Healthy Control', 'DXsimplified'] =
ppmi merge.loc[ppmi merge['Enroll.Diagnosis'] == 'Healthy Control', 'DXsimplified'] =
ppmi_merge.loc[ppmi_merge['Enroll.Diagnosis'] == 'Parkinson\'s Disease', 'DXsimplified']
ppmi_merge.loc[ppmi_merge['Enroll.Diagnosis'] == 'SWEDD', 'DXsimplified'] = "nonPDorMSA"
ppmi_merge.loc[ppmi_merge['Enroll.Diagnosis'] == "Prodromal", 'DXsimplified'] =
ppmi_merge.loc[ppmi_merge['Consensus.Subtype'] == "GBA", 'DXsimplified'] = "GBA_HC"
ppmi_merge.loc[ppmi_merge['Consensus.Subtype'] == "Genetic : GBA", 'DXsimplified'] =

→ "GBA_PD"

ppmi_merge.loc[ppmi_merge['Consensus.Subtype'] == "Genetic : GBA not Prodromal",

    'DXsimplified'] = "GBA_PD"

ppmi merge.loc[ppmi merge['Consensus.Subtype'] == "Genetic : GBA Prodromal",
→ 'DXsimplified'] = "GBA Pro"
ppmi_merge.loc[ppmi_merge['Consensus.Subtype'] == "Genetic : LRRK2", 'DXsimplified'] =
→ "LRRK2 PD"
ppmi_merge.loc[ppmi_merge['Consensus.Subtype'] == "Genetic : LRRK2 + GBA",
→ 'DXsimplified'] = "LRRK2 PD"
ppmi_merge.loc[ppmi_merge['Consensus.Subtype'] == "Genetic : LRRK2 + GBA not Prodromal",
→ 'DXsimplified'] = "LRRK2_PD"
ppmi_merge.loc[ppmi_merge['Consensus.Subtype'] == "Genetic : LRRK2 + GBA Prodromal",
→ 'DXsimplified'] = "LRRK2_Pro"
ppmi_merge.loc[ppmi_merge['Consensus.Subtype'] == "Genetic : LRRK2 not Prodromal",
→ 'DXsimplified'] = "LRRK2_PD"
ppmi_merge.loc[ppmi_merge['Consensus.Subtype'] == "Genetic : LRRK2 Phenoconverted",
→ 'DXsimplified'] = "LRRK2 Pro"
ppmi_merge.loc[ppmi_merge['Consensus.Subtype'] == "Genetic : LRRK2 Prodromal",
→ 'DXsimplified'] = "LRRK2 Pro"
ppmi_merge.loc[ppmi_merge['Consensus.Subtype'] == "Genetic : SNCA", 'DXsimplified'] =
→ "SNCA PD"
ppmi_merge.loc[ppmi_merge['Consensus.Subtype'] == "Genetic : SNCA Prodromal",
→ 'DXsimplified'] = "SNCA Pro"
ppmi_merge.loc[ppmi_merge['Consensus.Subtype'] == "Hyposmia" , 'DXsimplified'] =
→ "Sporadic Pro"
ppmi_merge.loc[ppmi_merge['Consensus.Subtype'] == "Hyposmia : Phenoconverted",
→ 'DXsimplified'] = "Sporadic_Pro"
ppmi_merge.loc[ppmi_merge['Consensus.Subtype'] == "No Mutation not Prodromal",
ppmi_merge.loc[ppmi_merge['Consensus.Subtype'] == "non-HC", 'DXsimplified'] = np.NaN
ppmi_merge.loc[ppmi_merge['Consensus.Subtype'] == "non-PD", 'DXsimplified'] =

→ "nonPDorMSA"

ppmi_merge.loc[ppmi_merge['Consensus.Subtype'] == "RBD", 'DXsimplified'] = "Sporadic_Pro"
```

```
ppmi_merge.loc[ppmi_merge['Consensus.Subtype'] == "RBD : Phenoconverted", 'DXsimplified']
ppmi_merge.loc[ppmi_merge['Consensus.Subtype'] == "RBD : Phenoconverted with GBA",

    'DXsimplified'] = "GBA_Pro"

ppmi_merge.loc[ppmi_merge['Consensus.Subtype'] == "Sporadic", 'DXsimplified'] =

→ "Sporadic_PD"

ppmi merge.loc[ppmi merge['Consensus.Subtype'] == "SWEDD/non-PD Active", 'DXsimplified']
ppmi_merge.loc[ppmi_merge['Consensus.Subtype'] == "SWEDD/PD Active", 'DXsimplified'] =
\rightarrow np.NaN
## Add in min PD Disease duration
# Create a simplified diagnosis group of just HC, PD, Prodromal, or nonPDorMSA
ppmi_merge['DXs2'] = ''
ppmi_merge['DXsimplified'].fillna('NA', inplace = True)
ppmi_merge.loc[ppmi_merge['DXsimplified'].str.contains('HC'), 'DXs2'] = 'HC'
ppmi_merge.loc[ppmi_merge['DXsimplified'].str.contains('_PD'), 'DXs2'] = 'PD'
ppmi merge.loc[ppmi merge['DXsimplified'].str.contains(' Pro'), 'DXs2'] = 'Pro'
ppmi_merge.loc[ppmi_merge['DXsimplified'].str.contains('nonPDorMSA'), 'DXs2'] =
→ 'nonPDorMSA'
# If HC, Prodromal or nonPDorMSA - fill in with 0 for PD.Min.Disease.Duration
ppmi_merge.loc[ppmi_merge['DXs2'] == 'HC', 'PD.Min.Disease.Duration'] = 0
ppmi_merge.loc[ppmi_merge['DXs2'] == 'Pro', 'PD.Min.Disease.Duration'] = 0
ppmi_merge.loc[ppmi_merge['DXs2'] == 'nonPDorMSA', 'PD.Min.Disease.Duration'] = 0
# For PD subjects, fill in PD.Min.Disease.Duration
print(ppmi_merge['PD.Diagnosis.Duration'])
ppmi merge.to csv('/Users/areardon/Desktop/ppmi merge1.csv')
ppmi merge['PD.Diagnosis.Duration'].fillna('', inplace = True)
uids = ppmi_merge['Subject.ID'].unique() # Unique subject ids
for u in uids :
   ww = ppmi_merge[(ppmi_merge['Subject.ID'] == u) & (ppmi_merge['DXs2'] == 'PD') &
if len(ww) > 0:
       print(ww['PD.Diagnosis.Duration'])
       mydd = min(ww['PD.Diagnosis.Duration'])
       ppmi_merge.loc[ppmi_merge['Subject.ID'] == u, 'PD.Min.Disease.Duration'] = mydd
## Add in Visit column
ppmi_merge['Visit'] = np.NaN # Initialize Visit col
searchfor = ['Baseline', 'Visit Month '] # Strings to search for
temp = ppmi_merge['Event.ID'].str.contains('|'.join(searchfor)) # locations of where row
→ contains str: baseline or visit month
ppmi_merge['Visit'].loc[temp == True] = ppmi_merge['Event.ID'].loc[temp == True] # Fill
→ in 'Visit' col with event.ID for baseline or visit month
ppmi_merge['Visit'] = ppmi_merge['Visit'].str.replace("Remote Visit Month ", "") #
→ Replace Remote visit month with '' (want only month number)
ppmi_merge['Visit'] = ppmi_merge['Visit'].str.replace("Visit Month ", "") # Replace Visit
→ month with '' (want only month number)
```

```
ppmi_merge['Visit'] = ppmi_merge['Visit'].str.replace("Baseline", "0") # Replace baseline
    with 0

ppmi_merge['Visit'] = ppmi_merge['Visit'].fillna(9999) # Filling with 9999 so we can
    change this col to int

ppmi_merge['Visit'] = ppmi_merge['Visit'].astype(int) # str to int

ppmi_merge['Visit'] = ppmi_merge['Visit'].replace(9999, np.nan) # Replace 9999 with nan

# Final re-organization of ppmi_merge and save

ppmi_merge.set_index('Subject.ID', inplace = True)

ppmi_merge.fillna('NA', inplace = True)

ppmi_merge.to_csv(userdir + 'ppmi_merge_v0.csv')
```