

Variable Definitions and Score Calculations

Summary

The following tables describe the calculation of the derived variables in the PPMI study.

Study Groups

Study Groups		
Characteristic	Variables	Dataset
Enrolled PD Subject	PATNO, APPRDX, ENROLLDT Merge SCREEN with RANDOM and find each unique PATNO with APPRDX = '1' that is not missing ENROLLDT	RANDOM (Consent and Enrollment), SCREEN
Enrolled Healthy Control	PATNO, APPRDX, ENROLLDT Merge SCREEN with RANDOM and find each unique PATNO with APPRDX = '2' that is not missing ENROLLDT	RANDOM (Consent and Enrollment), SCREEN
Enrolled SWEDD Subject	PATNO, APPRDX, ENROLLDT Merge SCREEN with RANDOM and find each unique PATNO with APPRDX = '3' that is not missing ENROLLDT	RANDOM (Consent and Enrollment), SCREEN
Enrolled Prodromal Subject	PATNO, APPRDX, ENROLLDT Merge SCREEN with RANDOM and find each unique PATNO with APPRDX = '4' that is not missing ENROLLDT	RANDOM (Consent and Enrollment), SCREEN
Enrolled Genetic Cohort Subject	PATNO, APPRDX, ENROLLDT Merge SCREEN with RANDOM and find each unique PATNO with APPRDX = '5' or '6' that is not missing ENROLLDT APPRDX = '5' are PD subjects APPRDX = '6' are Unaffected subjects	RANDOM (Consent and Enrollment), SCREEN
Enrolled Genetic Registry Subject	PATNO, APPRDX, ENROLLDT Merge SCREEN with RANDOM and find each unique PATNO with APPRDX = '7' or '8' that is not	RANDOM (Consent and Enrollment),



missing ENROLLDT	SCREEN
APPRDX = '7' are PD subjects APPRDX = '8' are Unaffected subjects	

Demographics and PD Characteristics

	Demographics and PD Characteristics	
Age	ENROLLDT - BIRTHDT	RANDOM (Consent and Enrollment)
Gender	GENDER = 2 is Male GENDER = 0 or 1 is Female	RANDOM (Consent and Enrollment)
Race	RAINDALS, RAASIAN, RABLACK, RAHAWOPI, RAWHITE, RANOS Other = RAINDALS, RAHAWOPI, RANOS, or more than one race specified	SCREEN
Family History of PD	BIOMOMPD, BIODADPD, FULSIBPD, HALFSIBPD, MAGPARPD, PAGPARPD, MATAUPD, PATAUPD, KIDSPD Subject has family history of PD if any one or more of these variables = '1'	FAMHXPD
Disease Duration	PDDXDT, ENROLLDT Duration = number of months between PDDXDT and ENROLLDT	PDFEAT, RANDOM (Consent and Enrollment)
TD / PIGD Classification	First calculate Tremor and PIGD scores: Tremor score = Mean of the following variables: NP2TRMR, NP3PTRMR, NP3PTRML, NP3KTRMR, NP3KTRML, NP3RTARU, NP3RTALU, NP3RTARL, NP3RTALL, NP3RTALJ, NP3RTCON PIGD score = Mean of the following variables: NP2WALK, NP2FREZ, NP3GAIT, NP3FRZGT, NP3PSTBL	NUPDRS2P, NUPDRS3
	Then calculate ratio = Tremor score / PIGD score.	



If ratio \geq 1.15, OR if PIGD score = 0 and Tremor	
score > 0, then subject is TD.	
If ratio ≤ 0.9 then subject is PIGD.	
If ratio > 0.9 and < 1.15, OR if Tremor score and	
PIGD score = 0, then subject is Indeterminate.	

Motor Assessments

	Wiotor Assessments	
MDS-UPDRS Part I	NP1COG, NP1HALL, NP1DPRS, NP1ANXS, NP1APAT,NP1DDS, NP1SLPN, NP1SLPD, NP1PAIN, NP1URIN, NP1CNST, NP1LTHD, NP1FATG	NUPDRS1, NUPDRS1P
	Part I Score = sum of these 13 variables	
MDS-UPDRS Part II	NP2SPCH, NP2SALV, NP2SWAL, NP2EAT, NP2DRES, NP2HYGN, NP2HWRT, NP2HOBB, NP2TURN, NP2TRMR, NP2RISE, NP2WALK, NP2FREZ	NUPDRS2P
	Part II Score = sum of these 13 variables	
MDS-UPDRS Part III	NP3SPCH, NP3FACXP, NP3RIGN, NP3RIGRU, NP3RIGLU, PN3RIGRL, NP3RIGLL, NP3FTAPR, NP3FTAPL, NP3FTAPL, NP3FTAPL, NP3PRSPR, NP3PRSPL, NP3TTAPR, NP3TTAPL, NP3LGAGR, NP3LGAGL, NP3RISNG, NP3GAIT, NP3FRZGT, NP3PSTBL, NP3POSTR, NP3BRADY, NP3PTRMR, NP3PTRML, NP3KTRMR, NP3KTRML, NP3RTALU, NP3RTALU, NP3RTALU, NP3RTALL, NP3RTALL, NP3RTALJ,	NUPDRS3
MDS-UPDRS Part IV	NP4WDYSK, NP4DYSKI, NP4OFF, NP4FLCTI, NP4FLCTX, NP4DYSTN Part IV Score = sum of these 6 variables	NUPDRS4
MDS-UPDRS Total Score	Sum of MDS-UPDRS Parts I, II, III	NUPDRS1, NUPDRS1P, NUPDRS2P, NUPDRS3



Non-Motor Assessments

Benton Judgment of Line Orientation Score	Sum of BJLOT1 – BJLOT30	LINEORNT
Epworth Sleepiness Scale	Sum of ESS1 - ESS8 Subjects with ESS < 10 are "Not Sleepy" Subjects with ESS ≥ 10 are "Sleepy"	EPWORTH
GDS Raw Score	Add 1 point for each response of "No" (0) to any of the following variables: GDSSATIS, GDSGSPIR, GDSHAPPY, GDSALIVE, GDSENRGY Add 1 point for each response of "Yes" (1) to any of the following variables: GDSDROPD, GDSEMPTY, GDSBORED, GDSAFRAD, GDSHLPLS, GDSHOME, GDSMEMRY, GDSWRTLS, GDSHOPLS, GDSBETER Subjects with GDS ≥ 5 are "Depressed" Subject with GDS < 5 are "Not Depressed"	GDSSHORT
HVLT Immediate/Total Recall	Sum of HVLTRT1 - HVLTRT3	HVLT
HVLT Discrimination Recognition	HVLTREC - (HVLTFPRL + HVLTFPUN)	HVLT
HVLT Retention	HVLTRDLY / max(HVLTRT2, HVLTRT3)	HVLT
Letter Number Sequencing (LNS)	Sum of LNS1A – LNS7C	LNSPD
MOCA Total Score	Unadjusted Score = sum of MCAALTTM, MCACUBE, MCACLCKC, MCACLCKN, MCACLCKH, MCALION, MCARHINO, MCACAMEL, MCAFDS, MCABDS, MCAVIGIL, MCASER7, MCASNTNC, MCAVF, MCAABSTR, MCAREC1, MCAREC2, MCAREC3, MCAREC4, MCAREC5, MCADATE, MCAMONTH, MCAYR,	MOCA, SOCIOECO



	MCADAY, MCAPLACE, MCACITY If EDUCYRS ≤ 12 and Unadjusted Score < 30, add 1 more point to score. If EDUCYRS > 12, do not add any more points to score.	
QUIP	For Sections A - D, add 1 point if either question has a response of "Yes" (1): Section A: CNTRLGMB, TMGAMBLE Section B: CNTRLSEX, TMSEX Section C: CNTRLBUY, TMBUY Section D: CNTRLEAT, TMEAT For Section E, add 1 point for each response of "Yes" (1): TMTORACT, TMTMTACT, TMTRWD	QUIPCS
REM Sleep Behavior Disorder (RBD)	Add 1 point for each response of "Yes" (1) to any of the following variables: DRMVIVID, DRMAGRAC, DRMNOCTB, SLPLMBMV, SLPINJUR, DRMVERBL, DRMFIGHT, DRMUMV, DRMOBJFL, MVAWAKEN, DRMREMEM, SLPDSTRB	REMSLEEP
	Add 1 point if <u>any</u> of the following variables has a response of "Yes" (1): STROKE, HETRA, PARKISM, RLS, NARCLPSY, DEPRS, EPILEPSY, BRNINFM, CNSOTH	
	If any of the previous variables are missing, then RBD score is missing. Subjects with score ≥ 5 are RBD Positive	
SCOPA-AUT Total	Subjects with score < 5 are RBD Negative SCAU1 - SCAU25	SCOPA
Score	For questions 1-21 (SCAU1 - SCAU21), add 3 points for each response of "9." Otherwise, add the number of points in response.	
	For questions 22-25 (SCAU22 - SCAU25), add 0 points for each response of "9." Otherwise, add the number of points in response.	



Semantic Fluency (SFT)	Sum of VLTANIM, VLTVEG, VLTFRUIT	SFT
State Trait Anxiety Total Score (STAI)	STAIAD1 - STAIAD40	STAI
	Add values for the following questions: 3, 4, 6, 7, 9, 12, 13, 14, 17, 18, 22, 24, 25, 28, 29, 31, 32, 35, 37, 38, 40	
	Use reverse scoring for the remaining questions and add to the first score (e.g., if value = 1, add 4 points to score; if value = 2, add 3 points to score, etc).	
STAI - State Subscore	STAIAD1 - STAIAD20	STAI
	Add values for the following questions: 3, 4, 6, 7, 9, 12, 13, 14, 17, 18	
	Use reverse scoring for the values of the remaining questions through question 20 and add to the first value.	
STAI - Trait Subscore	STAIAD21 - STAIAD40	STAI
	Add values for the following questions: 22, 24, 25, 28, 29, 31, 32, 35, 37, 38, 40	
	Use reverse scoring for the values of the remaining questions and add to the first value.	
UPSIT Raw Score	Sum of UPSITBK1 - UPSITBK4	UPSIT



Cognitive

Mild Cognitive Impairment (MCI) COGDECLN, FNCDTCOG, DVT_TOT DVT_RECOG_DISC_INDEX, DVS_JI DVS_LNS, DVT_SFTANIM, DVT_SD Subject has MCI if the following 3 criter 1) Cognitive decline marked as 'Yes' (€ 2) Any 2 or more of the following cogn SD below the standardized mean: • HVLT Total Recall (DVT_TOTAL • HVLT Recognition Discrimination (DVT_RECOG_DISC_INDEX ≤ 3 • Benton Judgment of Line Orientati (DVS_JLO_MSSAE ≤ 6) • Letter Number Sequencing (DVS_J	LO_MSSAE, DM eria are met: (COGDECLN = '1') nitive tests are >1.5 L_RECALL \le 35) n 35)	
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DaTSCAN

Contralateral	For Healthy Controls, no contralateral side is identified. Instead, the average of the left and right values is calculated.	SBR, PDFEAT
	For PD and SWEDD subjects: Use Left value if DOMSIDE = '2' (Right) Use Right value if DOMSIDE = '1' (Left) Use the average of the left and right values if DOMSIDE = '3' (Symmetric)	
Ipsilateral	For Healthy Controls, no ipsilateral side is identified. Instead, the average of the left and right values is calculated.	SBR, PDFEAT
	For PD and SWEDD subjects: Use Right value if DOMSIDE = '2' (Right) Use Left value if DOMSIDE = '1' (Left) Use the average of the left and right values if DOMSIDE = '3' (Symmetric)	



Mean Caudate	(CAUDATE_R + CAUDATE_L) / 2	SBR
Mean Putamen	(PUTAMEN_R + PUTAMEN_L) / 2	SBR
Mean Striatum	(CAUDATE_R + CAUDATE_L + PUTAMEN_R + PUTAMEN_L) / 4	SBR
Count Density Ratio	Caudate / Putamen	SBR, PDFEAT
Asymmetry Index	$\left \frac{left - right}{mean(left + right)} \times 100 \right $	SBR, PDFEAT

CSF

Note: In cases where multiple CSF results are available for the same visit, the statistical core uses the results with the most recent date (variable: RUNDATE).		
Abeta 1-42	TESTNAME, TESTVALUE	
	Create a subset of observations that have TESTNAME = "Abeta 42". The value of Abeta 42 is the variable TESTVALUE.	BIOSPECAN
t-tau	TESTNAME, TESTVALUE	
	Create a subset of observations that have TESTNAME = "Total tau". The value of Total Tau is the variable TESTVALUE.	BIOSPECAN
p-tau	TESTNAME, TESTVALUE	
	Create a subset of observations that have TESTNAME = "p-Tau 181P". The value of p-Tau 181P is the variable TESTVALUE.	BIOSPECAN
Alpha-synuclein	TESTNAME, TESTVALUE	
	Create a subset of observations that have TESTNAME = "CSF Alpha-synuclein". The value of CSF Alpha-synuclein is the variable TESTVALUE.	BIOSPECAN
t-tau/Abeta 1-42	"Total Tau" / "Abeta 42"	BIOSPECAN
p-tau/Abeta 1-42	"p-Tau 181P" / "Abeta 42"	BIOSPECAN
p-tau/t-tau	"p-Tau 191P" / "Total tau"	BIOSPECAN



CSF Hemoglobin	TESTNAME, TESTVALUE	
	Create a subset of observations that have TESTNAME = "CSF Hemoglobin". The value of CSF Hemoglobin is the variable TESTVALUE.	BIOSPECAN

Biospecimen

Urate	LTSTNAME, LSIRES, LSIUNIT	COVANCE
	Create a subset of observations that have LTSTNAME = "Serum Uric Acid". The value of Urate is the variable LSIRES, and the units are the variable LSIUNIT.	

Genetics

ApoE	TESTNAME, TESTVALUE	BIOSPECAN
	Create a subset of observations that have TESTNAME = "APOE GENOTYPE" or "ApoE_Genotype". The genotype is the variable TESTVALUE.	
SNCA rs3910105	TESTNAME, TESTVALUE	BIOSPECAN
	Create a subset of observations that have TESTNAME = "rs3910105". The genotype is the variable TESTVALUE.	
SNCA rs356181	TESTNAME, TESTVALUE	BIOSPECAN
	Create a subset of observations that have TESTNAME = "rs356181". The genotype is the variable TESTVALUE.	
MAPT H1/H2	Extract the following 5 SNPs from the genetics data and count the number of minor alleles in each SNP: rs17652121, minor allele = C rs8070723, minor allele = G rs1052587, minor allele = C rs16940799, minor allele = C rs17652748, minor allele = T If the sum of minor alleles for all 5 SNPs is:	NeuroX SNP Data
	 the sum of minor affeles for all 5 SNPs is: < 5, then genotype = H1/H1 = 5, then genotype = H1/H2 	
	• = 10, then genotype = H2/H2	



Levodopa Equivalent Daily Dose

Levodopa Equivalent Daily Dose (LEDD)	The variable LEDD shows the Levodopa equivalent daily dose for each individual PD medication. To find the total LEDD at a specific time point, add all values of the variable LEDD for each PD medication being taken at that time point.	CONMED
	Anticholinergics and other medications that are not included in the calculation of the total LEDD will have a missing value for the variable LEDD.	
	For COMT inhibitors, the variable LEDD will read "LD x 0.5" or "LD x 0.33". To find the LEDD for COMT inhibitors, first find the total dose of <u>Levodopa only</u> , and then multiply that value by either 0.5 or 0.33 as instructed.	
	Note: The variable LEDD is only updated quarterly, so this variable will not necessarily be up-to-date with the other data in the CONMED data file.	

About the Authors

This document was prepared by Christopher S. Coffey, Chelsea J. Caspell-Garcia, and Eric D. Foster, University of Iowa, Department of Biostatistics Clinical Trials Statistical and Data Management Center. For more information please contact Eric D. Foster at (319)384-4188 or by e-mail at eric-foster@uiowa.edu.

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