

Differences in the presentation and progression of Parkinson's disease by sex

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Complete List of Authors:	<p>Iwaki, Hirotaka; National Institutes of Health, NIA/LNG Blauwendraat, Cornelis; National Institute of Neurological Disorders and Stroke, Leonard, Hampton; National Institutes of Health, NIA/LNG Makarious, Mary; NIH Kim, Jonggeol; NIH, Laboratory of Neurogenetics Liu, Ganqiang; School of Medicine Maple Grødem, Jodi; Stavanger University Hospital, The Norwegian Centre for Movement Disorders; University in Stavanger, Centre for Organelle Research Corvol, Jean-Christophe ; Hopital Pitie-Salpetriere, Pihlstrøm, Lasse; Oslo University Hospital, Department of Neurology van Nimwegen, Marlies; Radboudumc, Neurology Smolensky, Luba; Michael J Fox Foundation for Parkinson's Research Amondikar, Ninad; Michael J Fox Foundation for Parkinson's Research Hutten, Samantha; Michael J Fox Foundation for Parkinson's Research Frasier, Mark; Michael J. Fox Foundation, Nguyen, Khanh-Dung; Biogen Inc, Translational Genome Science Rick, Jacqueline; University of Pennsylvania, Neurology Eberly, Shirley; University of Rochester, Biostatistics Faghri, Faraz; National Institute of Neurological Disorders and Stroke, Auinger, Peggy; University of Rochester, Neurology Scott, Kirsten; University of Cambridge, John Van Geest Centre for Brain Repair, Department of Clinical Neurosciences Wijeyekoon, Ruwani; University of Cambridge, John van Geest Centre for Brain Repair, Clinical Neurosciences Van Deerlin, Vivianna; University of Pennsylvania, Pathology and Laboratory Medicine Hernandez, Dena; NIH, Laboratory of Neurogenetics Gibbs, J; National Institutes of Health, NIA/LNG; University College London, Department of Molecular Neuroscience and Reta Lila Weston Institute of Neurological Studies Day-Williams, Aaron; Flagship Labs 60 Inc Brice, A.; Institut du cerveau et de la moelle épinière, Alves, Guido; Helse Stavanger HF, Norwegian Centre for Movement Disorders; Universitetet i Stavanger, Department of Chemistry, Bioscience and Environmental Engineering Noyce, Alastair; Queen Mary University of London, Wolfson Institute of Preventive Medicine; Tysnes, Ole Evans, Jonathan Breen, David; Cambridge Centre for Brain Repair, University of Cambridge Estrada, Karol; Biogen Inc, Translational Genome Sciences Wegel, Claire; Indiana University Purdue University at Indianapolis,</p>

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	<p>Department of Medical and Molecular Genetics, Indiana University Danjou, Fabrice; Institut du cerveau et de la moelle épinière Simon, David; Beth Israel Deaconess Medical Center, Neurology Andreassen, Ole; University of Oslo Ravina, Bernard; Voyager Therapeutics Inc Toft, Mathias; Oslo University Hospital Heutink, Peter; Hertie Institute for Clinical Brain Research, University of Tübingen, and DZNE, German Center for Neurodegenerative Diseases, Department of Neurodegenerative Diseases Bloem, Bastiaan; Radboud University Donders Institute for Brain Cognition and Behaviour Weintraub, Daniel; University of Pennsylvania, Psychiatry Barker, Roger; university of cambridge, neuroscience Williams-Gray, Caroline; University of Cambridge, Centre for Brain Repair van de Warrenburg, Bart; Radboud University Nijmegen Medical Centre, Neurology van Hilten, Jacobus; LUMC, Neurology Scherzer, Clemens; Harvard Medical School and Brigham and Women's Hospital, Center for Neurologic Diseases; Harvard Medical School, Harvard NeuroDiscovery Center Biomarker Program Singleton, Andrew; NIH Nalls, Mike; Data Tecnica International</p>
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Differences in the presentation and progression of Parkinson's disease by sex

Hirota Iwaki MD^{1,2}, Cornelis Blauwendraat PhD¹, Hampton L. Leonard MS^{1,2}, Mary B. Makarious BA¹, Jonggeol J. Kim BA¹, Ganqiang Liu PhD^{3,4,5}, Jodi Maple-Grødem PhD^{6,7}, Jean-Christophe Corvol MD⁸, Lasse Pihlstrøm MD⁹, Marlies van Nimwegen PhD¹⁰, Luba Smolensky MS¹¹, Ninad Amondikar BA¹¹, Samantha J. Hutten PhD¹¹, Mark Frasier PhD¹¹, Khanh-Dung H. Nguyen PhD¹², Jacqueline Rick PhD¹³, Shirley Eberly MS¹⁴, Faraz Faghri PhD¹, Peggy Auinger MS¹⁵, Kirsten M. Scott MRCP¹⁶, Ruwani Wijeyekoon MRCP¹⁶, Vivianna M. Van Deerlin MD¹⁷, Dena G. Hernandez PhD¹, J. Raphael Gibbs PhD¹, Aaron G. Day-Williams PhD^{18,19}, Alexis Brice MD^{20,21,22}, Guido Alves MD^{6,7,23}, Alastair J. Noyce MRCP^{24,25}, Ole-Bjørn Tysnes MD^{26,27}, Jonathan R. Evans MRCP²⁸, David P. Breen MRCP^{29,30,31}, Karol Estrada PhD¹², Claire E. Wegel MPH³², Fabrice Danjou MD²⁰, David K. Simon MD^{33,34}, Ole A. Andreassen MD^{35,36}, Bernard Ravina MD^{37,38}, Mathias Toft MD^{9,39}, Peter Heutink PhD^{40,41}, Bastiaan R. Bloem MD¹⁰, Daniel Weintraub MD^{42,43}, Roger A. Barker MRCP⁴⁴, Caroline H. Williams-Gray MRCP⁴⁵, Bart P. van de Warrenburg MD¹⁰, Jacobus J. Van Hilten MD⁴⁶, Clemens R. Scherzer MD^{4,5}, Andrew B. Singleton PhD¹, Mike A. Nalls PhD^{1,2}

Affiliations:

¹ Laboratory of Neurogenetics, National Institute on Aging, National Institutes of Health, Bethesda, MD, USA,

² Data Tecnica International, Glen Echo, MD, USA,

³ School of Medicine, Sun Yat-sen University, Guangzhou, Guangdong, China,

⁴ Advanced Center for Parkinson's Disease Research, Brigham and Women's Hospital and Harvard Medical School, Boston, MA, USA,

⁵ Precision Neurology Program, Harvard Medical School, Brigham and Women's Hospital, Boston, MA, USA,

⁶ The Norwegian Centre for Movement Disorders, Stavanger University Hospital, Stavanger, Norway,

⁷ Department of Chemistry, Bioscience and Environmental Engineering, University in Stavanger, Stavanger, Norway,

⁸ Assistance-Publique Hôpitaux de Paris, ICM, INSERM UMRs 1127, CNRS 7225, ICM, Department of Neurology and CIC Neurosciences, Pitié-Salpêtrière Hospital, Paris, France,

⁹ Department of Neurology, Oslo University Hospital, Oslo, Norway,

¹⁰ Department of Neurology, Donders Institute for Brain, Cognition, and Behaviour, Radboud University Medical Centre, Nijmegen, The Netherlands,

¹¹ The Michael J. Fox Foundation for Parkinson's Research, New York, NY, USA,

¹² Translational Genome Sciences, Biogen, Cambridge, MA, USA,

¹³ Department of Neurology University of Pennsylvania, Philadelphia, PA, USA,

¹⁴ Department of Biostatistics and Computational Biology, University of Rochester, Rochester, NY, USA,

¹⁵ Department of Neurology, Center for Health + Technology, University of Rochester, Rochester, NY, USA,

¹⁶ Department of Clinical Neurosciences, University of Cambridge, John van Geest Centre for Brain Repair, Cambridge, UK,

¹⁷ Department of Pathology and Laboratory Medicine, Center for Neurodegenerative Disease Research, Perelman School of Medicine at the University of Pennsylvania, Philadelphia, PA, USA,

¹⁸ Flagship Labs 60 Inc, Cambridge, MA, USA,

¹⁹ Statistical Genetics, Biogen, Cambridge, MA, USA,

²⁰ Institut du cerveau et de la moelle épinière ICM, Paris, France,

²¹ Sorbonne Université SU, Paris, France,

²² INSERM UMR1127, Paris, France,

- 1
2
3 23 Department of Neurology, Stavanger University Hospital, Stavanger, Norway,
4 24 Preventive Neurology Unit, Wolfson Institute of Preventive Medicine, Queen Mary University of London, London,
5 UK,
6 25 Department of Clinical and Movement Neurosciences, UCL Institute of Neurology, London, UK,
7 26 Department of Neurology, Haukeland University Hospital, Bergen, Norway,
8 27 Department of Clinical Medicine, University of Bergen, Bergen, Norway,
9 28 Department of Neurology, Nottingham University NHS Trust, Nottingham, UK,
10 29 Centre for Clinical Brain Sciences, University of Edinburgh, Edinburgh, Scotland,
11 30 Anne Rowling Regenerative Neurology Clinic, University of Edinburgh, Edinburgh, Scotland,
12 31 Usher Institute of Population Health Sciences and Informatics, University of Edinburgh, Edinburgh, Scotland,
13 32 Department of Medical and Molecular Genetics, Indiana University, Indianapolis, IN, USA,
14 33 Department of Neurology, Beth Israel Deaconess Medical Center, Boston, MA, USA,
15 34 Harvard Medical School, Boston, MA, USA,
16 35 NORMENT; Institute of Clinical Medicine, University of Oslo, Oslo, Norway, Norway,
17 36 Division of Mental Health and Addiction, Oslo University Hospital, Oslo, Norway, Norway,
18 37 Voyager Therapeutics, Cambridge, MA, USA,
19 38 Department of Neurology, University of Rochester School of Medicine, Rochester, NY, USA,
20 39 Institute of Clinical Medicine, University of Oslo, Oslo, Norway,
21 40 German Center for Neurodegenerative Diseases-Tuebingen, Tuebingen, Germany,
22 41 HIH Tuebingen, Tuebingen, Tuebingen, Germany,
23 42 Department of Psychiatry, University of Pennsylvania School of Medicine, Philadelphia, PA, USA,
24 43 Department of Veterans Affairs, Philadelphia, PA, USA,
25 44 Department of Clinical Neurosciences and WT-MRC Cambridge Stem Cell Institute, University of Cambridge,
26 Cambridge, UK,
27 45 Department of Clinical Neurosciences, University of Cambridge, Cambridge, UK,
28 46 Department of Neurology, Leiden University Medical Center, Leiden, The Netherlands.
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30
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32
33
34
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36
37

Corresponding author:

38 Mike A. Nalls Ph.D.

39 Laboratory of Neurogenetics, National Institute on Aging, National Institutes of Health

40 35 Convent Drive, Bethesda, MD 20892, USA

41 +1-202-468-1533

42 nallsm@mail.nih.gov

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For Peer Review

Abstract

Background: Previous studies reported various symptoms of Parkinson's disease (PD) associated with sex. Some were conflicting or confirmed only in one study.

Objectives: To examine sex associations to PD phenotypes cross-sectionally and longitudinally in large scale data.

Methods: We tested 40 clinical phenotypes, using longitudinal, clinic-based patient cohorts, consisting of 5,946 patients, with a median follow-up of 3.1 years. For continuous outcomes, we used linear regressions at baseline to test sex-associated differences in presentation, and linear mixed-effects models to test sex-associated differences in progression. For binomial outcomes, we used logistic regression models at baseline and Cox regression models for survival analyses. We adjusted for age, disease duration, and medication use. In the secondary analyses, data from 17,719 PD patients and 7,588 non-PD participants from an online-only, self-assessment PD cohort were cross-sectionally evaluated to determine whether the sex-associated differences identified in the primary analyses were consistent and unique to PD.

Results: Female PD patients had a higher risk of developing dyskinesia early during the follow-up period, with a slower progression in activities of daily living difficulties, and a lower risk of developing cognitive impairments compared with male patients. The findings in the longitudinal, clinic-based cohorts were mostly consistent with the results of the online-only cohort.

Conclusions: We observed sex-associated contributions to PD heterogeneity. These results highlight the necessity of future research to determine the underlying mechanisms and importance of personalized clinical management.

Keyword:

Parkinson's disease; gender; sex; dyskinesias; cognitive impairment; activities of daily livings;

Main text

Introduction

The prevalence of Parkinson's disease (PD) is 1.5–2.0 times higher in men than in women. This discrepancy suggests the potential existence of sex-associated factors that modify the disease process. Identifying the interplay between sex and PD has the potential to assist the development of disease-modifying therapy, inform patient management strategies, and allow the planning of more efficient clinical trials. Researchers have previously investigated sex-associated differences in phenotypes among patients with PD.^{1–3} Male PD patients have been reported to present akinesia/rigid features,⁴ cognitive impairment,^{5–7} daytime sleepiness,⁸ and rapid eye movement (REM) sleep behavioral disorder (RBD) more frequently than female PD patients.^{9,10} In contrast, anxiety disorder/depression^{11–14} and dyskinesia^{11,15–17} were documented to occur more frequently in female PD patients than in male PD patients. However, these studies were generally small in sample size and predominantly performed in a cross-sectional setting.

In this study, we analyzed longitudinal data from 12 PD cohorts, representing 5,946 participants, with a median of 3.1 years of follow-up. This study had two objectives: (1) to identify the baseline differences between men and women, in terms of disease presentation, and (2) to identify the influences of sex on longitudinal symptom trajectory. Further, we analyzed the Fox Insight dataset, an online-only, PD research cohort, to assess whether the observations made using the longitudinal datasets were consistent in an independent dataset. Moreover, by analyzing the data from both PD participants and non-PD participants in the Fox Insight dataset, we were able to evaluate differences in the prevalence of self-reported outcomes between participants with and without PD. This analysis further illustrated that some of the identified differences may be influenced by general differences between males and females, whereas others are disease-specific.

Methods

Participants

12 longitudinal cohorts

We analyzed data from 12 longitudinal PD cohorts, from North America, Europe, and Australia, in this study (Table 1). Among these cohorts, the following four studies enrolled people with early-phase PD who were not being treated at the time of study enrollment (de novo cohorts): Parkinson's Progression Markers Initiative (PPMI), Parkinson Research Examination of CEP-1347 Trial study and its subsequent prospective study (PreCEPT/PostCEPT), the Norwegian ParkWest study (PARKWEST), and Deprenyl and Tocopherol Antioxidative Therapy of Parkinsonism (DATATOP). Other cohorts included Parkinsonism Incidence and Cognitive and Non-motor heterogeneity In Cambridgeshire (PICNICS), National Institutes of Health Exploratory Trials in Parkinson's Disease Large Simple Study 1 (NET_PD_LS1), Drug Interaction With Genes in Parkinson's Disease (DIGPD), Parkinson's Disease Biomarker Program (PDBP), Harvard Biomarkers Study (HBS), ParkFit Study (PARKFIT), Profiling Parkinson's Disease Study (PROPARK), and Udall Centers program (UDALL_PENN). Participants' information was obtained under appropriate written consent and with local institutional and ethical approval. The summary of the designs and inclusion/exclusion criteria applied to these cohorts are documented in the Supplemental Materials. The study protocols were approved at the local institutional review boards and the participants provided written informed consent.

Fox Insight

To evaluate the consistency of results from the longitudinal dataset, we explored an independent dataset, Fox Insight. Fox Insight is an online-only, PD research cohort.¹⁸ The details of the study are available online (<https://foxinsight.michaeljfox.org/>). Individuals, aged 18 or older, with and without PD, were enrolled through in-person referral or online advertisements. The participants provided online informed consent, and self-reported demographic,

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3 characteristics, symptoms, medical history, and PD medication data were collected. Although Fox Insight is a longitudinal
4 study, we analyzed the data cross-sectionally for the present study because the follow-up periods were relatively short
5 (e.g., the median follow-up period was 0.4 years for Non-Motor Symptoms Questionnaire). During the analysis step, we
6 adjusted for age and disease duration. To limit the impacts of the extreme data points, we included participants from the
7 middle 80% of the age distribution and the disease duration distribution (only among PD participants), which excluded
8 any participants younger than the lower 10th percentile (< 46.8 years old) or older than the 90th percentile (> 77.4 years
9 old) and PD patients with a disease duration shorter than one year (10th percentile) and longer than 13.5 years (90th
10 percentile).

21 Measurements

26 Clinical Data Harmonization Among the 12 cohorts

29 Twenty-three measurements, 11 binomial and nine continuous measurements, were analyzed as outcome measures.

31 Binomial outcomes included constipation, mild cognitive impairment, depression, daytime sleepiness, hyposmia,
32 insomnia, wearing off, dyskinesias, RBD, restless-leg syndrome, and modified Schwab and England Activities of Daily
33 Living Scale scores of 70 or lower (SEADL70). Some binomial outcomes had study-specific outcomes, and these criteria
34 are summarized in the Supplemental Materials. For continuous outcomes, we collected the Hoehn and Yahr (HY) stage
35 scale, total and sub-scores for the Unified Parkinson's Disease Rating Scale (UPDRS) or the Movement Disorder Society–
36 revised version (MDS-UPDRS), Mini-Mental State Examination, Montreal Cognitive Assessment (MoCA), and modified
37 Schwab and England Activities of Daily Living Scale (SEADL). UPDRS scores were normalized to the z-values
38 (UPDRS*_scaled).

Fox Insight

The February 2020 data was downloaded from <https://foxden.michaeljfox.org>. The demographic and disease status data were obtained from enrollment and registration questionnaires. For clinical outcomes of interest, we obtained the responses from the following questionnaires: Geriatric Depression Scale (GDS) for depression (score of six or higher);¹⁹ Non-Motor Symptoms Questionnaire (NMS-QUEST) for constipation, depressed mood (Mood depressed) and a proxy for lack of the sense of smell/taste;²⁰ MDS-UPDRS Part II questionnaire; REM Sleep Behavior Disorder Single-Question Screen;²¹ 15-item Penn Parkinson's Disease Daily Activities Questionnaire (PDAQ-15) for cognition-related instrumental functional abilities;²² and Understanding the Impact of Off and On in Parkinson's Patients Questionnaire for dyskinesia and wearing off.

Statistical analysis

Linear and logistic models were used to analyze baseline differences in PD presentation between male and female patients, per cohort. For binomial outcomes, a minimum of 25 outcomes should be observed in the analyzed cohort. Covariates were the linear and square terms of age and disease duration, to adjust for linear and non-linear effects. In addition, we adjusted for levodopa and dopamine agonist use. To test differences in the progression rates among continuous outcomes, we used linear mixed-effects models, with the same covariates as the baseline models and random effects on the individual intercept and slope (change per year). We evaluated sex-associated differences in progression rates by testing the interaction between sex and disease duration. Survival analyses were conducted among those who did not have an outcome at baseline. Cox regression models were used, adjusting for the same covariates as those used in the baseline models. Any outcomes with fewer than 20 events over the follow-up period were not analyzed. The R model statements for these analyses are summarized in the Supplemental Materials.

Then, we combined the cohort-level results with an inverse variance-weighted random-effect model. We focused on robust associations throughout the cohorts; therefore, meta-analyses with p-values less than 0.05 for a test of homogeneity

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3 were excluded from further evaluations. Any associations with a two-sided p-value of 0.05, after Bonferroni-correction
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5 for the number of total analyses, were considered significant.
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9 For the analysis of the Fox Insight dataset, we tested two terms: the mean difference between males and females (main
10 term) and the interaction between sex and disease duration (interaction term). The adjusted covariates were linear and
11 square age, linear and square disease duration, and indicators of levodopa and dopamine agonist usage. We further
12 analyzed the association between sex and outcomes among non-PD participants, adjusted for linear and square age. Then,
13 we conducted a test of homogeneity between sex-associated differences identified among PD cases and non-PD
14 participants, to evaluate whether the sex differences were PD-specific or reflected differences observed in the non-PD
15 population. In the analyses for this dataset, we used a significance level of 0.05 for the raw p-value because the purpose of
16 these analyses was to evaluate consistency with the longitudinal analyses.
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28 All the statistical analyses and drawings were executed using R version 3.6 and python version 3.7. The analysis scripts
29 are available at https://github.com/neurogenetics/PDpheno_by_sex.
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34 Results

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38 The cohort participants are summarized in Table 1. Participants in these cohorts varied in age and PD stage; however,
39 most participants were in relatively early PD phases. The majority of participants were of European descent. Fox Insight
40 included more female participants than the other cohorts, and the ratio of females to males was especially high among
41 non-PD participants, as previously described.²³ Moreover, we did not observe a significant difference in age of diagnosis
42 between the men and the women among each cohort except for Fox Insight, in which the female patients had on average
43 0.61 (SD: 0.12) years younger age of diagnosis than the male patients. Interestingly, the age of non-PD participants in Fox
44 Insight was also younger than male non-PD participants. The younger age of onset may be reflecting different age
45 distributions of the study population by sex in Fox Insight. In the following analyses, we adjusted for age, disease duration
46 and medications.
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5 In total, we conducted 40 meta-analyses, using the clinic-based longitudinal data, three of which were rejected following a
6 test of heterogeneity, with a significance level of 0.05. Using the Bonferroni correction of multiple comparisons, we set
7 our p-value (P) threshold to $0.05/37 = 0.00135$. Among these associations, nine were significant, and the direction and
8 magnitude of associations linked to being female compared with being male are shown in Table 2 and Figure 1/2. (All
9 meta-analysis results can be found in Supplemental Materials).
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18 Female PD patients were less likely to develop cognitive impairments over time {hazard ratio (HR) 0.65 [0.53, 0.79]
19 (mean [95% confidence interval]), $P = 2.1E-5$ } than male PD patients, and an even stronger association was observed
20 when we adjusted for years of education (HR 0.59 [0.48, 0.73], $P = 4.6E-7$, Supplemental Material). This association
21 remain significant when we further adjusted for the baseline MoCA score (HR 0.56 [0.37, 0.86], $P = 0.007$) or the
22 baseline MMSE score (HR 0.67 [0.51, 0.90], $P = 0.007$, Supplemental Material) at the significance level of 0.05.
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28 Additionally, the baseline MoCA scores were higher in female patients (0.63 [0.27, 1.00]) than in male patients, whereas
29 the baseline MMSE score was not significantly different between sexes ($P = 0.97$, Supplemental Materials).
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35 Female patients presented with a higher rate of developing dyskinesia (HR 1.29 [1.16, 1.44]). To assess the impacts of
36 weight, body mass index (BMI) and medication on this association, we conducted ad hoc analyses on a subset of data
37 (PDBP, PPMI, and NET_PD_LS1: 2,281 participants) for which height at baseline, weight at baseline, and medication at
38 visits were recorded. We adjusted the analyses for each of these factors. With the “weight” adjustment, the association
39 was no longer significant ($P = 0.058$), whereas the magnitude of the association became larger when adjusted for levodopa
40 dosages or levodopa equivalent dosages. Adjusting for BMI did not substantially change the magnitude of the association
41 (Beta: from 0.284 to 0.249), and the sex difference remained still significant (Supplemental Materials). Consistent with
42 the higher incidence rate of dyskinesia in female patients, female PD patients in non-de novo cohorts also presented more
43 dyskinesia at baseline than male patients.
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3 Activities of daily living (ADL), captured in the UPDRS Part II, were better in female PD patients than in male PD
4 patients in the baseline analysis (-0.12 [-0.18, -0.06], in the z-score), and the progression rate was slower in female
5 patients than in male patients (-0.14 [-0.20, -0.08] in z-score per year). We added post-hoc analyses of UPDRS Part II
6 scores in the different versions separately. The baseline score differences (female-male) were -0.57 [-1.20, 0.06] (P =
7 0.07) in MDS-UPDRS and -0.52 [-0.82, -0.21] (P = 7.9E-4) in the original UPDRS. The differences in the progression
8 rate were -0.81 [-1.18, -0.44] (P = 1.4E-5) in MDS-UPDRS and -0.43 [-0.71, -0.15] (P = 2.5E-3) in the original UPDRS. A
9 more detailed analysis of the forest plots of the UPDRS Part II scores at baseline showed that the associations between sex
10 and UPDRS Part II were not apparent among the de novo cohorts but, rather, were driven by differences observed in the
11 non-de novo cohorts (Figure 1). Although we did not find significant sex-associated differences in progression rates in the
12 UPDRS Parts I/III/IV, the rate of change for the total UPDRS scores was significantly milder in female patients than in
13 male patients (-0.11 [-0.16, -0.06] per year, in the z-score). In the raw scores, the sex-associated difference (female-male)
14 in rate of change in MDS-UPDRS total score (female-male) was -2.7 [-3.47, -1.95] (P = 2.3E-12) and that of the original
15 UPDRS total score was -0.91 [-1.33, -0.49], (P = 2.66E-05). When only considering the de novo cohorts, similar results
16 were reported for UPDRS part III, with a slower progression rate in female patients than in male patients (-0.14 [-0.21, -
17 0.07] in z-score per year, P = 2.6E-5, Supplemental Materials). This was corresponding to -1.59 [-2.47, -0.71] (P = 4.6E-
18 4) per year difference (female-male) in the rate of change in MDS-UPDRS Part III or -1.01 [-1.78, -0.24] (P = 0.01) per
19 year in the original UPDRS Part III.
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41 Finally, female patients also had lower scores on the UPDRS Part III and the UPDRS total score compared with male
42 patients during the baseline analyses.
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47 When analyzing similar phenotypes within the Fox Insight dataset, we generally confirmed the results of the longitudinal
48 dataset analyses (Table 3). In the Fox Insight dataset analysis, the interaction terms between sex and disease duration
49 indicated the average sex-associated differences in the longitudinal trajectories for the outcomes. For example, a positive
50 association for the interaction between disease duration and PDAQ-15 indicated that the PDAQ-15 scores for female
51 patients were higher than those in male patients (i.e., better cognition-related instrumental functional abilities) among
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3 patients with longer disease durations in the Fox Insight dataset. To illustrate this, we visualized the sex differences,
4 stratified by disease duration (Supplemental Materials). The results are consistent with those for the longitudinal dataset
5 analysis, indicating that female patients had a lower risk of developing cognitive impairments during the disease course.
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7 Similarly, the results from the Fox Insight dataset were consistent with the increased rate of dyskinesia development
8 among female patients compared with male patients, and the lower scores and a slower deterioration rate in UPDRS Part
9 II among female patients, as observed in the longitudinal analyses.
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18 In addition, null differences between male and female patients in the presentation and progression of wearing off,
19 depression, and hyposmia were also supported by the Fox Insight dataset. In contrast, the loss of the sense of smell/taste
20 was significantly more frequently reported in males among the control participants. Having PD might diminish the general
21 sex difference associated with this phenotype.
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26 Single question answers for RBD and some NMSQuest questionnaire questions regarding “difficult to stay awake”
27 (NMSQ_Awake), “difficulty in getting to sleep” (NMSQ_Sleep), “feeling sad, low or blue” (NMSQ_Feel), and
28 NMSQ_Constipation were significantly different according to sex in the Fox Insight dataset. The prevalences of similar
29 outcomes, such as possible RBD, daytime sleepiness, insomnia, depression, and constipation, were not significantly
30 associated with sex in the meta-analyses of 12 longitudinal cohorts. However, the test for these associations gives raw p-
31 values less than 0.05, with the same directions as the Fox Insight results. The primary analyses may not have included
32 large enough sample sizes to detect these associations. All of the sex-phenotype associations among PD participants, not
33 significant in the longitudinal dataset but significant in the Fox Insight dataset, were also significant among non-PD
34 participants. In addition, based on the test of homogeneity between the results from PD and non-PD participants,
35 suggesting that the magnitudes of these sex-associated differences in PD participants did not differ from those in non-PD
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Discussion

We analyzed clinic-based, longitudinal data from 5,946 participants and meta-analyzed the differences in presentation and progression of phenotypes between men and women with PD. We also used web-based, online cohorts and analyzed data from 17,719 PD patients and 7,588 non-PD participants to confirm our results. The results suggested that female PD patients develop dyskinesia early, progress more slowly with respect to ADL restrictions, and are less likely to develop cognitive impairments. For some non-motor symptoms explored in the online questionnaires (such as possible RBD, daytime sleepiness, insomnia, depressive mood, and constipation), we found significant sex-associated differences among PD participants, only in the Fox Insight dataset. These unconfirmed sex-associated differences may not be specific to PD, as we also observed the same associations in the non-PD participants.

Some studies have previously reported that female patients demonstrated an increased risk of developing earlier and more severe dyskinesia^{11,15} and a longer duration of dyskinesia.¹⁶ These reports are consistent with the faster development of dyskinesia among female patients and the large rate of UPDRS Part IV score increases observed in our study. The reasons for this phenomenon are not fully understood, but the relatively higher levodopa dosages with respect to body weight in females may be partially responsible.¹⁷ For example, the commonly used levodopa tablet contains 50 mg or 100 mg levodopa and this is relatively a larger jump for those with less weight, and that may result in stronger treatment for them compared with those with more weight. Our ad hoc analyses also suggested that body weight plays a role in the association between sex and the early development of dyskinesia.

Contradictory results have been reported previously with regards to sex-associated differences in ADL impacts. Two studies evaluated patients who underwent surgical treatment for PD. One study observed no differences in the UPDRS Part II scores between males and females, whereas the other study reported that females had worse scores than males. In these studies, females had a longer duration of disease, which may have affected the results. Another cross-sectional study also reported worse UPDRS Part II scores among female patients.¹¹ They reported that, among the five categories of overall ADL capacity, the two most-severe categories were more frequent among females than males, based on the results

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3 of a chi-squared test, whereas our analyses used UPDRS Part II scores and multivariable regression models. These
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5 different outcome measurements and statistical approaches may account for different results.

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7 The slower development of cognitive declines in female patients was reported by some longitudinal studies.^{5,6,24} The
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9 executive and attention features were primarily affected in PD patients. While Alzheimer's disease, for which women
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11 confer more risk, is emphasized as disability in the memory feature, the executive and attention features are primarily
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13 affected in PD patients. MoCA is more sensitive for detecting dysfunctions in these areas than MMSE,²⁵ and this may be
14
15 one of the reasons that we observed baseline difference in MoCA but not MMSE. In contrast, the longitudinal differences
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17 in the rates of decline for either the MoCA or MMSE were not significantly different between the two sexes, in our data.
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19 Interestingly, MoCA scores were sometimes reported to be higher in healthy aging women than in men.²⁶⁻²⁸ The slower
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21 development of cognitive impairment observed in female patients may reflect their relatively high baseline abilities in the
22
23 areas that are susceptible to PD, although the baseline MoCA score nor MMSE score were able to completely explain the
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25 association between sex and the development of cognitive impairment in the current data.
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31 Several associations that were previously reported were not observed in the current analysis. RBD was reported to be
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33 more prevalent in males with PD than in females with PD,^{9,10} although some studies have disagreed.^{29,30} We were unable
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35 to confirm this association in the current longitudinal dataset. Although the prevalence of possible RBD, as detected by
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37 single-question screening was higher in male patients among the Fox Insight cohort, a similarly increased prevalence in
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39 possible RBD for non-PD male participants makes the PD-specific nature of this association questionable. Female PD
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41 patients were more depressed, according to previous reports.¹¹⁻¹⁴ We were not able to confirm a sex-associated difference
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43 in the presentation or progression of depression, in either the longitudinal data or the Fox Insight dataset. However, female
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45 PD patients expressed a depressive mood more frequently than male patients, in response to the related NMSQuest
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47 question ('feeling sad, 'low' or 'blue') from the Fox Insight dataset. However, the magnitude of the association was not
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49 different between PD and non-PD participants, indicating that the sex difference associated with this outcome may not be
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51 PD-specific. Regarding the NMSQ items evaluated, the similar null results except for NMSQ_Smell were reported
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53 previously in a cross-sectional analysis of de novo PD patients.³¹ Regarding the discrepancy in NMSQ_Smell, it may be
54
55 possible that the sex-difference in reported loss of smell/taste may be only detectable in the de novo PD stage.
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3 The current study has some limitations. Fox Insight is an online-only cohort, which is inherently different from a clinic-
4 based cohort; however, our analyses were mostly consistent across these two different settings. Additionally, because the
5 study participants were almost all of European descent, the generalizability of these observations across different
6 ancestrally distinct groups should be verified. In this study, we focused on the overall associations between sex and
7 phenotypes and did not separate the biological mechanisms from the environmental mechanisms. [For example, the effect
8 of estrogen on PD has been investigated frequently and the conflicting results were reported.](#)³² but we did not collect
9 necessary data to rigorously evaluate the impact of estrogen on the differences. Similarly, we did not have enough data to
10 investigate environmental factors such as smoking, alcohol, diet, physical activity levels, and socio-economic factors. The
11 different distribution of these factors by sex may explain the differences we observed in the current study. Well-designed
12 studies are warranted to dissect the overall differences into each underlying pathway.
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26 Despite some limitations, the current study has some strengths. First, the total number of participants examined in our
27 longitudinal analysis was one of the largest populations studied. Second, although each study had different cohort
28 characteristics, we controlled for heterogeneity and multiple comparisons to detect robust signals. Most of the associations
29 identified between sex and disease presentation and progression were consistent between the longitudinal cohort and
30 analyses performed using the independent Fox Insight dataset. Thus, our results could be generalized to PD patients across
31 various disease stages in different contexts, given the range of studies incorporated. Third, by comparing PD patients with
32 non-PD individuals, we obtained insight into whether sex-associated phenotypes in PD were disease-specific or reflected
33 more general sex differences. Finally, female PD patients have been an underrepresented population in clinical trials.³³
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43 The current work emphasizes the importance of recognizing gender biases when developing treatments for PD in the real
44 world.
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For Peer Review

Figure legends

Figure 1: Forest plots depicting sex differences in outcomes in progression analyses

DATATOP, Deprenyl and Tocopherol Antioxidative Therapy of Parkinsonism; DIGPD, Drug Interaction with Genes in Parkinson's Disease; HBS, Harvard Biomarkers Study; NET-PD_LS1, NIH Exploratory Trials in Parkinson's Disease Large Simple Study 1; PARKFIT, ParkFit study; PARKWEST, The Norwegian ParkWest study; PDBP, Parkinson's Disease Biomarker Program; PICNICS, Parkinsonism Incidence and Cognitive and Non-motor heterogeneity In Cambridgeshire; PPMI, Parkinson's progression markers initiative; PreCEPT_PostCEPT, Parkinson Research Examination of CEP-1347 Trial and PostCEPT; PROPARK, Profiling Parkinson's disease study; and UDALL_PENN, Morris K. Udall Centers for Parkinson's Research.

P, non-adjusted p-values; I^2_{sq} , I^2 statistic; QE_p , test of heterogeneity. “_scaled” scores were normalized (mean 0, standard deviation of 1) to the baseline distributions as the original scores.

Figure 2: Forest plots depicting sex differences in outcomes in baseline analyses

DATATOP, Deprenyl and Tocopherol Antioxidative Therapy of Parkinsonism; DIGPD, Drug Interaction with Genes in Parkinson's Disease; HBS, Harvard Biomarkers Study; NET-PD_LS1, NIH Exploratory Trials in Parkinson's Disease Large Simple Study 1; PARKFIT, ParkFit study; PARKWEST, The Norwegian ParkWest study; PDBP, Parkinson's Disease Biomarker Program; PICNICS, Parkinsonism Incidence and Cognitive and Non-motor heterogeneity In Cambridgeshire; PPMI, Parkinson's progression markers initiative; PreCEPT_PostCEPT, Parkinson Research Examination of CEP-1347 Trial and PostCEPT; PROPARK, Profiling Parkinson's disease study; and UDALL_PENN, Morris K. Udall Centers for Parkinson's Research.

P, non-adjusted p-values; I^2_{sq} , I^2 statistic; QE_p , test of heterogeneity. “_scaled” scores were normalized (mean 0, standard deviation of 1) to the baseline distributions as the original scores.

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Table 1. Baseline characteristics of study cohorts

Cohort	N	f-u, year	European %	Female, %	Stratum	Age, year old	Duration, y	LD, %	DA, %
PPMI	408	7.0	95.1	34.6	Male	62.15 (9.86)	0.53 (0.49)	-	-
					Female	60.76 (9.60)	0.60 (0.63)		
PreCEPT_PostCEPT	390	6.9	97.7	33.8	Male	59.83 (9.51)	0.79 (0.80)	-	-
					Female	60.96 (9.56)	0.84 (0.85)		
PARKWEST	181	5.0	100	37.8	Male	67.82 (9.21)	0.16 (0.10)	-	-
					Female	68.36 (9.10)	0.20 (0.14)*		
DATATOP	796	1.1	97.7	33.7	Male	61.45 (9.35)	1.16 (1.14)	-	-
					Female	60.34 (9.80)	1.10 (1.05)		
PICNICS	122	3.5	98.4	35.2	Male	67.85 (8.40)	0.30 (0.49)	30.4	17.7
					Female	67.93 (10.28)	0.12 (0.50)*	27.9	23.3
NET_PD_LS1	1705	4.0	92.7	35.7	Male	62.07 (9.32)	1.55 (1.08)	57.5	60.4
					Female	61.20 (10.06)	1.54 (1.10)	55.3	63.3
DIGPD	350	3.0	85.8	39.4	Male	61.45 (10.34)	2.55 (1.52)	65.6	77.8
					Female	62.40 (9.61)	2.46 (1.59)	62.3	63.0
PDBP	486	3.0	93.0	39.7	Male	65.03 (9.13)	5.31 (4.74)	81.9	50.2*
					Female	64.87 (8.67)	5.22 (4.78)	76.9	56.8
HBS	482	1.9	96.3	35.3	Male	65.79 (9.67)	4.28 (4.79)	73.7	39.4
					Female	66.60 (9.40)	3.97 (4.30)	70.0	42.4
PROPARK	327	5.0	NA	33.9	Male	59.56 (10.29)	6.48 (5.00)	67.1	69.9
					Female	59.51 (11.63)	6.98 (4.18)	64.0	79.3
PARKFIT	466	2.0	NA	33.3	Male	65.28 (7.41)	4.97 (4.25)	NA	NA
					Female	65.49 (7.60)	5.38 (4.76)		
UDALL_PENN	233	4.0	94.4	30.9	Male	70.53 (7.29)	5.73 (4.96)	84.5	46.0
					Female	70.14 (8.15)	6.64 (5.80)	90.3	56.9
Fox Insight (non-PD)	7588	-	95.8	78.8	Male	63.55 (7.89)	-	-	-
					Female	62.51 (7.39)*			
Fox Insight (PD)	17719	-	96.4	45	Male	66.72 (7.16)	4.61 (3.24)	80.3	29.4
					Female	66.00 (7.10)*	4.50 (3.25)*	76.8*	35.0*

f-u, median follow-up period; European, European descent; Duration, mean disease duration; LD, levedopa use; DA, dopamine agonist use. Age, mean (standard deviation). * p<0.05 for t-test comparing with male vs female.

DATATOP, Deprenyl and Tocopherol Antioxidative Therapy of Parkinsonism; DIGPD, Drug Interaction with Genes in Parkinson's Disease; HBS, Harvard Biomarkers Study; NET-PD_LS1, NIH Exploratory Trials in Parkinson's Disease Large Simple Study 1; PARKFIT, ParkFit study; PARKWEST, The Norwegian ParkWest study; PDBP, Parkinson's Disease Biomarker Program; PICNICS, Parkinsonism Incidence and Cognitive and Non-motor heterogeneity In Cambridgeshire; PPMI, Parkinson's progression markers initiative; PreCEPT_PostCEPT, Parkinson Research Examination of CEP-1347 Trial and PostCEPT; PROPARK, Profiling Parkinson's disease study; and UDALL_PENN, Morris K. Udall Centers for Parkinson's Research.

Table 2. Meta-analysis results for significant associations with sex and phenotypes (reference: male)

Outcome	Beta	SE	P	P-adj	Mean [95%CI]
<u>Progression analysis</u>					
Cognitive_Impairment	-0.436	0.102	2.1E-05	7.7E-4	0.65 [0.53, 0.79] (HR)
Dyskinesia	0.255	0.055	4.1E-06	1.6E-4	1.29 [1.16, 1.44] (HR)
UPDRS2_scaled	-0.139	0.029	1.1E-06	4.1E-5	-0.14 [-0.20, -0.08]
UPDRS_scaled	-0.113	0.025	5.3E-06	2.0E-4	-0.11 [-0.16, -0.06]
<u>Baseline analysis</u>					
Dyskinesia	0.434	0.129	7.3E-04	0.0277	1.54 [1.20, 1.99] (OR)
MoCA	0.634	0.186	6.8E-04	0.0251	0.63 [0.27, 1.00]
UPDRS2_scaled	-0.124	0.031	6.5E-05	0.0024	-0.12 [-0.18, -0.06]
UPDRS3_scaled	-0.114	0.031	2.5E-04	0.0093	-0.11 [-0.17, -0.05]
UPDRS_scaled	-0.107	0.027	6.9E-05	0.0026	-0.11 [-0.16, -0.05]

Progression analyses test the association between incidence rates (binomial) or rates of change per years (continuous) and sex. The models were adjusted for age and disease duration (both linear and square terms), indicators for levodopa and/or agonist usages. “_scaled” scores were normalized (mean 0, standard deviation of 1) to the baseline distributions as the original scores.

SE, standard error; P-adj, Bonferroni adjusted P (raw-P times 37 [the number of multiple-comparison]).

Mean [95%CI], Mean and 95% confidence interval of the difference in each scale. HR, hazard ratio; OR, Odds Ratio, UPDRS, unified Parkinson's disease rating scale; MoCA, Montreal Cognitive Assessment.

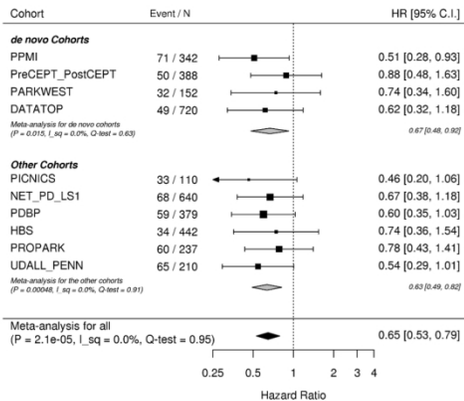
Table 3. Analysis results for sex difference in main term and interaction term with sex and disease duration in replication cohort (reference: male)

Outcome	PD				Control				Test of Homogeneity in main effect			
	Interaction term			Consistency with LT analysis	Main term			Consistency with LT analysis				
	Beta	SE	P		Beta	SE	P			Beta	SE	P
UPDRS2 total	-0.185	0.033	3.2E-08	Consistent	-0.487	0.184	8.1E-03	Consistent				
Cognitive IADL (PDAQ-15)	0.219	0.049	6.8E-06	Consistent	-0.024	0.266	0.928	Consistent	-0.668	0.297	0.025	0.106
Dyskinesia	0.073	0.032	0.024	Consistent	-0.311	0.205	0.129					
Wearing Off	-0.001	0.041	0.977	Consistent	0.259	0.217	0.232	Consistent				
Depression (GDS total >5)	-0.013	0.010	0.203	Consistent	-0.030	0.056	0.596	Consistent	-0.165	0.066	0.012	0.120
pRBD (single question)	-0.003	0.011	0.767	Consistent	-0.698	0.059	1.2E-32		-0.696	0.080	4.6E-18	0.990
NMSQ_Awake	0.001	0.012	0.963	Consistent	-0.216	0.072	2.5E-03		-0.292	0.091	1.3E-03	0.514
NMSQ_Sleep	-0.003	0.011	0.789	Consistent	0.406	0.058	2.4E-12		0.492	0.064	1.4E-14	0.317
NMSQ_Feel	-0.004	0.010	0.676	Consistent	0.384	0.055	4.3E-12		0.342	0.065	1.6E-07	0.628
NMSQ_Constipation	-0.018	0.010	0.072	Consistent	0.173	0.055	1.7E-03		0.310	0.075	3.8E-05	0.143
NMSQ_Smell	-0.011	0.011	0.314	Consistent	0.034	0.059	0.568	Consistent	-0.326	0.111	3.3E-03	4.2E-03

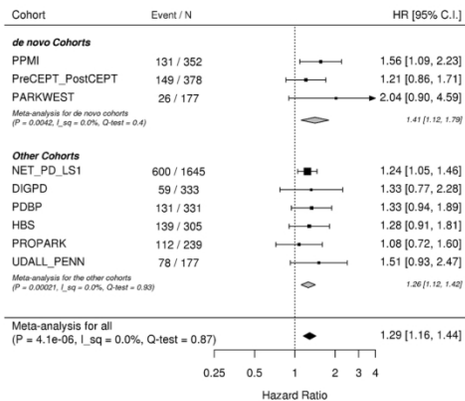
“Main term” is the average difference between the females and the males (reference: males). “Interaction term” is the interaction between disease duration and sex. The adjusted covariates were linear and square age for non-PD participants. For PD participants, linear and square disease duration, and indicators of levodopa and dopamine agonist usage were further adjusted.

SE, standard error; PDAQ-15, the Penn Parkinson's Daily Activities Questionnaire-15 (15-item measure of cognitive instrumental activities of daily living (IADL) for Parkinson's disease patients derived from the original 50-item PDAQ), ranging 0-60 (the lower the worse); Depression, Geriatric Depression Scale score more than 5. Consistency with longitudinal dataset analyses were evaluated for outcomes (Consistency with LT analysis). NMSQ, Non Motor Symptom Questionnaire; NMSQ_ awake: difficult to stay awake; NMSQ_Sleep, difficulty getting sleep at night; NMSQ_Feel, feeling sad, 'low' or 'blue'; NMSQ_Smell, loss or change in your ability to taste or smell.

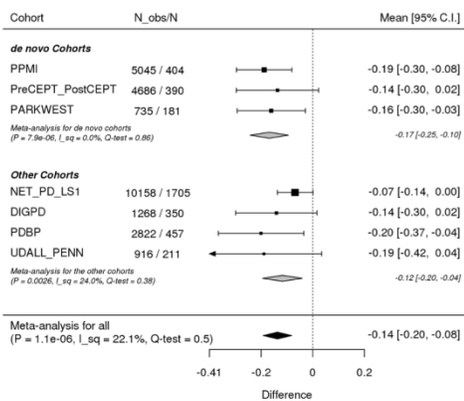
Hazard Ratio (female/male) in Developing Cognitive_Impairment



Hazard Ratio (female/male) in Developing Dyskinesia



Sex Difference (female-male) in Rate of Change in UPDRS2_scaled



Sex Difference (female-male) in Rate of Change in UPDRS2_scaled

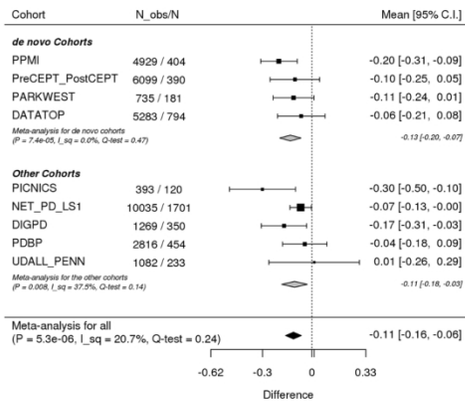


figure 1

152x152mm (300 x 300 DPI)

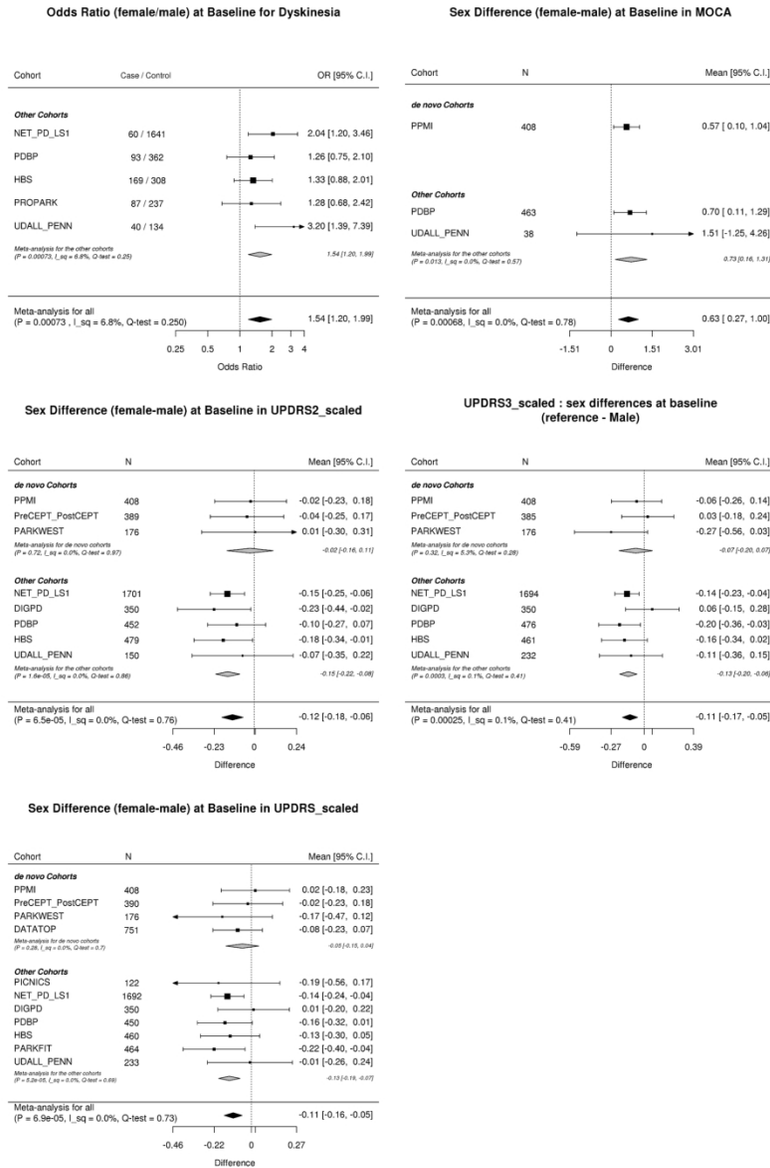


figure 2

152x228mm (300 x 300 DPI)

Supplemental Materials

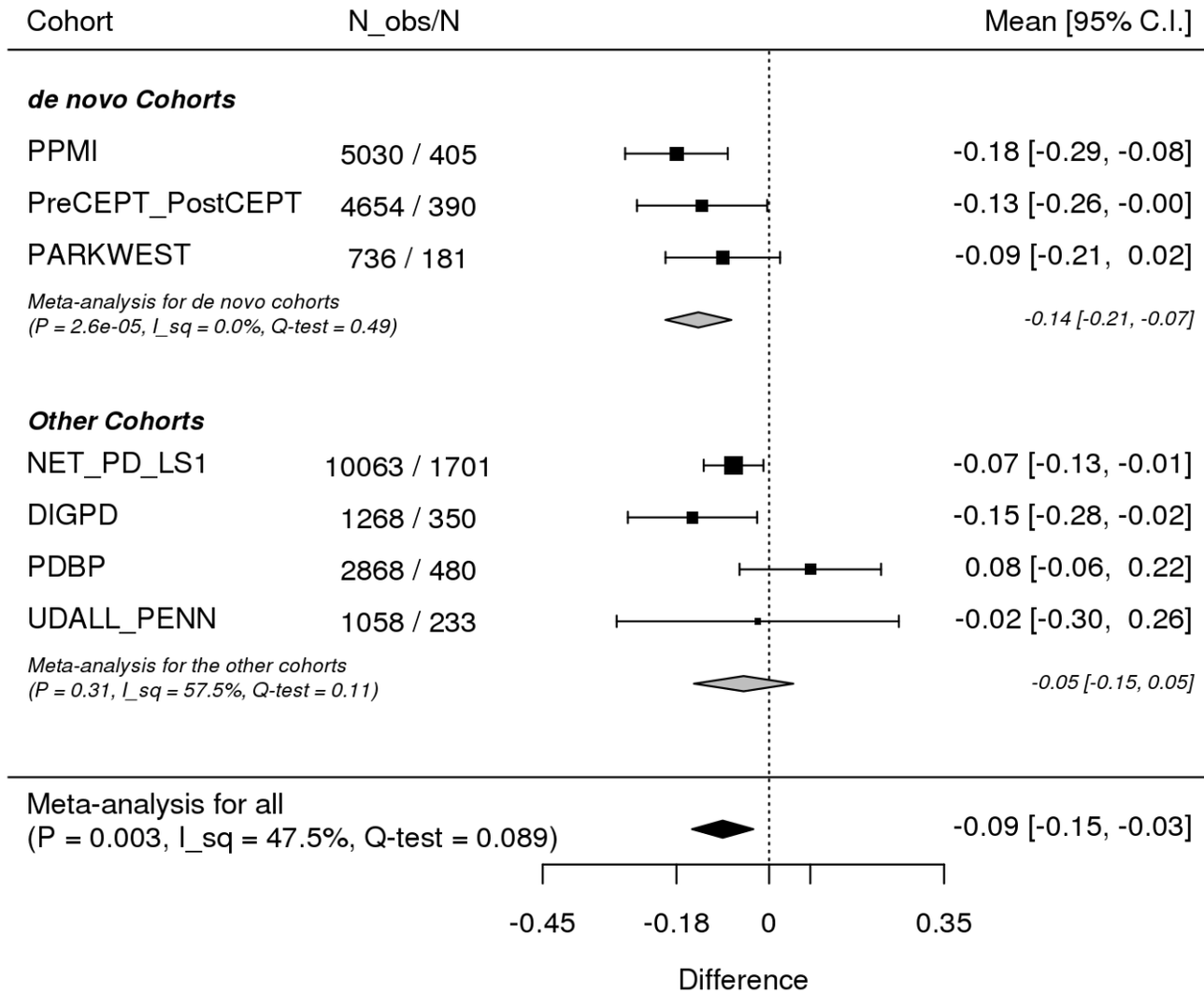
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Supplemental Figures

Forest plots for the sex differences in rate of change in UPDRS part III

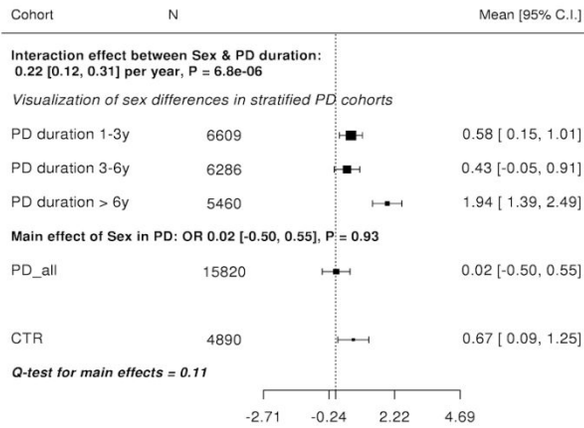
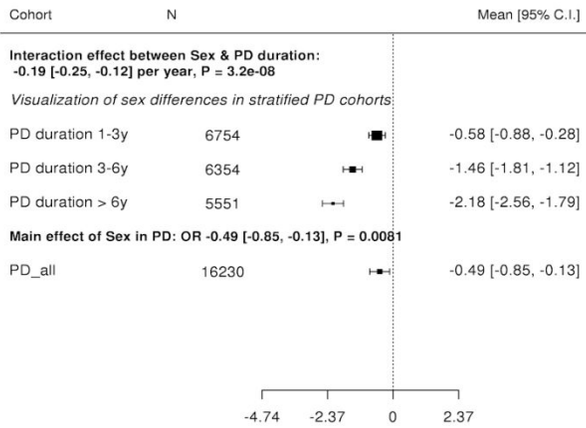
**UPDRS3_scaled : sex differences in rate of change per year
(reference - Male)**



Visualization of the sex differences in FI dataset - 1

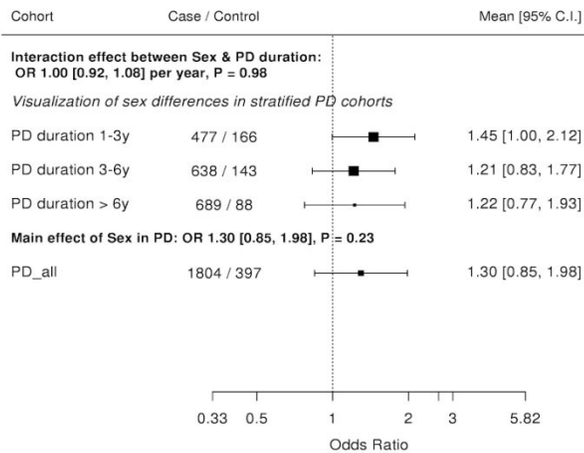
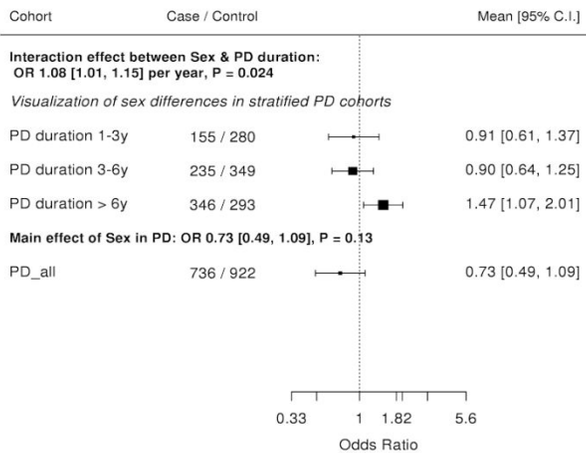
Sex differences in UPDRS2 (continuous)

Sex differences in PDAQ15_total (continuous)



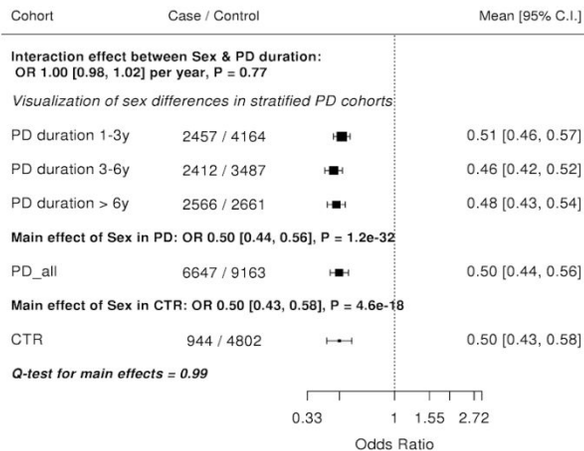
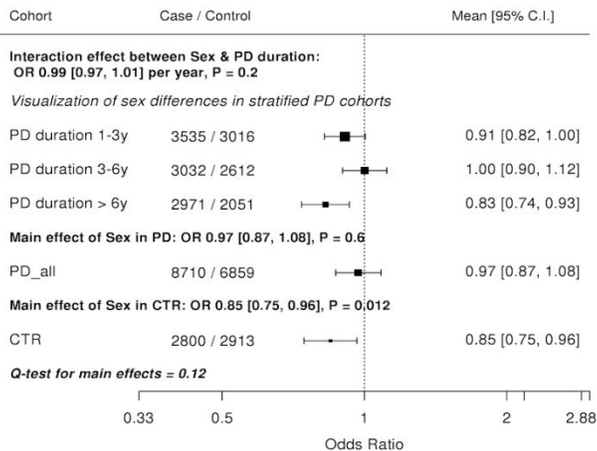
Sex differences in Dyskinesia (binomial)

Sex differences in Wearing_Off (binomial)



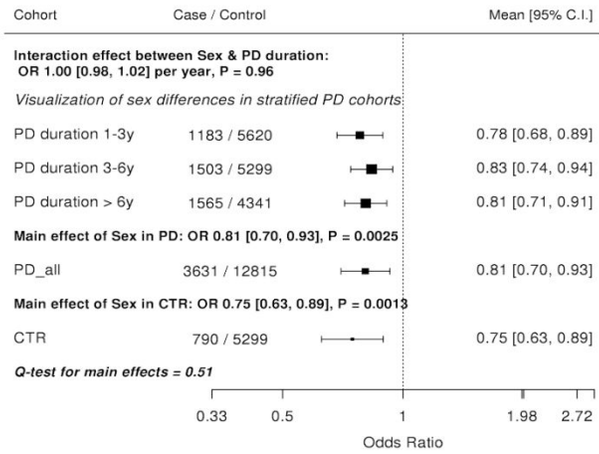
Sex differences in Depression (binomial)

Sex differences in pRBD (binomial)

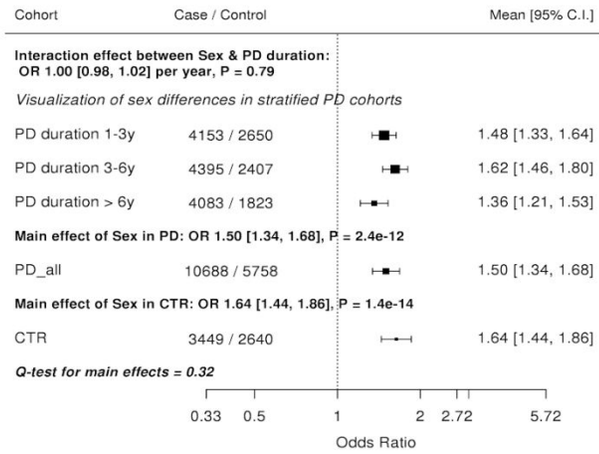


Visualization of the sex differences in FI dataset - 2

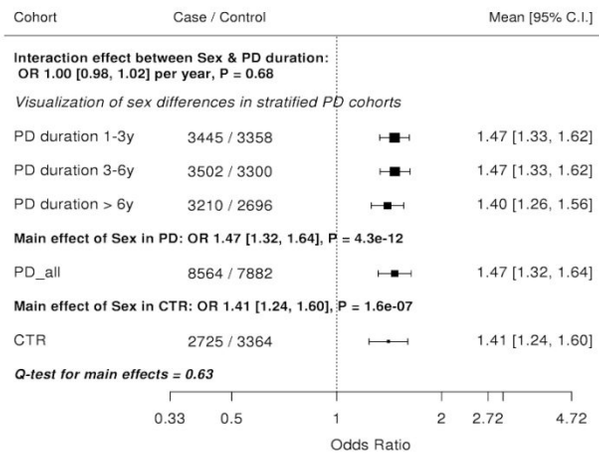
Sex differences in NMSQ_Awake (binomial)



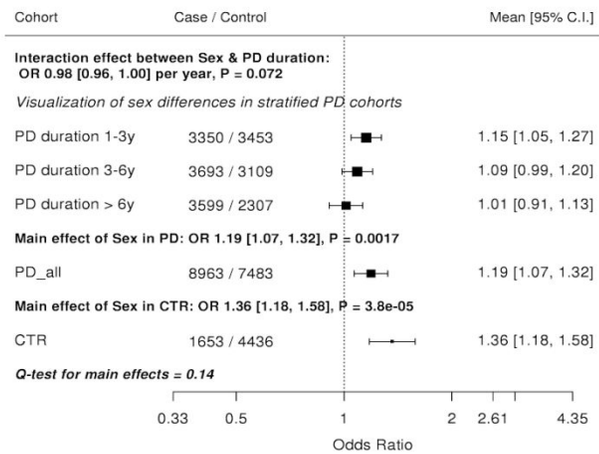
Sex differences in NMSQ_Sleep (binomial)



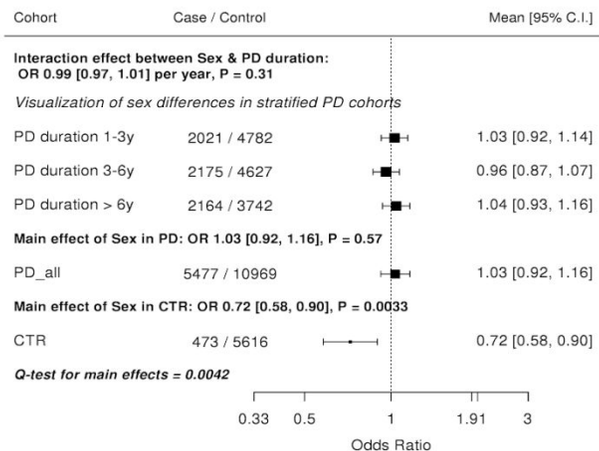
Sex differences in NMSQ_Feel (binomial)



Sex differences in NMSQ_Constipation (binomial)



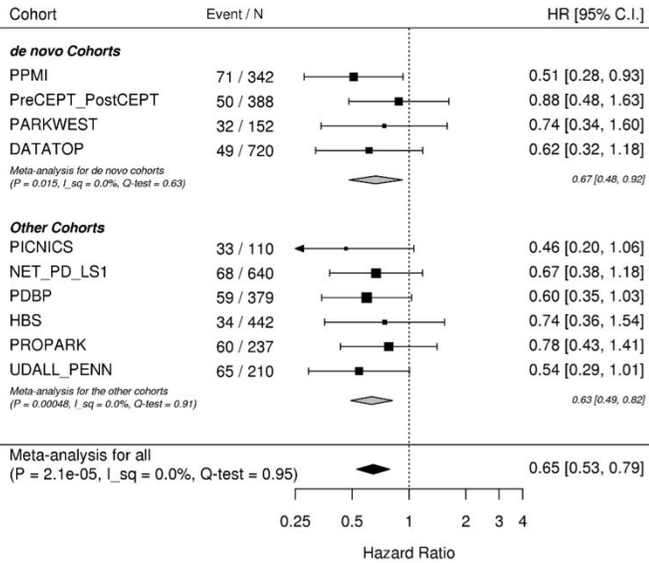
Sex differences in NMSQ_Smell (binomial)



Sex differences in developing cognitive impairment further adjusted for years of education and baseline cognitive test results.

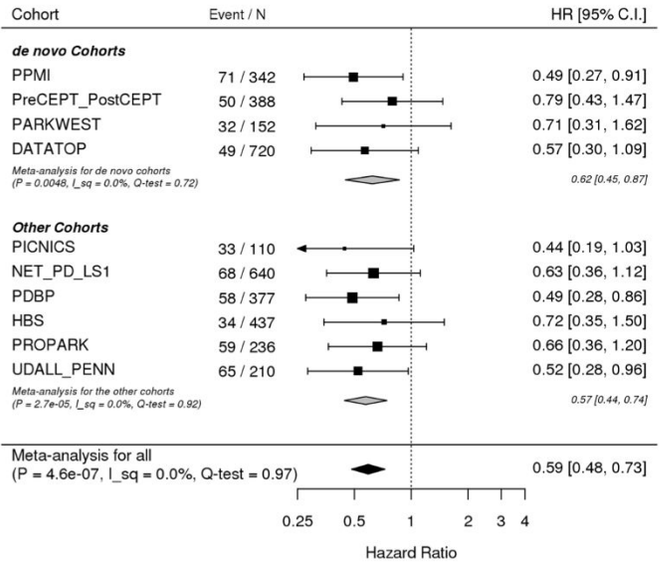
Hazard Ratio (female/male) in Developing Cognitive_Impairment

Base model (adjusted for age, disease duration and medications)



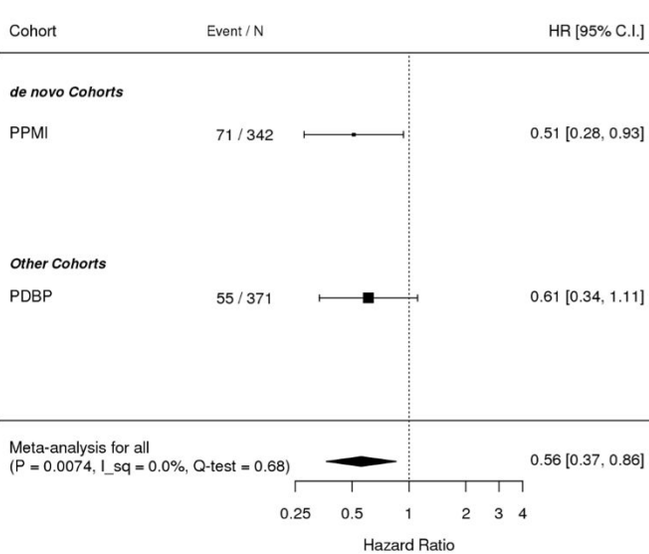
Hazard Ratio (female/male) in Developing Cognitive_Impairment

Base model + Years of education



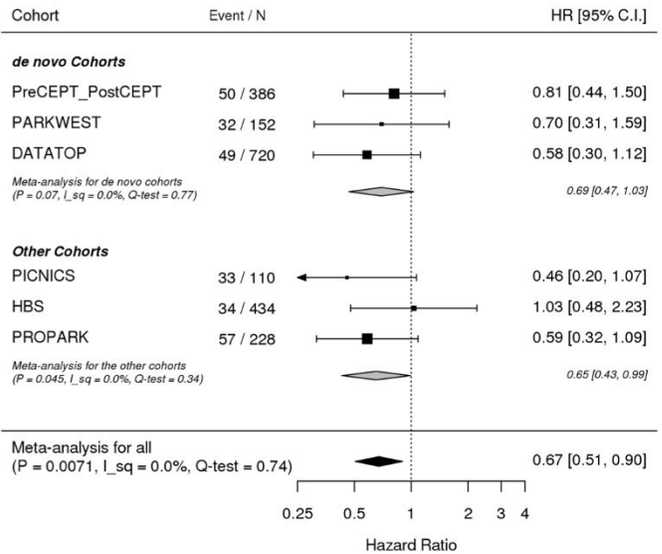
Hazard Ratio (female/male) in Developing Cognitive_Impairment

Base model + Years of education + MoCA at baseline



Hazard Ratio (female/male) in Developing Cognitive_Impairment

Base model + Years of education + MMSE at baseline



Supplemental Tables

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Cohort specific definitions of binomial outcomes

Supplemental Table. Cohort specific definitions of binomial outcomes

	DATATOP	DIGPD	HBS	NET-PD_LSI	ParkFit	ParkWest	PDBP	PICNICS	PPMI	PreCEPT/ PostCEPT	ProPark	Udall
Hyposmia	-	-	-	-	-	-	UPSIT \leq p15	-	UPSIT \leq p15	UPSIT \leq p15	-	UPSIT \leq p15
Cognitive Impairment	MMSE<27	-	MMSE<27	SCOPA_COG<25	MMSE<27	MMSE<27	MoCA<24	MMSE<27	MoCA<24	MMSE<27	MMSE <27	MoCA<24
Wearing off	-	Neurologist diagnosis	any>0 in UPDRS Q36-Q39	UPDRS4 Q39>0	-	UPDRS4 Q39>0	MDS-UPDRS4.3>0	MDS-UPDRS4.3>0	MDS-UPDRS4.3>0	MDS-UPDRS4.3>0	SPES/SCOPA item 20 >0	MDS-UPDRS4.3>0
dyskinesia	AE report	Neurologist diagnosis	any>0 in UPDRS Q32-Q35>0 at any of them	UPDRS4 Q32>0	-	UPDRS4 Q32>0	MDS-UPDRS4 .1>0	MDS-UPDRS4 .1>0	MDS-UPDRS4 .1>0	MDS-UPDRS4 .1>0	SPES/SCOPA item 18 >0	MDS-UPDRS4 .1>0
depression	AE report	Neurologist diagnosis	GDS15 >5	BDI>14	-	UPDRS Q3 \geq 2	HDRS>9	BDI>14 or GDS15>5	GDS15 >5	UPDRS Q3 >0	BDI>14	GDS15>4
restless legs syndrome	-	RLS criteria	Medical history	-	-	-	MSQ3 yes	-	RBDSQ, RLS yes	-	-	-
Constipation	AE report	NMSQuest, Q5 yes	-	-	-	-	MDS-UPDRS1.11>0	MDS-UPDRS1.11>0	MDS-UPDRS1.11>0	MDS-UPDRS1.11>0	SCOPA-AUT item 5>0	-
pRBD	-	-	-	-	-	-	RBDSQ>5	MSQ1 yes	-	RBDSQ>5	-	-
Daytime sleepiness	AE report	Neurologist diagnosis	-	-	-	-	Epworth > 9	Epworth > 9	ESS>9 OR MDS-UPDRS1.8>2	Epworth > 9	MDS-UPDRS1.8>0	SCOPA-SLEEP daytime sleepiness (section D) >4
Insomnia	AE report	Neurologist diagnosis	UPDRS Q41>0	-	-	-	"Do you have problems sleeping?" Yes	UPDRS Q41>0	MDS-UPDRS1.7>0	MDS-UPDRS1.7>0	MDS-UPDRS1.7>0	SCOPA-SLEEP Nighttime (section B) >6

BDI, Beck's Depression Inventory; GDS, Geriatric Depression Scale; HDRS, Hamilton Depression Rating Scale; MMSE, the Mini-Mental State Examination; MoCA Montreal Cognitive Assessment; MSQ, Mayo Sleep Questionnaire; NMS, Non-motor Symptoms Questionnaire; RBD, REM sleep Behavior Disorder; UPDRS Unified Parkinson's Disease Rating Scale (original); MDS-UPDRS, Movement disorder society revised UPDRS version; UPSIT, The University of Pennsylvania Smell Identification Test, 15 percentile (p15) were determined by the sex/age table described in the manual.

Cutoff scores for cognitive impairment were derived from Neurology. 2010 Nov 9;75(19):1717-25. doi: 10.1212/WNL.0b013e3181fc29c9; AE report, report of the symptom as an adverse event during the study;

All meta-analysis results

Supplemental Table. All meta-analysis results

Outcome	Model	Beta	SE	P	Qep	P-adj	Mean [95%CI]
UPDRS2_scaled	mixed	-0.139	0.029	1.1E-06	0.50	4.1E-05	-0.139 [-0.195, -0.083]
Dyskinesia	survival	0.256	0.055	4.1E-06	0.87	1.5E-04	1.291 [1.158, 1.439] (HR)
UPDRS_scaled	mixed	-0.113	0.025	5.3E-06	0.24	2.0E-04	-0.113 [-0.161, -0.064]
Cognitive_Impairment	survival	-0.436	0.102	2.1E-05	0.95	7.7E-04	0.647 [0.529, 0.790] (HR)
UPDRS2_scaled	linear	-0.124	0.031	6.5E-05	0.76	2.4E-03	-0.124 [-0.185, -0.063]
UPDRS_scaled	linear	-0.107	0.027	6.9E-05	0.73	2.6E-03	-0.107 [-0.160, -0.054]
UPDRS3_scaled	linear	-0.114	0.031	2.5E-04	0.41	9.3E-03	-0.114 [-0.175, -0.053]
MoCA	linear	0.634	0.186	6.8E-04	0.78	0.025	0.634 [0.268, 0.999]
Dyskinesia	logistic	0.434	0.129	7.3E-04	0.25	0.027	1.544 [1.200, 1.986] (OR)
UPDRS3_scaled	mixed	-0.092	0.031	3.0E-03	0.09	0.111	-0.092 [-0.153, -0.031]
Daytime_Sleepiness	survival	-0.276	0.095	3.6E-03	0.79	0.132	0.759 [0.630, 0.914] (HR)
UPDRS4_scaled	linear	0.103	0.037	4.8E-03	0.21	0.178	0.103 [0.032, 0.175]
RSL	survival	0.357	0.137	9.2E-03	0.57	0.342	1.429 [1.092, 1.871] (HR)
Insomnia	logistic	0.243	0.096	0.011	0.56	0.413	1.275 [1.057, 1.539] (OR)
MoCA	mixed	0.257	0.102	0.012	0.87	0.429	0.257 [0.057, 0.456]
Constipation	survival	0.227	0.092	0.013	0.48	0.490	1.255 [1.049, 1.503] (HR)
Depression	logistic	0.215	0.087	0.014	0.79	0.505	1.240 [1.045, 1.471] (OR)
Constipation	logistic	0.248	0.107	0.021	0.90	0.782	1.281 [1.038, 1.581] (OR)
UPDRS4_scaled	mixed	0.040	0.017	0.022	0.27	0.821	0.040 [0.006, 0.074]
MMSE	mixed	0.120	0.056	0.033	0.59	1.000	0.120 [0.009, 0.231]
Daytime_Sleepiness	logistic	-0.294	0.140	0.036	0.26	1.000	0.745 [0.566, 0.980] (OR)
Wearing_Off	survival	0.103	0.054	0.057	0.47	1.000	1.109 [0.997, 1.233] (HR)
Wearing_Off	logistic	0.255	0.142	0.072	0.07	1.000	1.291 [0.978, 1.705] (OR)
pRBD	survival	-0.212	0.134	0.115	0.32	1.000	0.809 [0.622, 1.053] (HR)
UPDRS1_scaled	mixed	-0.032	0.020	0.117	0.60	1.000	-0.032 [-0.072, 0.008]
Depression	survival	0.065	0.074	0.386	0.11	1.000	1.067 [0.922, 1.234] (HR)
UPDRS1_scaled	linear	0.026	0.035	0.450	0.45	1.000	0.026 [-0.042, 0.094]
SEADL70	logistic	0.192	0.277	0.488	0.98	1.000	1.212 [0.704, 2.086] (OR)
SEADL	mixed	0.129	0.221	0.560	0.26	1.000	0.129 [-0.305, 0.563]
Hyposmia	logistic	0.105	0.181	0.563	0.44	1.000	1.111 [0.778, 1.585] (OR)
Hyposmia	survival	-0.125	0.223	0.575	0.76	1.000	0.882 [0.570, 1.366] (HR)
HY	mixed	0.007	0.012	0.590	0.98	1.000	0.007 [-0.017, 0.031]
SEADL	linear	-0.100	0.214	0.640	0.65	1.000	-0.100 [-0.519, 0.319]
pRBD	logistic	-0.093	0.242	0.702	0.16	1.000	0.912 [0.567, 1.465] (OR)
Insomnia	survival	0.024	0.076	0.750	0.57	1.000	1.025 [0.882, 1.190] (HR)
RLS	logistic	-0.027	0.137	0.843	0.89	1.000	0.973 [0.743, 1.274] (OR)
MMSE	linear	-0.003	0.070	0.968	0.31	1.000	-0.003 [-0.140, 0.135]
Cognitive_Impairment	logistic	-0.124	0.201	0.538	0.03		0.883 [0.596, 1.311] (OR)
SEADL70	survival	0.058	0.161	0.717	0.01		1.060 [0.773, 1.453] (HR)
HY	linear	0.000	0.026	0.992	0.01		-0.000 [-0.051, 0.051]

SE, standard error; P-adj, Bonferroni adjusted P; QEp, test of homogeneity; Mean [95%CI], Mean and 95% confidence interval of the difference in each scale. HR, hazard ratio; OR, odds ratio.

* Test of homogeneity rejected (<0.05).

UPDRS, Unified Parkinson's Disease Rating Scale; MoCA, Montreal Cognitive Assessment; MMSE, Mini-Mental State examination; RLS, restless legs syndrome; RBD, REM behavior disorder; HY Hoehn and Yahr scale; SEADL Modified Schwab and England Activities of Daily Living Scale.

Associations between sex and dyskinesia in survival models with further adjustment

Supplemental Table. Associations between sex and dyskinesia in survival models with further adjustment

Further adjusted variable	Beta	SE	P	Test of homogeneity
None (Base Model)	0.284	0.082	0.0005	0.37
BMI, kg/m ²	0.249	0.073	0.0007	0.45
Weight at baseline, kg	0.156	0.083	0.0583	0.48
Levodopa dosage, mg/day	0.380	0.117	0.0012	0.15
Levodopa equivalent dose, /day	0.360	0.104	0.0006	0.21

Participants were 2281 people and 845 incidences of dyskinesia were observed during follow-up periods (PPMI, PDBP and NET_PD_LS1.)

The baseline model was adjusted for a linear and a square age; a linear and a square disease duration; a levodopa usage indicator; and a dopamine agonist usage indicator.

R model specifications

Supplemental Table. R model specifications

Study	Analysis	Outcome	Model
PPMI	Baseline analysis	Hyposmia	$Y \sim \text{FEMALE} + \text{Age} + \text{DiseaseDuration} + I(\text{DiseaseDuration}^2) + I(\text{Age}^2)$
PDBP	Baseline analysis	Hyposmia	$Y \sim \text{FEMALE} + \text{Age} + \text{DiseaseDuration} + I(\text{DiseaseDuration}^2) + I(\text{Age}^2)$
UDALL_PENN	Baseline analysis	Hyposmia	$Y \sim \text{FEMALE} + \text{Age} + \text{DiseaseDuration} + \text{LEVODOPA} + \text{AGONIST} + I(\text{DiseaseDuration}^2) + I(\text{Age}^2)$
PARKWEST	Baseline analysis	Cognitive_Impairment	$Y \sim \text{FEMALE} + \text{Age} + \text{DiseaseDuration} + \text{LEVODOPA} + \text{AGONIST} + I(\text{DiseaseDuration}^2) + I(\text{Age}^2)$
DATATOP	Baseline analysis	Cognitive_Impairment	$Y \sim \text{FEMALE} + \text{Age} + \text{DiseaseDuration} + \text{LEVODOPA} + \text{AGONIST} + I(\text{DiseaseDuration}^2) + I(\text{Age}^2)$
NET_PD_LS1	Baseline analysis	Cognitive_Impairment	$Y \sim \text{FEMALE} + \text{Age} + \text{DiseaseDuration} + \text{LEVODOPA} + \text{AGONIST} + I(\text{DiseaseDuration}^2) + I(\text{Age}^2)$
PDBP	Baseline analysis	Cognitive_Impairment	$Y \sim \text{FEMALE} + \text{Age} + \text{DiseaseDuration} + I(\text{DiseaseDuration}^2) + I(\text{Age}^2)$
HBS	Baseline analysis	Cognitive_Impairment	$Y \sim \text{FEMALE} + \text{Age} + \text{DiseaseDuration} + \text{LEVODOPA} + \text{AGONIST} + I(\text{DiseaseDuration}^2) + I(\text{Age}^2)$
PARKFIT	Baseline analysis	Cognitive_Impairment	$Y \sim \text{FEMALE} + \text{Age} + \text{DiseaseDuration} + I(\text{DiseaseDuration}^2) + I(\text{Age}^2)$
PROPARK	Baseline analysis	Cognitive_Impairment	$Y \sim \text{FEMALE} + \text{Age} + \text{DiseaseDuration} + \text{LEVODOPA} + \text{AGONIST} + I(\text{DiseaseDuration}^2) + I(\text{Age}^2)$
NET_PD_LS1	Baseline analysis	Wearing_Off	$Y \sim \text{FEMALE} + \text{Age} + \text{DiseaseDuration} + \text{LEVODOPA} + \text{AGONIST} + I(\text{DiseaseDuration}^2) + I(\text{Age}^2)$
DIGPD	Baseline analysis	Wearing_Off	$Y \sim \text{FEMALE} + \text{Age} + \text{DiseaseDuration} + \text{LEVODOPA} + \text{AGONIST} + I(\text{DiseaseDuration}^2) + I(\text{Age}^2)$
PDBP	Baseline analysis	Wearing_Off	$Y \sim \text{FEMALE} + \text{Age} + \text{DiseaseDuration} + I(\text{DiseaseDuration}^2) + I(\text{Age}^2)$
HBS	Baseline analysis	Wearing_Off	$Y \sim \text{FEMALE} + \text{Age} + \text{DiseaseDuration} + \text{LEVODOPA} + \text{AGONIST} + I(\text{DiseaseDuration}^2) + I(\text{Age}^2)$
PROPARK	Baseline analysis	Wearing_Off	$Y \sim \text{FEMALE} + \text{Age} + \text{DiseaseDuration} + \text{LEVODOPA} + \text{AGONIST} + I(\text{DiseaseDuration}^2) + I(\text{Age}^2)$
UDALL_PENN	Baseline analysis	Wearing_Off	$Y \sim \text{FEMALE} + \text{Age} + \text{DiseaseDuration} + \text{LEVODOPA} + \text{AGONIST} + I(\text{DiseaseDuration}^2) + I(\text{Age}^2)$
NET_PD_LS1	Baseline analysis	Dyskinesia	$Y \sim \text{FEMALE} + \text{Age} + \text{DiseaseDuration} + \text{LEVODOPA} + \text{AGONIST} + I(\text{DiseaseDuration}^2) + I(\text{Age}^2)$
PDBP	Baseline analysis	Dyskinesia	$Y \sim \text{FEMALE} + \text{Age} + \text{DiseaseDuration} + I(\text{DiseaseDuration}^2) + I(\text{Age}^2)$
HBS	Baseline analysis	Dyskinesia	$Y \sim \text{FEMALE} + \text{Age} + \text{DiseaseDuration} + \text{LEVODOPA} + \text{AGONIST} + I(\text{DiseaseDuration}^2) + I(\text{Age}^2)$
PROPARK	Baseline analysis	Dyskinesia	$Y \sim \text{FEMALE} + \text{Age} + \text{DiseaseDuration} + \text{LEVODOPA} + \text{AGONIST} + I(\text{DiseaseDuration}^2) + I(\text{Age}^2)$
UDALL_PENN	Baseline analysis	Dyskinesia	$Y \sim \text{FEMALE} + \text{Age} + \text{DiseaseDuration} + \text{LEVODOPA} + \text{AGONIST} + I(\text{DiseaseDuration}^2) + I(\text{Age}^2)$
PPMI	Baseline analysis	Depression	$Y \sim \text{FEMALE} + \text{Age} + \text{DiseaseDuration} + I(\text{DiseaseDuration}^2) + I(\text{Age}^2)$
PreCEPT_PostCEPT	Baseline analysis	Depression	$Y \sim \text{FEMALE} + \text{Age} + \text{DiseaseDuration} + \text{LEVODOPA} + \text{AGONIST} + I(\text{DiseaseDuration}^2) + I(\text{Age}^2)$
PARKWEST	Baseline analysis	Depression	$Y \sim \text{FEMALE} + \text{Age} + \text{DiseaseDuration} + \text{LEVODOPA} + \text{AGONIST} + I(\text{DiseaseDuration}^2) + I(\text{Age}^2)$
PICNICS	Baseline analysis	Depression	$Y \sim \text{FEMALE} + \text{Age} + \text{DiseaseDuration} + \text{LEVODOPA} + \text{AGONIST} + I(\text{DiseaseDuration}^2) + I(\text{Age}^2)$
NET_PD_LS1	Baseline analysis	Depression	$Y \sim \text{FEMALE} + \text{Age} + \text{DiseaseDuration} + \text{LEVODOPA} + \text{AGONIST} + I(\text{DiseaseDuration}^2) + I(\text{Age}^2)$
DIGPD	Baseline analysis	Depression	$Y \sim \text{FEMALE} + \text{Age} + \text{DiseaseDuration} + \text{LEVODOPA} + \text{AGONIST} + I(\text{DiseaseDuration}^2) + I(\text{Age}^2)$
PDBP	Baseline analysis	Depression	$Y \sim \text{FEMALE} + \text{Age} + \text{DiseaseDuration} + I(\text{DiseaseDuration}^2) + I(\text{Age}^2)$
HBS	Baseline analysis	Depression	$Y \sim \text{FEMALE} + \text{Age} + \text{DiseaseDuration} + \text{LEVODOPA} + \text{AGONIST} + I(\text{DiseaseDuration}^2) + I(\text{Age}^2)$

1	HBS	Survival analysis	Depression	Surv(TSTART, TSTOP, event == 1) ~ FEMALE + Age + DiseaseDuration + LEVODOPA + AGONIST + I(Age^2) + I(DiseaseDuration^2)
2	PROPARK	Survival analysis	Depression	Surv(TSTART, TSTOP, event == 1) ~ FEMALE + Age + DiseaseDuration + LEVODOPA + AGONIST + I(Age^2) + I(DiseaseDuration^2)
3	UDALL_PENN	Survival analysis	Depression	Surv(TSTART, TSTOP, event == 1) ~ FEMALE + Age + DiseaseDuration + LEVODOPA + AGONIST + I(Age^2) + I(DiseaseDuration^2)
4	PPMI	Survival analysis	RLS	Surv(TSTART, TSTOP, event == 1) ~ FEMALE + Age + DiseaseDuration + LEVODOPA + I(Age^2) + I(DiseaseDuration^2)
5	PARKWEST	Survival analysis	RLS	Surv(TSTART, TSTOP, event == 1) ~ FEMALE + Age + DiseaseDuration + LEVODOPA + AGONIST + I(Age^2) + I(DiseaseDuration^2)
6	DIGPD	Survival analysis	RLS	Surv(TSTART, TSTOP, event == 1) ~ FEMALE + Age + DiseaseDuration + LEVODOPA + AGONIST + I(Age^2) + I(DiseaseDuration^2)
7	PDBP	Survival analysis	RLS	Surv(TSTART, TSTOP, event == 1) ~ FEMALE + Age + DiseaseDuration + LEVODOPA + AGONIST + I(Age^2) + I(DiseaseDuration^2)
8	HBS	Survival analysis	RLS	Surv(TSTART, TSTOP, event == 1) ~ FEMALE + Age + DiseaseDuration + LEVODOPA + AGONIST + I(Age^2) + I(DiseaseDuration^2)
9	PPMI	Survival analysis	Constipation	Surv(TSTART, TSTOP, event == 1) ~ FEMALE + Age + DiseaseDuration + I(Age^2) + I(DiseaseDuration^2)
10	PreCEPT_PostCEPT	Survival analysis	Constipation	Surv(TSTART, TSTOP, event == 1) ~ FEMALE + Age + DiseaseDuration + LEVODOPA + AGONIST + I(Age^2) + I(DiseaseDuration^2)
11	PARKWEST	Survival analysis	Constipation	Surv(TSTART, TSTOP, event == 1) ~ FEMALE + Age + DiseaseDuration + LEVODOPA + AGONIST + I(Age^2) + I(DiseaseDuration^2)
12	PICNICS	Survival analysis	Constipation	Surv(TSTART, TSTOP, event == 1) ~ FEMALE + Age + DiseaseDuration + LEVODOPA + AGONIST + I(Age^2) + I(DiseaseDuration^2)
13	DIGPD	Survival analysis	Constipation	Surv(TSTART, TSTOP, event == 1) ~ FEMALE + Age + DiseaseDuration + LEVODOPA + AGONIST + I(Age^2) + I(DiseaseDuration^2)
14	PDBP	Survival analysis	Constipation	Surv(TSTART, TSTOP, event == 1) ~ FEMALE + Age + DiseaseDuration + LEVODOPA + AGONIST + I(Age^2) + I(DiseaseDuration^2)
15	PROPARK	Survival analysis	Constipation	Surv(TSTART, TSTOP, event == 1) ~ FEMALE + Age + DiseaseDuration + LEVODOPA + AGONIST + I(Age^2) + I(DiseaseDuration^2)
16	PPMI	Survival analysis	pRBD	Surv(TSTART, TSTOP, event == 1) ~ FEMALE + Age + DiseaseDuration + I(Age^2) + I(DiseaseDuration^2)
17	PARKWEST	Survival analysis	pRBD	Surv(TSTART, TSTOP, event == 1) ~ FEMALE + Age + DiseaseDuration + LEVODOPA + AGONIST + I(Age^2) + I(DiseaseDuration^2)
18	PDBP	Survival analysis	pRBD	Surv(TSTART, TSTOP, event == 1) ~ FEMALE + Age + DiseaseDuration + LEVODOPA + AGONIST + I(Age^2) + I(DiseaseDuration^2)
19	PPMI	Survival analysis	Daytime_Sleepiness	Surv(TSTART, TSTOP, event == 1) ~ FEMALE + Age + DiseaseDuration + I(Age^2) + I(DiseaseDuration^2)
20	PreCEPT_PostCEPT	Survival analysis	Daytime_Sleepiness	Surv(TSTART, TSTOP, event == 1) ~ FEMALE + Age + DiseaseDuration + LEVODOPA + AGONIST + I(Age^2) + I(DiseaseDuration^2)
21	PARKWEST	Survival analysis	Daytime_Sleepiness	Surv(TSTART, TSTOP, event == 1) ~ FEMALE + Age + DiseaseDuration + LEVODOPA + AGONIST + I(Age^2) + I(DiseaseDuration^2)
22	PICNICS	Survival analysis	Daytime_Sleepiness	Surv(TSTART, TSTOP, event == 1) ~ FEMALE + Age + DiseaseDuration + LEVODOPA + AGONIST + I(Age^2) + I(DiseaseDuration^2)
23	DIGPD	Survival analysis	Daytime_Sleepiness	Surv(TSTART, TSTOP, event == 1) ~ FEMALE + Age + DiseaseDuration + LEVODOPA + AGONIST + I(Age^2) + I(DiseaseDuration^2)
24	PDBP	Survival analysis	Daytime_Sleepiness	Surv(TSTART, TSTOP, event == 1) ~ FEMALE + Age + DiseaseDuration + LEVODOPA + AGONIST + I(Age^2) + I(DiseaseDuration^2)
25	PROPARK	Survival analysis	Daytime_Sleepiness	Surv(TSTART, TSTOP, event == 1) ~ FEMALE + Age + DiseaseDuration + LEVODOPA + AGONIST + I(Age^2) + I(DiseaseDuration^2)
26	PPMI	Survival analysis	Insomnia	Surv(TSTART, TSTOP, event == 1) ~ FEMALE + Age + DiseaseDuration + LEVODOPA + I(Age^2) + I(DiseaseDuration^2)
27	PreCEPT_PostCEPT	Survival analysis	Insomnia	Surv(TSTART, TSTOP, event == 1) ~ FEMALE + Age + DiseaseDuration + LEVODOPA + AGONIST + I(Age^2) + I(DiseaseDuration^2)
28	PARKWEST	Survival analysis	Insomnia	Surv(TSTART, TSTOP, event == 1) ~ FEMALE + Age + DiseaseDuration + LEVODOPA + AGONIST + I(Age^2) + I(DiseaseDuration^2)
29	DATATOP	Survival analysis	Insomnia	Surv(TSTART, TSTOP, event == 1) ~ FEMALE + Age + DiseaseDuration + LEVODOPA + AGONIST + I(Age^2) + I(DiseaseDuration^2)
30	PICNICS	Survival analysis	Insomnia	Surv(TSTART, TSTOP, event == 1) ~ FEMALE + Age + DiseaseDuration + LEVODOPA + AGONIST + I(Age^2) + I(DiseaseDuration^2)
31	DIGPD	Survival analysis	Insomnia	Surv(TSTART, TSTOP, event == 1) ~ FEMALE + Age + DiseaseDuration + LEVODOPA + AGONIST + I(Age^2) + I(DiseaseDuration^2)
32	PDBP	Survival analysis	Insomnia	Surv(TSTART, TSTOP, event == 1) ~ FEMALE + Age + DiseaseDuration + LEVODOPA + AGONIST + I(Age^2) + I(DiseaseDuration^2)
33	HBS	Survival analysis	Insomnia	Surv(TSTART, TSTOP, event == 1) ~ FEMALE + Age + DiseaseDuration + LEVODOPA + AGONIST + I(Age^2) + I(DiseaseDuration^2)

1	PROPARK	Survival analysis	Insomnia	Surv(TSTART, TSTOP, event == 1) ~ FEMALE + Age + DiseaseDuration + LEVODOPA + AGONIST + I(Age^2) + I(DiseaseDuration^2)
2	PPMI	Survival analysis	SEADL70	Surv(TSTART, TSTOP, event == 1) ~ FEMALE + Age + DiseaseDuration + LEVODOPA + I(Age^2) + I(DiseaseDuration^2)
3	PreCEPT_PostCEPT	Survival analysis	SEADL70	Surv(TSTART, TSTOP, event == 1) ~ FEMALE + Age + DiseaseDuration + LEVODOPA + AGONIST + I(Age^2) + I(DiseaseDuration^2)
4	PARKWEST	Survival analysis	SEADL70	Surv(TSTART, TSTOP, event == 1) ~ FEMALE + Age + DiseaseDuration + LEVODOPA + AGONIST + I(Age^2) + I(DiseaseDuration^2)
5	DATATOP	Survival analysis	SEADL70	Surv(TSTART, TSTOP, event == 1) ~ FEMALE + Age + DiseaseDuration + LEVODOPA + AGONIST + I(Age^2) + I(DiseaseDuration^2)
6	NET_PD_LS1	Survival analysis	SEADL70	Surv(TSTART, TSTOP, event == 1) ~ FEMALE + Age + DiseaseDuration + LEVODOPA + AGONIST + I(Age^2) + I(DiseaseDuration^2)
7	DIGPD	Survival analysis	SEADL70	Surv(TSTART, TSTOP, event == 1) ~ FEMALE + Age + DiseaseDuration + LEVODOPA + AGONIST + I(Age^2) + I(DiseaseDuration^2)
8	PDBP	Survival analysis	SEADL70	Surv(TSTART, TSTOP, event == 1) ~ FEMALE + Age + DiseaseDuration + LEVODOPA + AGONIST + I(Age^2) + I(DiseaseDuration^2)
9	UDALL_PENN	Survival analysis	SEADL70	Surv(TSTART, TSTOP, event == 1) ~ FEMALE + Age + DiseaseDuration + LEVODOPA + AGONIST + I(Age^2) + I(DiseaseDuration^2)

UPDRS, unified parkinson's disease rating scale; MOCA, Montreal Cognitive Assessment; MMSE, Mini-Mental State examination; RLS, restless legs syndrome; RBD, REM behavior disorder; HY Hoehn and Yahr scale; SEADL Modified Schwab and England Activities of Daily Living Scale; TSTART, TSTOP, survival observation (start and stop).

For Peer Review

Supplemental documents about the longitudinal cohorts

Descriptions

Deprenyl and Tocopherol Antioxidative Therapy of Parkinsonism (DATATOP) was a randomized clinical trial conducted between September 1987 and November 1989 at 28 sites across US and Canada. The primary objective was to test the efficacy of deprenyl and/or tocopherol. 800 patients with Parkinson's disease diagnosed within 5 years and not requiring symptomatic treatment were observed for up to 2 years.¹ The study was supported by a Public Health Service grant (NS24778) from the National Institute of Neurological Disorders and Stroke; by grants from the General Clinical Research Centers Program of the National Institutes of Health at Columbia University (RR00645), the University of Virginia (RR00847), the University of Pennsylvania (RR00040), the University of Iowa (RR00059), Ohio State University (RR00034), Massachusetts General Hospital (RR01066), the University of Rochester (RR00044), Brown University (RR02038), Oregon Health Sciences University (RR00334), Baylor College of Medicine (RR00350), the University of California, San Diego (RR00827), Johns Hopkins University (RR00035), the University of Michigan (RR00042), and Washington University (RR00036); the Parkinson's Disease Foundation at Columbia-Presbyterian Medical Center, New York; the National Parkinson Foundation, Miami; the Parkinson Foundation of Canada, Toronto; the United Parkinson Foundation, Chicago; the American Parkinson's Disease Association, New York; and the University of Rochester, Rochester, N.Y.

Drug Interaction with Genes in Parkinson's Disease (DIGPD) is a cohort with 413 patients with Parkinson's disease diagnosed by UK Parkinson's disease society brain bank clinical diagnostic (UKPDSBB) criteria with disease duration less than 5 years at the entry.² It is an ongoing study since 2009, and the participants are followed for up to 7 years at eight sites in France. (Corvol et al., in press in Neurology). DNA samples were collected from all of them. DIGPD is sponsored by Assistance Publique Hôpitaux de Paris, funded by a grant from the French Ministry of Health (PHRC 2008, AOM08010) and a grant from the Agence Nationale pour la Sécurité des Médicaments (ANSM 2013).

Harvard Biomarkers Study (HBS) is a longitudinal case-control study. More than 2,700 individuals with early-stage PD, patients with memory impairment, and controls without neurological disease were enrolled and longitudinally phenotyped since 2008.³ HBS was supported by the Harvard NeuroDiscovery Center, MJFF, NINDS U01NS082157, U01NS100603, and the Massachusetts Alzheimer's Disease Research Center NIA P50AG005134.

1 NIH Exploratory Trials in Parkinson's Disease Large Simple Study 1 (NET-PD LS1) was a randomized study conducted
2 between March 2007 and September 2013 to determine if the nutritional supplement creatine slows the clinical
3 progression of Parkinson's disease over time. 1741 patients from 50 sites in the US and Canada participated.⁴ They were
4 within 5 years from diagnosis. The plan was for them to be followed for at least 5 years, but the study ended early for
5 futility based on an interim analysis at which point the median follow-up time was 4 years. Financial support for the LS-1
6 study was provided by National Institute of Neurological Disorders and Stroke (NINDS) grant U01NS43128.

7 Oslo PD study[Citation error] (Oslo) is an ongoing study since 2007, with 317 patients diagnosed with ULPDSBB criteria
8 with modification of allowing family history. The participants are being followed up to 6 years in prospective (30 years in
9 retrospective) at Oslo University Hospital in Norway.⁵ Oslo PD is supported by the Research Council of Norway and
10 South-Eastern Norway Regional Health Authority.

11 ParkFit cohort was originally a randomized trial evaluating a multifaceted behavioural change programme to increase
12 physical activities in patients with Parkinson's disease.⁶ The study conducted from September 2008 to February 2012 at a
13 single center in the Netherlands, with 586 patients with Parkinson's disease diagnosed by UKPDSBB, with Hoehn Yahr
14 stage 3 or lower, and with sedentary lifestyle at the entry. They were followed up for 2 years. The primary objective was
15 concluded as not significant⁶. ParkFit is supported by ZonMw (the Netherlands Organization for Health Research and
16 Development (75020012)) and the Michael J Fox Foundation for Parkinson's research, VGZ (health insurance company),
17 GlaxoSmithKline, and the National Parkinson Foundation.

18 The Norwegian ParkWest study (ParkWest) is an ongoing prospective longitudinal multicenter cohort study of patients
19 with incident Parkinson's disease from Western and Southern Norway, designed to study the incidence, neurobiology and
20 prognosis of PD.⁷ Between November 1st 2004 and 31st of August 2006, all new cases of Parkinson Disease within the
21 study area (Sogn and Fjordane, Hordaland, Rogaland and Aust-Agder) were recruited, and since the start of the study 212
22 of these patients and their age-/sex-matched control group were followed. The Norwegian ParkWest study is supported by
23 the Research Council of Norway, the Western Norway Regional Health Authority, Stavanger University Hospital
24 Research Funds, and the Norwegian Parkinson's Disease Association.

25 The National Institute of Neurological Disorders and Stroke (NINDS) Parkinson's Disease Biomarker Program (PDBP) is
26 aiming to discover new diagnostic and progression biomarkers for Parkinson's disease.⁸ It is a combined cohort of 9
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1 PDBP-funded research studies. The members have various stages of Parkinson's disease and recruited throughout the
2 United States.

3
4 Parkinsonism: Incidence and Cognitive and Non-motor heterogeneity In Cambridgeshire (PICNICS) is a population-based
5 longitudinal study of 282 incident PD cases recruited between 2008 and 2013 with ongoing follow-up at 18 month
6 intervals.^{9,10} PD cases were diagnosed based on the UKPDSBB criteria, and followed up at a single center in the UK.
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8 PICNICS has received funding from the Cure Parkinson's Trust, the Van Geest Foundation and is supported by the
9
10 National Institute of Health Research Cambridge Biomedical Research Centre.
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15 Parkinson's progression markers initiative (PPMI) is an ongoing study started on July 2010, enrolling 424 patients with
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17 Parkinson's disease diagnosed within 2 years from the study entry date.¹¹ The study sites are located in 33 sites across the
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19 US, Europe, Israel and Australia¹¹. PPMI is supported by the Michael J Fox Foundation for Parkinson's Research.
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22 Parkinson Research Examination of CEP1348 Trial (PreCEPT) is a clinical trial of the mixed lineage kinase inhibitor
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24 CEP-1357,4 sponsored by Cephalon, Inc. (West Chester, PA) and H. Lundbeck A/S (Valby-Copenhagen, Denmark). The
25
26 study was conducted at 65 sites in North America. The trial enrolled 806 early, untreated PD patients within one year
27
28 from the onset. The original trial was started in April 2002 and terminated in August 2005 due to the futility, but the
29
30 participants were continuously followed-up in the prospective observational study (PostCEPT).¹²
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33 The studies were funded by NINDS 5U01NS050095-05, Department of Defense Neurotoxin Exposure Treatment
34
35 Parkinson's Research Program. Grant Number: W23RRYX7022N606, the Michael J Fox Foundation for Parkinson's
36
37 research, Parkinson's Disease Foundation, Lundbeck Pharmaceuticals, Cephalon Inc, Lundbeck Inc, John Blume
38
39 Foundation, Smart Family Foundation, RJG Foundation, Kinetics Foundation, National Parkinson Foundation, Amarin
40
41 Neuroscience LTD, CHDI Foundation Inc, National Institutes of Health (NHGRI, NINDS), Columbia Parkinson's Disease
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43 Research Center.
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46 Profiling Parkinson's disease study (ProPark) is an ongoing study started from May 2003. Initially, 420 patients recruited
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48 in several sites in the Netherlands by March 2006.¹³ Patients were diagnosed with UKPDSBB criteria and in various
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50 disease durations at the enrollment. They are evaluated annually with the SCOPA scale. This study is funded by the
51
52 Alkemade-Keuls Foundation, Stichting Parkinson Fonds, Parkinson Vereniging, The Netherlands Organisation for Health
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54 Research and Development.
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Study investigators

Harvard Biomarkers Study. Co-Directors: Harvard NeuroDiscovery Center: Clemens R. Scherzer, Bradley T. Hyman, Charles G. Jennings; Investigators and Study Coordinators: Harvard NeuroDiscovery Center: Yuliya Kuras, Daly Franco, Frank Zhu; Brigham and Women's Hospital: Lewis R. Sudarsky, Michael T. Hayes, Chizoba C. Umeh, Reisa Sperling; Massachusetts General Hospital: John H. Growdon, Michael A. Schwarzschild, Albert Y. Hung, Alice W. Flaherty, Deborah Blacker, Anne-Marie Wills, U. Shivraj Sohur, Vivek K. Unni, Nicta I. Mejia, Anand Viswanathan, Stephen N. Gomperts, Vikram Khurana, Mark W. Albers, Maria Allora-Palli, Alireza Atri, David Hsu, Alexandra Kimball, Scott McGinnis, Nutan Sharma, John Becker, Randy Buckner, Thomas Byrne, Maura Copeland, Bradford Dickerson, Matthew Frosch, Theresa Gomez-Isla, Steven Greenberg, James Gusella, Julius Hedden, Elizabeth Hedley-Whyte, Keith Johnson, Raymond Kelleher, Aaron Koenig, Maria Marquis-Sayagues, Gad Marshall, Sergi Martinez-Ramirez, Donald McLaren, Olivia Okereke, Elena Ratti, Christopher William, Koene Van Dij, Shuko Takeda, Anat Stemmer-Rachaminov, Jessica Kloppenburg, Catherine Munro, Rachel Schmid, Sarah Wigman, Sara Wlodarczyk; University of Ottawa: Michael G. Schlossmacher; Scientific Advisory Board: Massachusetts General Hospital: John H. Growdon; Brigham and Women's Hospital: Dennis J. Selkoe, Reisa Sperling; Harvard School of Public Health: Alberto Ascherio; Data Coordination: Harvard NeuroDiscovery Center: Thomas Yi, Massachusetts General Hospital: Joseph J. Locascio, Haining Li; Biobank Management Staff: Harvard NeuroDiscovery Center: Gabriel Stalberg, Zhixiang Liao.

Parkinson Study Group DATATOP investigators: *Steering Committee* — Ira Shoulson, M.D. (principal investigator), University of Rochester, Rochester, N.Y.; Stanley Fahn, M.D. (co-principal investigator), Columbia–Presbyterian Medical Center, New York; David Oakes, Ph.D. (chief biostatistician, 1987 to present), University of Rochester; Charles Odoroff Ph.D. (deceased) (chief biostatistician, 1985–1987), University of Rochester; Anthony Lang, M.D., Toronto Western Hospital, Toronto; J. William Langston, M.D., California Parkinson's Foundation, San Jose, Calif.; Peter LeWitt, M.D., Sinai Hospital, Detroit; Warren Olanow, M.D., University of South Florida, Tampa; John B. Penney, M.D. (deceased), University of Michigan, Ann Arbor; and Caroline Tanner, M.D., Rush–Presbyterian–St. Luke's Medical Center, Chicago.

Participating Investigators — William Koller, M.D. (deceased), University of Kansas, Kansas City; Warren Olanow, M.D., University of South Florida; Robert Rodnitzky, M.D., University of Iowa, Iowa City; J. Stephen Fink, M.D., Ph.D. (deceased), and John H. Growdon, M.D., Massachusetts General Hospital, Boston; George Paulson, M.D., Ohio State

University, Columbus; Roger Kurlan, M.D., University of Rochester; Joseph H. Friedman, M.D., Roger Williams General Hospital, Providence; Stephen Gancher, M.D., and John Nutt, M.D., Oregon Health Sciences University, Portland; Ali H. Rajput, M.D., University of Saskatchewan, Saskatoon; James Bennett, M.D., Ph.D., and George F. Wooten, M.D., University of Virginia, Charlottesville; Peter LeWitt, M.D., Sinai Hospital, Detroit; Christopher Goetz, M.D., Caroline Tanner, M.D., Kathleen Shannon, M.D., and Harold Klawans, M.D. (deceased), Rush–Presbyterian–St. Luke's Medical Center, Chicago; Oksana Suchowersky, M.D., University of Calgary, Calgary, Alta.; Mitchell Brin, M.D., and Susan Bressman, M.D., Columbia–Presbyterian Medical Center, New York; William J. Weiner, M.D. (deceased), and Juan Sanchez-Ramos, M.D., Ph.D., University of Miami, Miami; Joseph Jankovic, M.D., Baylor College of Medicine, Houston; John B. Penney, M.D. (deceased), University of Michigan, Ann Arbor; Anthony Lang, M.D., Toronto Western Hospital, Toronto; Margaret Hoehn, M.D. (deceased), St. Luke's Hospital, Denver; James Tetrud, M.D., California Parkinson's Foundation, San Jose; J. David Grimes, M.D. (deceased), Ottawa Civic Hospital, Ottawa, Ont.; Ronald Pfeiffer, M.D., University of Nebraska and Creighton University, Omaha; Cliff Shults, M.D. (deceased), and Leon Thal, M.D. (deceased), University of California, San Diego; Serge Gauthier, M.D., Montreal General Hospital and McGill University, Montreal; Lawrence I. Golbe, M.D., University of Medicine and Dentistry of New Jersey, New Brunswick; Joel S. Perlmutter, M.D., Washington University, St. Louis; Hamilton Moses III, M.D., Johns Hopkins University, Baltimore; and Howard I. Hurtig, M.D., and Matthew Stern, M.D., The Graduate Hospital, Philadelphia.

Site Coordinators — Ruth Barter, R.N., and Bridget Vetere-Over-field, R.N., Kansas City, Kans.; Lisa Gauger, B.A., and Terresita Malapira, R.N., Tampa, Fla.; Judith Dobson, R.N., Iowa City, Iowa; Susan Atamian, R.N., Marsh Tennis, R.N., Jennifer B. Cohen, B.A., and Gena Desclos, B.A., Boston; Lena Denio, M.T., Steven Huber, Ph.D., and Teresa Woike, R.N., Columbus, Ohio; Jill Behr, R.N., M.S., and Irenita Gardiner, R.N., Rochester, N.Y.; Margaret Lannon, R.N., M.S., Providence, R.I.; Julie Carter, R.N., and Susanne Northrup, Portland, Ore.; Bernice Kanigan, R.N., Saskatoon, Sask.; Margaret Turk, R.N., M.S., and Elke Landow, R.N., Charlottesville, Va.; Patricia Schlick, R.N., and Kathie Mistura, R.N., Detroit; V. Susan Carrol, R.N., M.S., Jeana Thelen, R.N., and Joan Lechner, Chicago; Carol Demong, R.N., Calgary, Alta.; Linda Winfield, R.N., and Carol Moskowitz, R.N., New York; Angela Ingenito, R.N., Carol Sheldon, R.N., and Lisa Cornelius, B.A., Miami; Dorothy Heiberg, R.N., Houston; Jan Brady, R.N., M.S., Ann Arbor, Mich.; Catherine Kierans, R.N., M.A., and Loretta Bell-Scantlebury, R.N., Toronto; Helena Weber, M.T., M.A., Denver; Deborah Savoini, R.N., Paula Lewis, R.N., and S. Jerome Kutner, Ph.D., San Jose, Calif.; Peggy Gray, R.N., Ottawa, Ont.; Ruth Hofman,

1 R.N., and Carolyn Glaeske, R.N., Omaha; Mary Margaret Pay, R.N., and David Salmon, Ph.D., San Diego; Frances
2 McFaul, R.N., and Donna Amyot, R.N., Montreal; Mary Bergen, R.N., New Brunswick, N.J.; Lori McGee-Minnich, R.N.,
3 St. Louis; Patricia O'Donnell, R.N., M.S., Baltimore; and Susie Ferrise, R.N., and Kathy Shallow, B.A., Philadelphia.
4
5
6 *Coordination and Data Center* (University of Rochester Medical Center, Suite 160, 1325 Mt. Hope Ave., Rochester, NY
7 14620) — Rita M. Pelusio, M.S.Ed. (program manager); Alice Rudolph, Ph.D. (chief study coordinator); Peter Como,
8 Ph.D. (neuropsychology consultant); Charlyne Miller, R.N., M.S. (nurse clinician); Michael Linsner, B.S., Joseph
9 Connorton, B.A., and Judith Nusbaum, B.A. (analyst programmers); Carrie Irvine, B.S., and Constance Orme, B.A.
10 (information analysts); Ruth Nobel, Deborah Baker, Donna LaDonna, Mary Ellen Rothfuss, and Lynn Doerr (deceased)
11 (secretarial staff); Jacqueline Wendel, B.T. (CLINFO manager); and Belinda Rodriguez, Virginia Collins, Scott Dalston,
12 and Paul Bivrell (student clerks).
13
14
15 *Biostatistics Center* (Division of Biostatistics, University of Rochester Medical Center, Rochester, NY 14642) — Charles
16 Odoroff, Ph.D. (deceased), and David Oakes, Ph.D. (chief biostatisticians); Michael McDermott, Ph.D., and Shirley
17 Eberly, M.S. (biostatisticians); Sandra Plumb, B.S. (lead programmer); Arthur Watts, B.S., Lori Yorkey, B.A., Anna
18 Choi, B.A., and Karen Gerwitz, B.S. (analyst programmers).
19
20
21 *Pharmacy Center* (Strong Memorial Hospital, Rochester, NY 14642) — Paul Evans, R.Ph. (chief pharmacist); Lori
22 Dellapena and Verna Singletary (pharmacy technicians).
23
24
25 *Safety Monitoring Committee* (Rochester, N.Y.) — Robert Herndon, M.D. (chair, January 1, 1987–June 30, 1988), Pierre
26 Tariot, M.D. (chair, July 1, 1988 to present), Edward Bell, Pharm.D., Robert C. Griggs, M.D., W. Jackson Hall, Ph.D.,
27 Sandra Plumb, B.S. (lead programmer), and Arthur Watts, B.S. (analyst programmer).
28
29
30 *Scientific Advisory Committee* — C. David Marsden, D.Sc. (deceased) (chair), London; Gerald Cohen, Ph.D. (deceased),
31 Joseph Fleiss, Ph.D. (deceased), and Richard Mayeux, M.D., New York; and Laurence Jacobs, M.D., and Arthur J. Moss,
32 M.D., Rochester, N.Y.
33
34
35 *Clinical Trials Monitoring Committee* (National Institute of Neurological Diseases and Stroke) — Emanuel M. Stadlan,
36 M.D. (chair), Bethesda, Md.; Milton Alter, M.D. (deceased), Philadelphia; Jesse Cedarbaum, M.D., White Plains, N.Y.;
37 Jonas Ellenberg, Ph.D., Bethesda, Md.; and Robert Kibler, M.D., Atlanta.
38
39
40
41
42
43
44
45
46
47
48
49
50
51
52
53
54
55
56
57
58
59
60

1 *Assay Standards Committee* — Robert Roth, Ph.D. (chair), New Haven, Conn.; Harvey Cohen, M.D., Ph.D., Rochester,
2 N.Y.; Matthew Galloway, Ph.D., Detroit; Ian Irwin, Ph.D., San Jose, Calif.; Peter LeWitt, M.D., Detroit; Govind
3
4 Vatassery, Ph.D., Minneapolis.
5

6 *Neuropsychological Testing Committee* — Richard Mayeux, M.D. (chair), New York; Peter Como, Ph.D., Rochester,
7 N.Y.; Jean St. Cyr, Ph.D., Toronto; Yaakov Stern, Ph.D., and Janet Williams, D.S.W., New York; and Robert Wilson,
8
9 Ph.D., Chicago.
10

11
12 *Cerebrospinal Fluid Assay Center* (Wayne State University, Detroit, MI 48207) — Matthew P. Galloway, Ph.D.
13
14 (director); Mike Kaplan (deceased) and Rashid Lodhi.
15

16
17 *Deprenyl Metabolites Assay Center* (Institute for Medical Research, San Jose, CA 95128) — Ian Irwin, Ph.D. (director).
18

19 *Blood Tocopherol Assay Center* (Our Lady of Mercy Medical Center, Bronx, NY 10466) — Edward Norkus, Ph.D.
20
21 (director).
22

23 *Specimen Repository* (Department of Neurology, University of Rochester, Rochester, NY 14642) — Dorothy Flood,
24
25 Ph.D. (director), Thomas McNeill, Ph.D., Norma Harary, Ph.D., and Laurie Koek, B.S.
26

27 *Laboratory Surveillance Testing* (SciCor Laboratories, Indianapolis, IN 46241) — Robert L. Creveling, M.D. (director).
28
29
30

31
32 Parkinson Study Group PRECEPT investigators: Steering committee: *University of Rochester, Rochester, NY*: Ira
33
34 Shoulson, MD (principal investigator), Steven Schwid, MD (medical monitor), Christopher Hyson, MD (medical
35
36 monitor), David Oakes, PhD (chief biostatistician), Emily Gorbold, BA, (project coordinator), Alice Rudolph, PhD
37
38 (project coordinator), Aileen Shinaman, JD (Parkinson Study Group executive director), Cornelia Kamp, MBA (director,
39
40 Clinical Research Operations), Karl Kieburtz, MD, MPH (director, Clinical Trials Coordination Center); *Toronto Western*
41
42 *Hospital, University Health Network, Toronto, Ontario, Canada*: Anthony Lang, MD (coprincipal investigator); *Columbia*
43
44 *University Medical Center, New York, NY*: Stanley Fahn, MD; *Duke University Medical Center, Durham, NC*: Lisa
45
46 Gauger, BA; *Rush University Medical Center, Chicago, IL*: Christopher Goetz, MD; *Institute for Neurodegenerative*
47
48 *Disorders, New Haven, CT*: Kenneth Marek, MD, John Seibyl, MD.
49

50
51 Participating investigators and coordinators: *Colorado Neurological Institute, Englewood, CO*: Lauren Seeberger, MD,
52

53 Rajeev Kumar, MD; *London Health Sciences Center, London, Canada*: Mandar Jog, MD, Cheryl Horn, RN; *Rush–*
54

55 *Presbyterian–St. Luke’s Medical Center, Chicago, IL*: Kathleen Shannon, MD, Lucia Blasucci, RN, CCRC; *University of*
56
57

1 *Colorado Health Sciences Center, Denver, CO: Maureen Leehey, MD, Teresa Derian, RN; Ottawa Hospital Civic Site,*
2 *Ottawa, Ontario, Canada: David Grimes, MD, Melodie Mortensen, BSCN, Keely Haas, RN; University of Minnesota/*
3 *Minnesota VA Medical Center, Minneapolis, MN: Paul Tuite, MD, Susan Rolandelli, RN; University of California Irvine,*
4 *Irvine, CA: Neal Hermanowicz, MD, Shari Niswonger, RN; University of Rochester, Rochester, NY: Roger Kurlan, MD,*
5 *Irenita Gardiner, RN, CCRC; Toronto Western Hospital, University Health Network, Toronto, Ontario, Canada: Janis*
6 *Miyasaki, MD, FRCPC, Lisa Johnston, RN, BSCN, CNN; The Parkinson's Institute, Sunnyvale, CA: James Tetrud, MD,*
7 *Tracy Stewart, RN; NeuroHealth PD Movement Disorders Center, Warwick, RI: Joseph Friedman, MD, Hubert*
8 *Fernandez, MD, Margaret Lannon, RN, MS; University of Iowa, Iowa City, IA: Robert Rodnitzky, MD, Judith Dobson,*
9 *RN, CCRC; Mayo Clinic Arizona, Scottsdale, AZ: Virgilio Evidente, MD, Marlene Lind, RN; Oregon Health & Science*
10 *University, Portland, OR: Julie Carter, RN, MN, ANP, Pamela Andrews, BS; Chum-Hotel Dieu/McGill Center for Studies*
11 *in Aging, Montreal, Quebec, Canada: Michel Panisset, MD; Washington University School of Medicine, St. Louis, MO:*
12 *Brad Racette, MD, Patricia Deppen, RN; Baylor College of Medicine, Houston, TX: Joseph Jankovic, MD, Christine*
13 *Hunter, RN, CCRC; Institute for Neurodegenerative Disorders, New Haven, CT: Danna Jennings, MD, Barbara Fussell,*
14 *RN; Albany Medical College, Albany, NY: Eric Molho, MD, Stewart Factor, MD; Indiana University School of Medicine,*
15 *Indianapolis, IN: Joanne Wojcieszek, MD; University of California–Davis, Sacramento, CA: Lin Zhang, MD, PhD, Lisa*
16 *Wilson, MS, CCRP, Teresa Tempkin, RNC, MSN; Duke University Medical Center, Durham, NC: Burton Scott, MD,*
17 *Joanne Field, BSN, RN; Cleveland Clinic, Cleveland, OH: Thyagarajan Subramanian, MD, Ruth Kolb, CCRP; University*
18 *of Pennsylvania, Philadelphia, PA: Andrew Siderowf, MD, Amy Colcher, MD, Heather Maccarone, RN, BSN; University*
19 *of South Florida, Tampa, FL: Robert Hauser, MD, Joanne Nemeth, RN; Johns Hopkins University, Baltimore, MD:*
20 *Joseph Savitt, MD, PhD, Kevin Biglan, MD, MPH, Melissa Gerstenhaber, RNC, MSN; University of*
21 *Cincinnati/Cincinnati Children's Hospital, Cincinnati, OH: Alok Sahay, MD, Arif Dalvi, MD, Maureen Gartner, RN,*
22 *Donna Schwieterman, MA, CCRC; Mayo Clinic Jacksonville, Jacksonville, FL: Ryan Uitti, MD, Margaret Turk, RN;*
23 *University of Sherbrooke, Sherbrooke, Quebec, Canada: Jean Rivest, MD, Daniel Soucy, RN; University of Virginia,*
24 *Charlottesville, VA: Frederick Wooten, MD, Elke Rost-Ruffner, RN, BSN; Massachusetts General Hospital, Boston, MA:*
25 *Michael Schwarzschild, MD, PhD, Marsha Tennis, RN; Medical College of Georgia, Augusta, GA: Kapil Sethi, MD, Lisa*
26 *Hatch, RN, BSN; University of Tennessee–Memphis, Memphis, TN: Ronald Pfeiffer, MD, Brenda Pfeiffer, RN, BSN;*
27 *North Shore–LIJ Health System, Manhasset, NY: Andrew Feigin, MD, Jean Ayan, RN, Barbara Shannon, RN;*

1 *Northwestern University, Chicago, IL: Tanya Simuni, MD, Karen Williams, BA, Michele Wolff, BA; Medical University*
2 *of Ohio, Toledo, OH: Lawrence Elmer, MD, PhD, Kathy Davis, RN; University of Connecticut, Glastonbury, CT:*
3
4 *Antonelle de Marcaida, MD, Sheila Thurlow, RN; Hotel-Dieu Hospital-CHUM, Montreal, Quebec, Canada: Sylvain*
5
6 *Chouinard, MD, Hubert Poiffaut, RN; Barrow Neurological Institute, Phoenix, AZ: Holly Shill, MD, Mark Stacy, MD,*
7
8 *Lynn Marlor, BSN, MSHS, Jill Danielson, RN; The Parkinson's & Movement Disorder Institute, Fountain Valley, CA:*
9
10 *Daniel Truong, MD, Jacky Vo, MS; LSU Health Science Center Shreveport, Shreveport, LA: Richard Zweig, MD,*
11
12 *Rhonda Feldt, RN; Columbia University Medical Center, NY, NY: Cheryl Waters, MD, Angel Figueroa, BBA, Anne Tam,*
13
14 *BS, CCRC; University of Kansas Medical Center, Kansas City, KS: Rajesh Pahwa, MD, Amy Parsons, RN, BSN;*
15
16 *University of Southern California, Los Angeles, CA: Jennifer Hui, MD, Allan Wu, MD, Connie Kawai, RN, BSN, CCRC;*
17
18 *University of Alberta, Edmonton, AB, Canada: Richard Camicioli, MD, Pamela King, BScN, RN; University of Chicago,*
19
20 *Chicago, IL: Arif Dalvi, MD, Un Jung Kang, MD, Elizabeth Shaviers, Barbara Harding-Clay, CMA, CCRC; University of*
21
22 *Maryland School of Medicine, Baltimore, MD: Stephen Reich, MD, Lisa Shulman, MD, Carol Dignon, RN, MSN, Kelly*
23
24 *Dustin, RN; UMDNJ Robert Wood Johnson Medical School, New Brunswick, NJ: Margery Mark, MD, Deborah Caputo,*
25
26 *RN, MSN; Saskatoon Dist Health Board Royal University Hospital, Saskatoon SK, Canada: Ali Rajput, MD; Boston*
27
28 *University, Boston, MA: Peter Novak, MD, Cathi-Ann Thomas, RN, MS; Pacific Neuroscience Medical Group, Oxnard,*
29
30 *CA: James Sutton, MD, Juanita Young, CCRC; University of California-San Diego, La Jolla, CA: David Song, MD,*
31
32 *Deborah Fontaine, RNCS, MS; Creighton University, Omaha, NE: John M. Bertoni, MD, PhD, Carolyn Peterson, RN;*
33
34 *Medical College of Wisconsin, Milwaukee, WI: Karen Blindauer, MD, Jeannine Petit, ANP; Scott & White Hospital/Texas*
35
36 *A&M University, Temple, TX: Bala Manyam, MD, Danielle McNeil-Keller, LMSW, Jacqueline Whetteckey, MD;*
37
38 *Clinical Neuroscience Center, Southfield, MI: Peter LeWitt, MD, Maryan DeAngelis, RN, CCRC; University of Calgary,*
39
40 *Calgary, AB, Canada: Ranjit Ranawaya, MD, Oksana Suchowersky, MD, Carol Pantella, RN; Brigham & Women's*
41
42 *Hospital, Boston, MA: Lewis Sudarsky, MD; Beth Israel Deaconess Medical Center, Boston, MA: Daniel Tarsy, MD,*
43
44 *Linda Paul, NP, Lisa Scollins, NP; Long Island Jewish Medical Center, New Hyde Park, NY: Mark Forrest Gordon, MD;*
45
46 *Beth Israel Medical Center, NY, NY: Susan Bressman, MD, Alessandro DiRocco, MD, Karyn Boyar, RN, CNS, FNP;*
47
48 *Stanford University Medical Center, Stanford, CA: Helen Bronte-Stewart, MD, Amy Andrzejewski, BS; UMDNJ School*
49
50 *of Osteopathic Medicine, Stratford, NJ: Gerald Podskalny, DO; Cleveland Clinic Florida-Weston, Weston, FL: Nestor*
51
52
53
54
55
56
57
58
59
60

Galvez- Jimenez, MD, Jose Alvarez, CCRC; *University of Arkansas for Medical Sciences, Little Rock, AR*: Sami Harik, MD, Samer Tabbal, MD, Jana Patterson, RN

Biostatistics and coordination center staff: *University of Rochester, Rochester, NY*: Arthur Watts, BS, Rory Doolan, BA, Michele Goldstein, BS, Connie Orme, BA, Larry Preston, BPS, Tina Winebrenner.

Data monitoring committee: *The Parkinson's Institute, Sunnyvale, CA*: Caroline Tanner, MD, PhD (chair); *Johns Hopkins University, Baltimore, MD*: Steven Piantadosi, MD, PhD; *University of British Columbia, Vancouver, BC, Canada*: Jon Stoessl, MD, Paul Keown, MD; *University of Pennsylvania, Philadelphia, PA*: Lynn Schuchter, MD.

The following employees of Cephalon, Inc. and H. Lundbeck A/S were substantively involved in the design, conduct, and analysis of PRECEPT: *Lundbeck A/S, Copenhagen, Denmark*: Misser Forrest, MD, Thomas Bisgaard, MPOlSc, Erik Bardrum Nielsen, PhD, Sissel Vorstrup, MD. *Cephalon, Inc., Fraser, PA*: Heather Snyder, PhD, John Ondrasik, PhD, Lilliam Kingsbury, PhD, Steve Mulcahy, MS, Coleen Myers, BSN, Lesley Russell, MD.

Parkinson Study Group PostCEPT investigators:

Steering Committee: *University of Rochester, Rochester, NY*: Ira Shoulson, MD (principal investigator), Karl Kieburtz, MD, MPH, Bernard Ravina, MD, MCSE, David Oakes, PhD (chief biostatistician), Emily Flagg, BA (project coordinator), Roger Kurlan, MD; *Toronto Western Hospital, University Health Network, Toronto, Ontario, Canada*: Anthony Lang, MD; *Rush University Medical Center, Chicago, IL*: Christopher Goetz, MD; *Institute for Neurodegenerative Disorders, New Haven, CT*: Kenneth Marek, MD; *The Parkinson's Institute, Sunnyvale, CA*: Caroline Tanner, PhD, Robin Elliot, MA; *Columbia University, New York, NY*: Stanley Fahn, MD.

Oversight Committee: *Cephalon, Inc., Fraser, PA*: Gilbert Block; *Lundbeck A/S, Copenhagen, Denmark*: Misser Forrest, MD; *Department of Defense, Washington D.C.*: Stephen Grate, DVM; *The Parkinson's Disease Foundation, New York, NY*: Robin Elliot, MA; *NIH/NINDS, Bethesda, MD*: Diane DiEuliis, PhD, Wendy R. Galpern, MD, PhD.

PostCEPT Participating Investigators and Coordinators: *Chum-Hotel Dieu/McGill Center for Studies in Aging, Montreal, Quebec, Canada*: Michel Panisset, MD, Sylvain Chouinard, MD, Johanne Blais; *London Health Sciences, London, Ontario, Canada*: Mandar Jog, MD; *Colorado Neurological Institute, Englewood CO*: Dawn Miracle, BS, MS; *Rush University Medical Center, Chicago, IL*: Kathleen M. Shannon, MD; *University of Colorado Health Sciences Center,*

1 Denver, CO: Maureen Leehey, MD, Teresa Derian, RN; University of California, Irvine, Irvine, CA and The Phillip &
2 Carol Traub Center for Parkinson's Disease, Eisenhower Medical Center, Rancho Mirage, CA: Neal Hermanowicz, MD,
3 Shari Niswonger, RN; Ottawa Hospital Civic Site, Ottawa, Ontario, Canada: David Grimes, MD, RN; University of
4 Rochester, Rochester, NY: Roger Kurlan, MD, Nancy Pearson, RN, MS; NeuroHealth Parkinson's Disease Movement
5 Disorders Center, Warwick, RI: Joseph Friedman, MD, Margaret Lannon, RN, MS; The Parkinson's Institute, Sunnyvale,
6 CA; Toronto Western Hospital, University Health Network, Toronto, Ontario, Canada; University of Minnesota/
7 Minnesota VA Medical Center, Minneapolis, MN: Paul Tuite, MD, Susan Rolandelli, RN; University of Iowa, Iowa City,
8 IA: Robert Rodnitzky, MD, Judith Dobson, RN; Duke University Medical Center, Durham, NC: Burton Scott, MD,
9 Joanne Field, BSN, RN; Oregon Health & Science University, Portland, OR: Julie Carter, RN, MN, ANP, Pamela
10 Andrews; University of Pennsylvania, Philadelphia, PA: Andrew Siderowf, MD, Lisa Altin, BS; University of
11 Sherbrooke, Sherbrooke, Quebec, Canada: Daniel Soucy, RN; Johns Hopkins University, Baltimore, MD: Joseph Savitt,
12 MD, PhD, Melissa Gerstenhaber, RNC, MSN; LSU Health Science Center Shreveport, Shreveport, LA: Richard Zweig,
13 MD, Collette Hilliard, MS; Baylor College of Medicine, Houston, TX: Joseph Jankovic, MD, Christine Hunter, RN,
14 CCRC; Columbia University Medical Center, New York, NY: Cheryl Waters, MD, Angel Figueroa, CCRC; Northwestern
15 University, Chicago, IL: Tanya Simuni, MD, Karen Williams; Saskatoon Dist Health Board Royal University Hospital,
16 Saskatoon SK, Canada: Ali Rajput, MD, Marilyn Martin, BSc, ADV; University of Cincinnati/Cincinnati Children's
17 Hospital, Cincinnati, OH: Alberto Espay, MD, MSC; Sun Health Research Institute, Sun City, AZ: Holly Shill, MD;
18 University of South Florida, Tampa, FL; University of Tennessee-Memphis, Memphis, TN: Ronald Pfeiffer, MD, Brenda
19 Pfeiffer, RN, BSN; University of Virginia, Charlottesville, VA: Frederick Wooten, MD, Margaret F. Keller, RN, MS,
20 CCRC; Albany Medical College, Albany, NY: Eric Molho, MD, Katy Regan; Boston University, Boston, MA: Marie H.
21 Saint-Hilaire, MD, Cathi-Ann Thomas, RN, MS; Mayo Clinic Jacksonville, Jacksonville, FL: Margaret Turk, RN;
22 Medical College of Georgia, Augusta, GA: Buff Dill, BS, ED; Milton S. Hershey Medical Center, Hershey, PA:
23 Thyagarajan Subramanian, MD, Donna Stuppy, LPN; Institute for Neurodegenerative Disorders, New Haven, CT;
24 UMDNJ Robert Wood Johnson Medical School, New Brunswick, NJ: Margery H. Mark, MD, Debbie Caputo, MSN,
25 FNP-BC; University of Maryland School of Medicine, Baltimore, MD; Massachusetts General Hospital, Charleston, MA:
26 Michael Schwarzschild, MD, PhD; University of Toledo Health Science Center, Toledo, OH: Lawrence Elmer, MD, PhD,
27 Kathy Davis, RN Stephanie Wilson, RN, MSN, CCRC; University of Alberta, Edmonton, AB, Canada: Richard

1 Camicioli, MD, Pamela King, BScN, RN; University of California Davis, Sacramento, CA: Lin Zhang, MD, PhD, John
2 Bautista; University of Southern California, Los Angeles, CA; Creighton University, Omaha, NE; Pacific Neuroscience
3 Medical Group, Oxnard, CA; University of Calgary, Calgary, AB, Canada: Ranjit Ranaway, MD; University of
4 California San Diego, La Jolla, CA: David Song, MD, Deborah Fontaine, RNCS, MS; University of Kansas Medical
5 Center, Kansas City, KS: Kelly Lyons, PhD, Carey Mack, RN; Beth Israel Deaconess Medical Center, Boston, MA:
6 Daniel Tarsy, MD, Peggy Rose, RN; Brigham & Women's Hospital, Boston, MA: Lewis Sudarsky, MD, Georgette Hage,
7 MD; Cleveland Clinic Florida-Weston, Weston, FL: Nestor Galvez- Jimenez, MD; The Parkinson's & Movement
8 Disorder Institute, Fountain Valley, CA: Daniel Truong, MD; University of Chicago, Chicago, IL: Un Jung Kang, MD,
9 Joan Young, CCRC; North Shore-LIJ Health System, Manhasset, NY: Andrew Feigin, MD, Jean Ayan, RN; Stanford
10 University Medical Center, Stanford, CA: Helen Bronte-Stewart, MD; Beth Israel Medical Center, New York, NY: Karyn
11 Boyar, RN, CNS, FNP; University of Arkansas for Medical Sciences, Little Rock, AR: Jana Patterson, RN.
12
13
14
15
16
17
18
19
20
21
22
23
24 Biostatistics and Coordination Center Staff: University of Rochester, Rochester, NY: Earl Westerlund, Lisa Lang, Tina
25 Winebrenner, Nicole McMullen, Sandra Plumb, Cindy Casaceli, MBA.
26
27
28
29
30

31 DIGPD Study group. Steering committee: Jean-Christophe Corvol, MD, PhD (Pitié-Salpêtrière Hospital, Paris, principal
32 investigator of DIGPD), Alexis Elbaz, MD, PhD (CESP, Villejuif, member of the steering committee), Marie Vidailhet,
33 MD (Pitié-Salpêtrière Hospital, Paris, member of the steering committee), Alexis Brice, MD (Pitié-Salpêtrière Hospital,
34 Paris, member of the steering committee and PI for genetic analysis) ; Statistical analyses: Alexis Elbaz, MD, PhD (CESP,
35 Villejuif, PI for statistical analyses), Fanny Artaud, PhD (CESP, Villejuif, statistician); Principal investigators for sites
36 (alphabetical order): Frédéric Bourdain, MD (CH Foch, Suresnes, PI for site), Jean-Philippe Brandel, MD (Fondation
37 Rothschild, Paris, PI for site), Jean-Christophe Corvol, MD, PhD (Pitié-Salpêtrière Hospital, Paris, PI for site), Pascal
38 Derkinderen, MD, PhD (CHU Nantes, PI for site), Franck Durif, MD (CHU Clermont-Ferrand, PI for site), Richard Levy,
39 MD, PhD (CHU Saint-Antoine, Paris, PI for site), Fernando Pico, MD (CH Versailles, PI for site), Olivier Rascol, MD
40 (CHU Toulouse, PI for site); Co-investigators (alphabetical order): Anne-Marie Bonnet, MD (Pitié-Salpêtrière Hospital,
41 Paris, site investigator), Cecilia Bonnet, MD, PhD (Pitié-Salpêtrière Hospital, Paris, site investigator), Christine Brefel-
42 Courbon, MD (CHU Toulouse, site investigator), Florence Cormier-Dequaire, MD (Pitié-Salpêtrière Hospital, Paris, site
43 investigator), Bertrand Degos, MD, PhD (Pitié-Salpêtrière Hospital, site investigator), Bérangère Debilly, MD (CHU
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Clermont-Ferrand, site investigator), Alexis Elbaz, MD, PhD (Pitié-Salpêtrière Hospital, Paris, site investigator), Monique Galitsky (CHU de Toulouse, site investigator), David Grabli, MD, PhD (Pitié-Salpêtrière Hospital, Paris, site investigator), Andreas Hartmann, MD, PhD (Pitié-Salpêtrière Hospital, Paris, site investigator), Stephan Klebe, MD (Pitié-Salpêtrière Hospital, Paris, site investigator), Julia Kraemmer, MD (Pitié-Salpêtrière Hospital, site investigator), Lucette Lacomblez, MD (Pitié-Salpêtrière Hospital, Paris, site investigator), Sara Leder, MD (Pitié-Salpêtrière Hospital, Paris, site investigator), Graziella Mangone, MD, PhD (Pitié-Salpêtrière Hospital, Paris, site investigator), Louise-Laure Mariani, MD (Pitié-Salpêtrière Hospital, Paris, site investigator), Ana-Raquel Marques, MD (CHU Clermont Ferrand, site investigator), Valérie Mesnage, MD (CHU Saint Antoine, Paris, site investigator), Julia Muellner, MD (Pitié-Salpêtrière Hospital, Paris, site investigator), Fabienne Ory-Magne, MD (CHU Toulouse, site investigator), Violaine Planté-Bordeneuve, MD (Henri Mondor Hospital, Créteil, site investigator), Emmanuel Roze, MD, PhD (Pitié-Salpêtrière Hospital, Paris, site investigator), Melissa Tir, MD (CH Versailles, site investigator), Marie Vidailhet, MD (Pitié-Salpêtrière Hospital, Paris, site investigator), Hana You, MD (Pitié-Salpêtrière Hospital, Paris, site investigator); Neuropsychologists: Eve Benchetrit, MS (Pitié-Salpêtrière Hospital, Paris, neuropsychologist), Julie Socha, MS (Pitié-Salpêtrière Hospital, Paris, neuropsychologist), Fanny Pineau, MS (Pitié-Salpêtrière Hospital, Paris, neuropsychologist), Tiphaine Vidal, MS (CHU Clermont-Ferrand, neuropsychologist), Elsa Pomies (CHU de Toulouse, neuropsychologist), Virginie Bayet (CHU de Toulouse, neuropsychologist); Genetic core: Alexis Brice (Pitié-Salpêtrière Hospital, Paris, PI for genetic studies), Suzanne Lesage, PhD (INSERM, ICM, Paris, genetic analyses), Khadija Tahiri, PhD (INSERM, ICM, Paris, lab technician) Hélène Bertrand, MS (INSERM, ICM, Paris, lab technician), Graziella Mangone, MD, PhD (Pitié-Salpêtrière Hospital, Paris, genetic analyses); Sponsor activities and clinical research assistants: Alain Mallet, PhD (Pitié-Salpêtrière Hospital, Paris, sponsor representative), Coralie Villeret (Hôpital Saint Louis, Paris, Project manager), Merry Mazmanian (Pitié-Salpêtrière Hospital, Paris, project manager), Hakima Manseur (Pitié-Salpêtrière Hospital, Paris, clinical research assistant), Mostafa Hajji (Pitié-Salpêtrière Hospital, Paris, data manager), Benjamin Le Toullec, MS (Pitié-Salpêtrière Hospital, Paris, clinical research assistant), Vanessa Brochard, PhD (Pitié-Salpêtrière Hospital, Paris, project manager), Monica Roy, MS (CHU de Nantes, clinical research assistant), Isabelle Rieu, PhD (CHU Clermont-Ferrand, clinical research assistant), Stéphane Bernard (CHU Clermont-Ferrand, clinical research assistant), Antoine Faurie-Grepon (CHU de Toulouse, clinical research assistant).

1
2 ParkWest: Principal investigators: Guido Alves (Norwegian Centre for Movement Disorders, Stavanger University
3 Hospital), Ole-Bjørn Tysnes (Haukeland University Hospital). Investigators and study coordinators: Karen Herlofson,
4 Solgunn Ongre, Siri Bruun (Sørlandet Hospital Arendal); Ineke HogenEsch, Marianne Kjerandsen, Liv Kari Håland
5 (Haugesund Hospital); Wenche Telstad, Aliaksei Labusau, Jane Kastet (Førde Hospital); Bernd Müller, Geir Olve Skeie,
6 Charalampos Tzoulis (Haukeland University Hospital); Kenn Freddy Pedersen, Michaela Dreetz Gjerstad, Elin Bjelland
7 Forsaa, Jodi Maple-Grødem, Johannes Lange, Veslemøy Hamre Frantzen, Anita Laugaland, Karen Simonsen, Ingvild
8 Dalen (Stavanger University Hospital).

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19 PICNICS: Principal investigator -Roger Barker; study team - Caroline H Williams-Gray, Jonathan R Evans, David P
20 Breen, Gemma Cummins, Marta Camacho, Ruwani Wijeyekoon, Kirsten M Scott, Thomas Stoker, Julia C Greenland.

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