

Using global team science to identify genetic Parkinson's disease worldwide

Eva-Juliane Vollstedt, MD¹, Meike Kasten, MD^{1,2}, Christine Klein, MD^{1*} on behalf of the MJFF Global Genetic Parkinson's Disease Study Group (Supplementary Table 1)

¹Institute of Neurogenetics, University of Luebeck, Luebeck, Germany ²Department of Psychiatry, University of Luebeck, Luebeck, Germany

*Corresponding author: Christine Klein, MD, Institute of Neurogenetics, University of Lübeck, Maria-Goeppert-Str. 1, 23562 Lübeck, Germany, phone +49 4513101 8200, e-mail: christine.klein@neuro.uni-luebeck.de

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Talks on rare diseases in the field of neurology often start with a statement like this: "About 80% of all rare diseases have a neurologic manifestation and about 80% of those are genetic in origin". Although these numbers probably represent more of an estimate than well-documented evidence, rapidly advancing and cost-effective sequencing technologies have led to the quickly growing identification of patients with hereditary neurological diseases. While the importance of genetics for diagnosis and genetic counseling is undisputed, the recent development of first gene-targeted therapies entering clinical trial^{1,2} is adding an important new layer to the (re-)consideration of genetic testing in neurology. However, establishing accurate genotype-phenotype and genotype-treatment relationships requires large sample sizes. Systematic reviews can serve as instruments to combine information from several small samples, but unfortunately, this is often complicated by inconsistent and incomplete reporting of clinical and genetic data across studies. Thus, large multi-center approaches are necessary to systematically and uniformly characterize patients with genetic neurologic conditions and to eventually establish sizable clinical trial-ready cohorts.

Using genetic Parkinson's disease (PD) as an example and Illustrating the magnitude of the issue, up to ~300,000 patients worldwide are estimated to have hereditary forms of PD, representing 5% of an estimated total of 6 million patients with PD in 2018³. Monogenic forms of Parkinson's disease (PD) can be caused by mutations in *SNCA, LRRK2, VPS35, Parkin, PINK1,* and *DJ1*⁴. These genes have been unequivocally linked to PD according to the criteria established by the International Parkinson and Movement Disorder Society Task Force on Genetic Nomenclature in Movement Disorders⁵. In addition, *GBA* variants represent the strongest known genetic risk factor for PD, with an age-dependent penetrance of ~30% at the age of 80 years⁶. However, *individual* clinical information for patients with genetic PD is only

reported for a fraction of cases (n=1,769; Movement Disorder Society Genetic mutation database; MDSGene, www.mdsgene.org) in the international medical literature and publications are often biased towards unusual presentations of gene mutations. Both clinical expression and penetrance of gene mutations may vary considerably across different populations and ethnicities^{7,8}, further challenging pooling of data and their interpretation. Finally, given the growing availability of diagnostic genetic testing and the increasing difficulty to publish case reports of mutation carriers in peer-reviewed journals, we expect the proportion of published vs. unpublished cases to rapidly shift towards the latter.

As a result, neurologists commonly lack reliable reference data to be able to offer tailored counseling and treatment to patients with genetic PD and other hereditary neurological diseases.

Since the 1990s, there has been a growing interest and investment in large-scale, teambased research initiatives to address complex and multifaceted problems that require collaboration across *different* disciplines⁹. Likewise, there is an increasing necessity for – ideally global-scale - team science approaches of clinicians and researchers with *similar* interests joining forces to promote advanced research and to improve patient care. Employing novel ways of team science, electronic databases, and global communication, we performed a worldwide survey of genetic PD with an emphasis on availability of demographic, clinical, omics and imaging data as well as of biomaterials, to both foster and exploit global collaboration. In order to identify possible participants for our survey, we compiled the names of corresponding authors from articles included in the Movement Disorder Society Genetic mutation database (MDSGene, www.mdsgene.com) that comprises the following PD genes: *SNCA, LRRK2, VPS35, Parkin, PINK1*, and *DJ1^{7,8}*. For *GBA*,

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we screened the literature according to MDSGene criteria to identify corresponding authors of eligible articles (articles published in English with clinical information available). In addition, the "Genetic Epidemiology of Parkinson's disease" consortium (https://geopd.lcsb.uni.lu) contributed names of members not already identified as a corresponding author of publications represented in MDSGene. Additional contacts were included upon recommendation of participants (Figure 2). We next developed an online survey and invited the previously identified researchers to report availability of information on their genetic PD patients. To avoid multiple reporting of the same cases, we asked participants to indicate sharing of samples and encouraged participants from the same center to nominate one person to report all cases or to divide up the cases between reporting researchers. The survey was open for contributions for 5 weeks from March until May 2018. Two rounds of email reminders were sent out after two weeks each to enhance the response rate. Participants received no financial compensation for their contribution. A total of 103 researchers from different centers in 43 countries reported n=8,453 PD patients with mutations in PD genes (Figure 1, for more details please see Supplementary Table 2). The overall response rate to our invitation was 45% and 79% of the respondents completed the survey. Of these, 98% indicated their interest in further collaboration, and 45% sent personal emails expressing their interest to contribute further to this project and/or suggesting additional collaborative projects (Figure 2). Reflecting the worldwide distribution of participating centers, the backgrounds of the reported patient samples were diverse and include Arab, Asian, Ashkenazi Jewish, Caucasian, Hispanic, Indian, Mennonites, and North

LRRK2 or a *GBA* risk allele, followed by pathogenic variants in *Parkin* (n= 1500), *SNCA* and *PINK1* (n= 250 each), *DJ1* and *VPS35* (n= 30 each). The total number of reported monogenic

African. Almost equal numbers of patients (n=~3000) were reported to carry mutations in

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PD cases in the present study exceeds published cases by a factor of 2.6 (n=5,299 vs. n=2,064 cases (www.mdsgene.org), both excluding *GBA*) indicating that the majority of patients with genetic PD are not reflected in the published literature. Regarding data quality and completeness, information on PD nonmotor signs is particularly scarce in the literature. Even for cognition, published information is available only for about a third of the cases with monogenic PD (www.mdsgene.org)^{7,8}. In contrast, our present approach indicates availability of nonmotor signs in general for two thirds of the reported cases (Table 1).

To fill these knowledge gaps, a new collaborative mindset is indispensable. Our worldwide collaborative effort revealed an unprecedented level of data completeness, a paramount prerequisite when aiming to perform meaningful genotype-phenotype correlations and when selecting patients for clinical studies and trials^{7,8}. Team science of clinicians and researchers in the field of rare neurological diseases will facilitate sharing of expertise, promote new research opportunities, offer a network to foster new training opportunities for young researchers, and, advance development of individualized therapies. As a future perspective, collecting very large cohorts of monogenic PD may result in powerful enough data sets to perform meaningful natural history studies of individual monogenic forms as well as modifier studies. These can be modeled after studies performed in well-characterized longitudinal cohorts unselected for genetic cause¹⁰. In a similar vein, our approach may be expanded to mutation-negative early-onset PD patients with a positive family history in order to uncover rare novel mutations.

Based on the enthusiastic responses to our survey and the eager willingness to collaborate, we are confident that we have successfully established a team science approach that will specifically enable i) successfully increasing sample sizes of patients with rare neurological

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diseases; ii) leveraging neurology expertise globally; and iii) fostering team science among neurologists worldwide.

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Author contributions

A full list of all authors contributing to this manuscript can be viewed in Supplementary Table 1. EJV and CK were responsible for the conception and design of the study; EJV, CK and MK conducted the analysis and interpretation of the data; EJV and CK drafted the manuscript; all authors and the members of the MJFF Global Genetic Parkinson's Disease Study Group contributed to the acquisition of data and to the revision of the manuscript.

Conflicts of interest

The authors have no conflict of interest to declare.

Figure 1: Worldwide centers reporting patients with monogenic PD and PD patients with variants in GBA.

Legend: Please find a detailed list of the participating members of the **MJFF Global Genetic Parkinson's Disease Study Group** in Supplementary Online Table 1.



Figure 2: Response analysis.

Legend: *publications on GBA were screened according to the MDSGene protocols.



Table 1: Availability of information.

Legend: The participating centers were asked to report availability of an item if the relating data was available for at least a subset of the reported PD patients registered at their center.

	number of centers
	reporting available
	information
	(percentage of total
	participating centers
	(n=103))
Age	102 (99%)
Sex	102 (99%)
Ethnicity	101 (98%)
Pedigree	94 (91%)
Age at onset	97 (94%)
Unified Parkinson Disease Rating Scale	
(UPDRS)	68 (66%)
Hoehn & Yahr Scale	75 (73%)
Dopaminergic medications	79 (77%)
Nonmotor signs	68 (66%)
Environmental exposures	48 (47%)
Life style variables	36 (35%)
Treatment response	76 (74%)
Omics data	17 (17%)
-Genomics	14 (14%)
-Transcriptomics	2 (2%)
-Proteomics	1 (1%)
-Metabolomics	1 (1%)
Imaging	38 (37%)
-MRI	33 (32%)
-SPECT/PET	19 (18%)
-TCS	7 (7%)
DNA	88 (85%)
RNA	22 (21%)
Serum	28 (27%)
Plasma	26 (25%)
Whole blood	23 (22%)
Cerebrospinal fluid	8 (8%)
Fibroblasts	12 (12%)
iPSCs	11 (11%)
Brain tissue	11 (11%)

Supplement

Supplement 1: The MJFF Global Genetic Parkinson's Disease Study Group

Supplement 2: Number of centers and reported genetic PD patients

Name	Title	Affiliations
Jan Aasly	MD	Department of Neurology, St. Olavs Hospital, Trondheim,
		Department of Neuroscience, Norwegian University of
		Science and Technology, Trondheim, Norway
Charlos Adlor	MD	Department of Neurology, Mayo Clinic College of Medicine
Charles Auler	PhD	Scottdale Arizona USA
Azlina Abmad	PhD	Biomodical Science, University of Malaya, Kuala Lumpur
Annuar		Malaysia
Alberto Albanese	MD	Department of Neurology, IRCCS Istituto Clinico Humanitas.
		Rozzano, Milano, Italy;
		Department of Neurology, Catholic University, Milano, Italy
Roy Alcalay	MD,	Department of Neurology, Columbia University, New York,
	MS	New York, USA
Bashayer Al-	PhD	Behavioural genetics Unit, Department of Genetics,
Mubarak		Research Centre, King Faisal Specialist Hospital and
		Research Center, Riyadh, Saudi Arabia
Victoria Alvarez	PhD	Laboratório de Genética, Hospital Universitario Central de
		Asturias, Oviedo, Asturias, Spain
Brennie Andree-	MD	Department of Neurology, Hospital Luis Vernaza, Guayaquil,
Muñoz		Equador
Grazia Annesi	PhD	Institute of Molecular Bioimaging and Physiology, National
		Research Council, Germaneto, Cantazaro, Italy
Silke Appel-	MD	Pacific Parkinson's Research Centre, Division of Neurology,
Cresswell		Department of Medicine, Vancouver, British Columbia,
		Canada
David Arkadir	MD,	Department of Neurology, Hadassah Medical Center and
	PhD	the Hebrew University, Jerusalem, Israel
Sebastian Armasu	MSc	Division of Biomedical Statistics and Informatics,
		Department of Health Sciences Research, Mayo Clinic,
		Rochester, Minnesota, USA
Thomas R. Barber	MD	Oxford Parkinson's Disease Centre, Nuffield Department of
		Clinical Neuroscience, University of Oxford, Oxford,
		Oxfordshire, United Kingdom
Soraya Bardien	MD	Division of Molecular Biology and Human Genetics, Faculty

Cape Town, South Africa

of Medicine and Health Sciences, Stellenbosch University,

Supplement 1: The MJFF Global Genetic Parkinson's Disease Study Group

Melinda	PhD	DST/NWU Preclinical Drug Development Platform, North-
Barkhuizen		West University, Potchefstroom, North-West, South Africa
Matthew J. Barrett	MD	Department of Neurology, University of Virginia,
		Charlottesville, Virginia, USA
A. Nazlı Başak	PhD	Department of Molecular Biology and
		Genetics, Koç University School of Medicine, KUTTAM, Suna
		and Inan Kıraç Foundation, NDAL, Istanbul, Turkey
Thomas Beach	MD,	Civin Laboratory of Neuropathology, Banner Sun Health
	PhD	Research Institute, Sun City, Arizona, USA
Bruno A. Benitez	MD	Department of Psychiatry, Washington University in Saint
		Louis, Saint Louis, Missouri, USA
Daniela Berg	MD	Department of Neurology, Christian-Albrechts-Universität,
		Kiel, Germany
Kailash Bhatia	MD	Clinical Movement Disorders, UCL Institute of Neurology,
		University College London, London, United Kingdom;
		Clinical Neurology, National Hospital for Neurology,
		University College London, London, United Kingdom
Ferdinand Binkofski	MD	Clinical Cognitive Research, Department of Neurology,
		University Hospital RWTH Aachen, Aachen, Germany;
		Institute for Neuroscience and Medicine (INM-4), Research
		Center Juelich, Juelich, Germany
Cornelis	PhD	Laboratory of Neurogenetics, National Institute on Ageing,
Blauwendraat		National Institutes of Health, Bethesda, Maryland, USA
Vincenzo Bonifati	MD,	Department of Clinical Genetics, Erasmus MC, University
	PhD	Medical Center Rotterdam, Rotterdam, Netherlands
Vanderci Borges	MD	Department of Neurology, Universidade Federal de São
		Paulo, São Paulo, Brazil
Maria Bozi	MD,	Parkinson's and Movement Disorders Unit, Attiko Hospital,
	PhD	University of Athens, Haidari, Athens, Greece;
		Department of Neurology, Athens Psychiatry Hospital, NHS,
		Haidari, Athens, Greece
Alexis Brice	MD	ICM, Institut du Cerveau et de la Moelle épinière, Paris,
		France;
		UMR 1127, NS-PARK/FCRIN, Inserm, Paris, France;
		Sorbonne Université, Paris, France;
		UMR 7225, CNRS, Paris, France;
		Department of Genetics, Hôpital Pitié-Salpêtrière,
		Assistance Publique Hôpitaux de Paris, Paris, France
Laura Brighina	MD,	Department of Neurology, Milan Center for Neuroscience,
	PhD	University of Milano-Bicocca/San Gerardo Hospital, Monza,
		Monza Brianza, Italy
Kathrin Brockmann	MD	Department of Neurodegenerative Diseases, University of
		Tuebingen, Tuebingen, Baden Wuerttemberg, Germany;
		Hertie Institute for Clinical Brain Research and German
		Centre for Neurodegenerative Diseases, Tuebingen, Baden
		Wuerttemberg, Germany

Norbert	MD	Institute of Neurogenetics, University of Luebeck, Luebeck,
Brüggemann		Germany;
		Department of Neurology, University of Luebeck, Luebeck,
		Germany
Marta Camacho	MSc	Department of Clinical Neurosciences, University of
		Cambridge, Cambridge, United Kingdom
Francisco Cardoso	MD,	Department of Internal Medicine, UFMG, Belo Horizonte,
	PhD	Minas Gerais, Brazil
Andrea Carmine	PhD	Department of Neuroscience, Karolinska Institutet,
Belin		Stockholm, Sweden
Jonathan Carr	MD	Division of Neurology, Department of Medicine, Faculty of
		Medicine and Health Sciences, Stellenbosch University,
		Cape Town, South Africa
Piu Chan	MD,	Department of Neurobiology and Neurology, Capital
	PhD	Medical University, Xuanwu Hospital, Beijing, Beijing, China
Jorge Chang-	MD	Department of Molecular Biology, Hospital Luis Vernaza,
Castello		Guayaquil, Ecuador
Bruce Chase	PhD	Department of Biology, University of Nebraska at Omaha,
		Omaha, Nebraska, USA
Alice Chen-Plotkin	MD	Department of Neurology, University of Pennsylvania,
		Philadelphia, Pennsylvania, USA
Sun Ju Chung	MD,	Medical Genetic Center, Department of Neurology, Asan
	PhD	Medical Center, University of Ulsan College of Medicine,
		Seoul, South Korea
Roberto Cilia	MD	ASST Pini-CTO, Parkinson Institute, Milan, Milan, Italy
Jordi Clarimon	PhD	Department of Neurology, Biomedical Research Institute
		IIB-Sant Pau, Hospital de la Santa Creu i Sant Pau, Barcelona,
		Spain;
		Centro de Investigación Biomédica en Red sobre
		Enfermedades Neurodegenerativas (CIBERNED), Madrid,
		Spain
Lorraine Clark	PhD	Department of Pathology and Cell Biology, Vagelos College
		of Physicians & Surgeons, Columbia University Irving
		Medical Center, New York, New York, USA;
		Taub Institute for Research on Alzheimer's Disease and the
		Aging Brain, Columbia University Irving Medical Center, New
		York, New York, USA;
		Laboratory of Personalized Genomic Medicine, Vagelos
		College of Physicians & Surgeons, Columbia University Irving
		Medical Center, New York, New York, USA
Mario Cornejo-	MD	Neurogenetics Research Center, Instituto Nacional de
Olivas		Ciencias Neurologicas, Lima, Peru;
		Center for Global Health, Universidad Peruana Cayetano
		Hereula, Lima, Peru
Jean-Christophe	IVID	ILIVI, INSTITUT du Cerveau et de la Moelle epiniere, Paris,
Corvoi		
1	1	UNIK 1127, NS-PARK/FCRIN, Inserm, Paris, France;

		Sorbonne Université, Paris, France;
		UMR 7225, CNRS, Paris, France;
		Department of Genetics, Hôpital Pitié-Salpêtrière,
		Assistance Publique Hôpitaux de Paris, Paris, France
Carlos Cosentino	MD	Neurogenetics Research Center, Instituto Nacional de
		Ciencias Neurologicas, Lima, Peru;
		School of Medicine, Universidad Nacional Mayor de San
		Marcos, Lima, Peru
Patrick Cras	MD,	Department of Neurology, Antwerp University Hospital,
	PhD	Edegem, Belgium;
		Born Bunge Institute, Department of Neurology, University
		of Antwerp, Wilrijk, Belgium
David Crosiers	MD,	Department of Neurology, Antwerp University Hospital,
	PhD	Edegem, Belgium;
		Born Bunge Institute, Department of Neurology, University
		of Antwerp, Wilrijk, Belgium;
		Center for Molecular Neurology, VIB, Wilrijk, Belgium
Joana Damásio	MD	Department of Neurology, Hospital de Santo António -
		Centro Hospitalar do Porto, Porto, Portugal
Parimal Das	PhD	Centre for Genetic Disorders, Institute of Science, Banaras
		Hindu University, Varanasi, Uttar Pradesh, India
Patricia de	MD,	Department of Neurology, Universidade Federal de São
Carvalho Aguiar	PhD	Paulo, São Paulo, Brazil;
		Instituto Israelita de Ensino e Pesquisa, Hospital Israelita
		Albert Einstein, São Paulo, Brazil
Giuseppe De	MD	Department of Neurosciences and Reproductive and
Michele		Odontostomatological Sciences, Federico II University,
		Naples, Italy
Anna De Rosa	MD,	Department of Neurosciences and Reproductive and
	PhD	Odontostomatological Sciences, Federico II University,
		Naples, Italy
Elena Dieguez	MD	Neurology Institute, Universidad de la Republica,
		Montevideo, Uruguay
Jolanta Dorszewska	MD,	Laboratory of Neurobiology, Department of Neurology,
	PhD	Poznan University of Medical Sciences, Poznan, Poland
Sevda Erer	MD	Department of Neurology, Uludag University, Neurology,
		Bursa, Turkey
Sibel Ertan	MD	Department of Neurology, Istanbul University, Cerrahpaşa
		School of Medicine, Istanbul, Turkey
Matthew Farrer	PhD	Centre for Applied Neurosciences, University of British
		Columbia, Vancouver, British Columbia, Canada
Ekaterina Fedotova	MD,	Department of Neurogenetics, Research Center of
	PhD	Neurology, Moscow, Russia
Rosangela Ferese	PhD	IRCCS Neuromed, Localita' Camerelle, Pozzilli, Isernia, Italy
Carlo Ferrarese	MD	Department of Neurology, Milan Center for Neuroscience,
		University of Milano-Bicocca/San Gerardo Hospital, Monza,
		Monza Brianza, Italy

Henrique Ferraz	MD	Department of Neurology, Universidade Federal de São Paulo, São Paulo, Brazil
Ondrej Fiala	MD,	Department of Neurology, Institute of Neuropsychiatric
	PND	Care (INEP), Prague, Czech Republic
Tatiana Foroud	PhD	Department of Medical and Molecular Genetics, Indiana University School of Medicine, Indianapolis, Indiana, USA
Andrzei Friedman	MD.	Department of Neurology, Medical University in Warsaw,
	PhD	Warsaw, Poland
Roberta Frigerio	MD	Department of Neurology, NorthShore University
		HealthSystem, Evanston, Illinois, USA
Manabu Funayama	PhD	Research Institute for Diseases of Old Age, Graduate School
		of Medicine, Juntendo University, Bunkyo, Tokyo, Japan
Stefano Gambardella	PhD	IRCCS Neuromed, Localita' Camerelle, Pozzilli, Isernia, Italy
Gaetan Garraux	MD,	Department of Neurology, Centre Hospitalier Universitaire
	PhD	(CHU) de Liège, Liège, Belgium;
		MoVeRe group, GIGA-CRC in vivo imaging, University of
		Liege, Liège, Belgium
Emilia M. Gatto	MD	Movement Disorders, Department of Neurology, Instituto
		de Neurosciencias Buenos Aires, Buenos Aires, Argentina
Gençer Genç	MD	Department of Neurology, Şişli Etfal Training and Research
		Hospital, Istanbul, Turkey
Stefano Goldwurm	MD,	ASST Pini-CTO, Parkinson Institute, Milan, Milan, Italy;
	PhD	Department of Neuroscience "Rita Levi Montalcini",
		University of Turin, Turin, Italy
Juan Carlos Gomez-	PhD,	Neurodegenerative diseases, Biocruces Research Institute,
Esteban	MD	Biocruces, Barakaldo, Bizkaia, Spain
Pilar Gómez-Garre	PhD	Centro de Investigación Biomédica en Red sobre
		Enfermedades Neurodegenerativas (CIBERNED), Madrid,
		Spain;
		Unidad de Trastornos del Movimiento, Servicio de
		Neurología y Neurofisiología Clínica, Hospital Universitario
		Virgen del Rocío/CSIC/Universidad de Sevilla, Instituto de
		Biomedicina de Sevilla, Seville, Seville, Spain
Ana Gorostidi	PhD	Genomics Platform, Biodonostia Research Institute, San
		Sebastian, Gipuzkoa, Spain;
		Neuroscience Area, Biodonostia Research Institute, San
		Sebastian, Gipuzkoa, Spain
Donald Grosset	MD	Institute of Neurological Sciences, NHS Greater Glasgow &
		Clyde, Queen Elizabeth University Hospital, Glasgow,
		Scotland, United Kingdom;
		Institute of Neuroscience & Psychology, University of
		Glasgow, Glasgow, Scotland, United Kingdom
Hasmet Hanagasi	MD	Department of Neurology, Istanbul Faculty of Medicine,
_		Istanbul University, Istanbul, Turkey
John Hardy	MD	Neurodegenerative Disease, University College London, UCL
		Institute of Neurology, London, United Kingdom

Anhar Hassan	MD	Department of Neurology, Mayo Clinic, Rochester, Minnesota, USA
Nobutaka Hattori	MD, PhD	Department of Neurology, Juntendo University School of Medicine, Bunkyo, Tokyo, Japan
Robert A. Hauser	MD	Parkinson's Disease and Movement Disorders Center, Department of Neurology, University of South Florida, Tampa, Florida, USA
Peter Hedera	MD, PhD	Division of Movement Disorders, Department of Neurology, Vanderbilt University, Nashville, Tennessee, USA
Faycal Hentati	MD	Mongi Ben Hamida National Institute of Neurology, Tunis, Tunisia
Jens Michael Hertz	MD	Department of Clinical Genetics, Odense University Hospital, Odense C, Denmark
Janice L. Holton	MD	Queen Square Brain Bank for Neurological Disorders, Department of Clinical and Movement Neurosciences, UCL Queen Square Institute of Neurology, University College London, London, United Kingdom
Henry Houlden	MD	Neuromuscular Diseases, UCL Institute of Neurology, University College London, London, United Kingdom
Mara H. Hutz	PhD	Departamento de Genética, Universidade Federal do Rio Grande do Sul, Porto Alegre, Brazil
Takeshi Ikeuchi	MD, PhD	Molecular Genetics, Brain Research Institute, Niigata University, Niigata, Niigata, Japan
Sergey Illarioshkin	MD, PhD	Department of Neurogenetics, Research Center of Neurology, Moscow, Russia
Miguel Inca- Martinez	BS	Neurogenetics Research Center, Instituto Nacional de Ciencias Neurologicas, Lima, Peru
Jon Infante	MD, PhD	Department of Neurology, University Hospital Marques de Valdecilla, IDIVAL, University of Cantabria, Santander, Cantabria, Spain
Joseph Jankovic	MD	Parkinson's Disease Center and Movement Disorders Clinic, Department of Neurology, Baylor College of Medicine, Houston, Texas, USA
Beom Seok Jeon	MD, PhD	Department of Neurology, Seoul National University Hospital, Seoul, South Korea
Silvia Jesús	MD, PhD	Centro de Investigación Biomédica en Red sobre Enfermedades Neurodegenerativas (CIBERNED), Madrid, Spain; Unidad de Trastornos del Movimiento, Servicio de Neurología y Neurofisiología Clínica, Hospital Universitario Virgen del Rocío/CSIC/Universidad de Sevilla, Instituto de Biomedicina de Sevilla, Seville, Seville, Spain
Marlene Jimenez- Del-Rio	MSc, DSc	Department of Neurosciences, Universidad de Antioquia, Medellin, Colombia
Valtteri Kaasinen	MD, PhD	Dept. of Neurology and Turku PET Centre, University of Turku and Division of Clinical Neurosciences, Turku University Hospital, Turku, Finland

Meike Kasten	MD	Institute of Neurogenetics, University of Luebeck, Luebeck,
		Germany
		Department of Psychiatry and Psychotherapy, University of
		Luebeck, Luebeck, Germany
Hiroshi Kataoka	MD	Department of Neurology, Nara Medical University,
		Kashihara, Nara, Japan
Hideshi Kawakami	MD,	Epidemiology, Research Institute for Radiation Biology &
	PhD	Medicine, Hiroshima University, Hiroshima, Hiroshima,
		Japan
Yun Joong Kim	MD,	ILSONG Institute of Life Science, Hallym University, Anyang-
	PhD	si, Gyeonggi-do, South Korea;
		Department of Neurology, Hallym University, Anyang-si,
		Gyeonggi-do, South Korea
Christine Klein	MD	Institute of Neurogenetics, University of Luebeck, Luebeck,
		Germany
Péter Klivényi	MD	Department of Neurology, University of Szeged, Szeged,
		Hungary
Sulev Koks	MD,	The Perron Institute for Neurological and Translational
	PhD	Science, Nedlands, Western Australia, Australia
Inke R. König	PhD	Institute of Medical Biometry and Statistics, University of
		Luebeck, Luebeck, Germany
Vladimir Kostić	MD	Department for Neurodegeneration, Clinic for Neurology
		CCS, Belgrade, Serbia
Dariusz	MD,	Department of Neurology, Medical University in Warsaw,
Koziorowski	PhD	Warsaw, Poland
Rejko Krüger	MD	University of Luxembourg, Esch-sur-Alzette, Luxembourg
Anna Krygowska-	MD	Department of Neurology, Collegium Medicum, Jagiellonian
Wajs		University, Krakow, Poland
Jaime Kulisevsky	MD,	Department of Neurology, Biomedical Research Institute
	PhD	IIB-Sant Pau, Hospital de la Santa Creu i Sant Pau, Barcelona,
		Spain;
		Centro de Investigación Biomédica en Red sobre
		Enfermedades Neurodegenerativas (CIBERNED), Madrid,
		Spain
Anthony Lang	MD	Edmond J. Safra Program in Parkinson's Disease, Division of
		Neurology, Department of Medicine, University of Toronto,
		Toronto Western Hospital, Toronto, Ontario, Canada
Mark LeDoux	MD,	Department of Psychology and School of Health Studies,
	PhD	University of Memphis, Memphis,
Suzanne Lesage	MD	ICM, Institut du Cerveau et de la Moelle épinière, Paris,
		France;
		UMR 1127, NS-PARK/FCRIN, Inserm, Paris, France;
		Sorbonne Université, Paris, France;
		UMR 7225, CNRS, Paris, France;
		Department of Genetics, Hôpital Pitié-Salpêtrière,
		Assistance Publique Hôpitaux de Paris, Paris, France

	1	
Shen-Yang Lim	MD	Division of Neurology and the Mah Pooi Soo & Tan Chin
		Nam Centre for Parkinson's & Related Disorders, Faculty of
		Medicine, University of Malaya, Kuala Lumpur, Malaysia
Chin-Hsien Lin	MD,	Department of Neurology, National Taiwan University
	PhD	Hospital, Taipei, Taiwan
Katja Lohmann	PhD	Institute of Neurogenetics, University of Luebeck, Luebeck,
		Germany
Francisco Lopera	MD	Department of Neurosciences, Universidad de Antioquia,
		Medellin, Colombia
Grisel Lopez	MD	Medical Genetics Branch, National Human Genome
		Research Institute, NIH, Bethesda, Maryland, USA
Chin-Song Lu	MD	Professor Lu Neurological Clinic, Taoyuan, Taiwan
Tim Lynch	MD	Department of Neurology, The Dublin Neurological Institute at the Mater Misericordiae University Hospital, Dublin, Ireland; School of Medicine and Medical Sciences, University College
		Dublin Dublin Ireland
Maciei Machaczka	MD	Department of Clinical Science and Education Stockholm
		South Hospital and Karolinska Institutet, Stockholm, Sweden;
		Medical Faculty, University of Rzeszow, Rzeszow, Poland
Harutyun Madoev	MSc	Institute of Neurogenetics, University of Luebeck, Luebeck, Germany
Marina Magalhães	MD	Department of Neurology, Hospital de Santo António -
		Centro Hospitalar do Porto, Porto, Portugal
Kari Majamaa	MD	Research Unit of Clinical Neuroscience, University of Oulu,
		Oulu, Finland;
		Department of Neurology, Medical Research Center, Oulu
		University Hospital, Oulu, Finland
Demetrius	MD	Department of Neurology, Northshore University Health
Maraganore		System, Glenview, Illinois, USA
Karen Marder	MD	Department of Neurology, Columbia University, New York, New York, USA
Katerina	MD,	Department of Neurology, Northshore University Health
Markopoulou	PhD	System, Glenview, Illinois, USA;
		Department of Neurology, University of Chicago, Chicago,
		Illinois, USA
Mika Henrik	MD,	Department of Neurology, Faculty of Medicine, University of
Martikainen	PhD	Turku, Turku, Finland;
		Division of Clinical Neurosciences, Department of
		Neurology, Turku University Hospital, Turku, Finland
Ignacio Mata	PhD	Genomic Medicine, Lerner Research Institute, Cleveland
		Clinic, Cleveland, Ohio, USA;
		Department of Neurology, University of Washington,
		Seattle, Washington, USA
Pilar Mazzetti	MD	Neurogenetics Research Center, Instituto Nacional de
		Ciencias Neurologicas, Lima, Peru;

		School of Medicine, Universidad Nacional Mayor de San
		Marcos, Lima, Peru
George Mellick	MD	Griffith Institute for Drug Discovery (GRIDD), School of
		Environment and Science, Griffith University, Brisbane,
		Queensland, Australia
Manuel Menéndez-	MD	Servicio de Neurología, Hospital Universitario Central de
González		Asturias, Oviedo, Asturias, Spain
Federico Micheli	MD	Movement Disorders, Hospital de Clínicas José de San
		Martin, Buenos Aires, Argentina
Anat Mirelman	PhD	Movement Disorders, Neurological Institute, Tel-Aviv
		Medical Center, Tel-Aviv, Israel;
		Sackler School of Medicine, Tel-Aviv University, Tel-Aviv,
		Israel;
		Sagol School of Neuroscience, Tel-Aviv University, Tel-Aviv,
		Israel
Pablo Mir	MD,	Centro de Investigación Biomédica en Red sobre
	PhD	Enfermedades Neurodegenerativas (CIBERNED), Madrid,
		Spain;
		Unidad de Trastornos del Movimiento, Servicio de
		Neurología y Neurofisiología Clínica, Hospital Universitario
		Virgen del Rocío/CSIC/Universidad de Sevilla, Instituto de
		Biomedicina de Sevilla, Seville, Seville, Spain
Hiroyuki Morino	MD,	Epidemiology, Research Institute for Radiation Biology &
-	PhD	Medicine, Hiroshima University, Hiroshima, Hiroshima,
		Japan
Huw Morris	MD	Queen Square Brain Bank for Neurological Disorders,
		Department of Clinical and Movement Neurosciences, UCL
		Queen Square Institute of Neurology, University College
		London, London, United Kingdom
Renato P. Munhoz	MD,	Parana Parkinson's Association, Curitiba, Paraná, Brazil;
	PhD	Movement Disorders Centre, Division of Neurology,
		Department of Medicine, University of Toronto, Toronto
		Western Hospital, Toronto, Ontario, Canada
Anna Naito	PhD	Research Programs, The Michael J. Fox Foundation for
		Parkinson's Research, New York, New York, USA;
		Research Programs, Parkinson's Foundation, Miami, Florida,
		USA
Diana Angelika	MD,	Department of Neurology, The Dublin Neurological Institute
Olszewska	PhD	at the Mater Misericordiae University Hospital, Dublin,
		Ireland:
		School of Medicine and Medical Sciences, University College
		Dublin. Dublin. Ireland
Laurie J. Ozelius	PhD	Department of Neurology. Massachusetts General Hospital
		Charlestown, Massachusetts. USA
Shalini	PhD	Research Programs. The Michael J. Fox Foundation for
Padmanabhan		Parkinson's Research, New York, New York, USA
	1	

Coro Paisán-Ruiz	PhD	Department of Neurology, Icahn School of Medicine at
		Mount Sinai, New York, New York, USA;
		Genetics and Genomic Sciences, Icahn School of Medicine at
		Mount Sinai, New York, New York, USA;
		Psychiatry, Icahn School of Medicine at Mount Sinai, New
		York. New York. USA:
		Friedman Brain Institute. Icahn School of Medicine at
		Mount Sinai, New York, New York, USA:
		Mindich Child Health and Development, Icahn School of
		Medicine at Mount Sinai. New York. New York. USA
Havdeh Pavami	MD	Departments of Neurology and Genetics, University of
		Alabama at Birmingham, Birmingham, Alabama, USA
Silvio Peluso	MD,	Department of Neurosciences and Reproductive and
	PhD	Odontostomatological Sciences, Federico II University.
		Naples, Italy
Sonia Petkovic	PhD	Institute of Neurogenetics. University of Luebeck. Luebeck.
		Germany
Simona Petrucci	MD.	Department of Clinical and Molecular Medicine, Sapienza
	PhD	University of Rome, Rome, Italy;
		Division of Medical Genetics, IRCCS-Casa Sollievo della
		Sofferenza, San Giovanni Rotondo, Italy
Gianni Pezzoli	MD	ASST Pini-CTO Parkinson Institute Milan Milan Italy
Márcia Pimentel	PhD	Department of Genetics Institute of Biology Roberto
	FIID	Alcantara Gomes, State University of Rio de Janeiro, Rio de
		lanoiro. Pio do lanoiro. Brazil
Maltor Dirkor		Department of Neurology, Wilhelminenspital, Vienna
		Austria
Peter P	MD	Institute for Biomedicine, Affiliated Institute of the
Pramstaller	NIL O	University of Lübeck Eurac Research Bolzano South Tyrol
		Italy.
		Department of Neurology Central Hospital Bolzano, South
		Tyrol. Italy
Teeratorn Pulkes	MD	Division of Neurology, Department of Medicine.
		Ramathibodi Hospital, Mahidol University, Raithevi.
		Bangkok. Thailand
Andreas	MD.	Department of Neurology, Clinical Sciences, Lund University,
Puschmann	PhD	Lund, Sweden:
		Department of Neurology, Skåne University, Lund, Sweden
Aldo Quattrone	MD	Institute of Molecular Bioimaging and Physiology, National
	IVID	Research Council Germaneto Cantazaro Italy
Victor Raggio	MD	Department of Genetics, Universidad de la Republica
Victor Naggio	IVID	Montevideo Uruguay
Gerhard Bansmaur	МП	Department of Neurology 2, Kenler University Hospital Linz
		Austria
Carlos Pieder		Movement Disorders Unit Hespital de Clínicas de Porte
	טואן	Alegre Porte Alegre Prezil
		Alegie, Pullo Alegie, Bld211

Olaf Riess	MD	Institute of Medical Genetics and Applied Genomics, University of Tuebingen, Tuebingen, Germany
Federico	MD	Department of Neurology, Medical University of South
Rodriguez-Porcel		Carolina, Charleston, South Carolina, USA
Ekaterina Rogaeva	PhD	Tanz Centre for Research in Neurodegenerative Disease,
		University of Toronto, Toronto, Ontario, Canada
Owen A. Ross	MD	Department of Neuroscience, Mayo Clinic, Jacksonville, Florida, USA
Javier Ruiz-	MD,	Neuroscience Area, Biodonostia Research Institute, San
Martinez	PhD	Sebastian, Gipuzkoa, Spain;
		Department of Neurology, University Hospital Donostia, San
		Sebastian, Gipuzkoa, Spain
Esther Sammler	MD,	Neurology Department, Ninewells Hospital and Medical
	PhD	School, Dundee, United Kingdom;
		MRC Protein Phosphorylation and Ubiquitylation Unit,
		University of Dundee, Dundee, United Kingdom
Marta San Luciano	MD,	Department of Neurology, Movement Disorders and
	MS	Neuromodulation Center, University of California San
		Francisco, San Francisco, California, USA
Wataru Satake	MD,	Division of Neurology/Molecular Brain Science, Kobe
	PhD	University Graduate School of Medicine, Kobe, Hyogo,
		Japan
Rachel Saunders-	MD	Department of Neurology, Icahn School of Medicine at
Pullman		Mount Sinai, New York, New York, USA
Ali Sazci	PhD	Department of Medical Biology and Genetics, Faculty of
		Medicine, University of Kocaeli, Kocaeli, Turkey
Clemens Scherzer	MD	Center for Advanced Parkinson's Disease Research,
		Department of Neurology, Brigham & Women's Hospital,
		Harvard Medical School, Boston, Massachusetts, USA
Anette Schrag	MD	Queen Square Brain Bank for Neurological Disorders,
		Department of Clinical and Movement Neurosciences, UCL
		Queen Square Institute of Neurology, University College
		London, London, United Kingdom
Artur Schumacher-	MD,	Departamento de Farmacologia, Universidade Federal do
Schuh	PhD	Rio Grande do Sul, Porto Alegre, Brazil
Manu Sharma	PhD	Institute for Clinical Epidemiology and Applied Biometry,
		Centre for Genetic Epidemiology, University of Tuebingen,
		Tuebingen, Germany
Ellen Sidransky	MD	Medical Genetics Branch, National Human Genome
		Research Institute, NIH, Bethesda, Maryland, USA
Andrew B.	PhD	Laboratory of Neurogenetics, National Institute on Ageing,
Singleton	 	National Institutes of Health, Bethesda, Maryland, USA
Maria Skaalum	PhD	Department of Occupational Medicine and Public Health,
Petersen		Ine Faroese Hospital System, Torshavn, Faroe Islands;
		Faculty of Health Sciences, Centre of Health Science,
		University of the Faroe Islands, Tórshavn, Faroe Islands

Stefanie Smolders		Born Bunge Institute, Department of Neurology, University
		of Antwerp, Wilrijk, Belgium;
		Center for Molecular Neurology, VIB, Wilrijk, Belgium;
		Biomedical Sciences, University of Antwerp, Wilrijk, Belgium
Mariana Spitz	MD,	Department of Neurology, State University of Rio de
	PhD	Janeiro, Rio de Janeiro, Rio de Janeiro, Brazil
Leonidas Stefanis	MD	First Department of Neurology, Medical School of the
		National and Kapodistrian University of Athens, Eginition
		Hospital, Athens, Greece;
		Translational Medicine, Biomedical Research Foundation of
		the Academy of Athens, Athens, Greece
Walter Struhal	MD	Department for Neurology, Karl Landsteiner Private
		University, Tulln, Lower Austria, Austria
Carolyn Sue	MD	Department of Neurogenetics, Kolling Institute, University
		of Sydney, Sydney, New South Wales, Australia;
		Department of Neurology, Royal North Shore Hospital, St
		Leonards, New South Wales, Australia
Matthew Swan	MD	Department of Neurology, Icahn School of Medicine at
		Mount Sinai, New York, New York, USA
Maria Swanberg	PhD	Translational Neurogenetics Unit, Wallenberg Neuroscience
		Center, BMC A10, Experimental Medical Science, Lund
		University, Lund, Sweden
Pille Taba	MD,	Department of Neurology and Neurosurgery, University of
	PhD	Tartu, Tartu, Estonia
Ricardo Taipa	MD,	Portuguese Brain Bank, Neuropathology Unit, Department
	PhD	of Neurosciences, Centro Hospitalar Universitário do Porto,
		Porto, Portugal
Manuela Tan	MD	Queen Square Brain Bank for Neurological Disorders,
		Department of Clinical and Movement Neurosciences, UCL
		Queen Square Institute of Neurology, University College
		London, London, United Kingdom
Ai Huey Tan	MD	Division of Neurology and the Mah Pooi Soo & Tan Chin
		Nam Centre for Parkinson's & Related Disorders, Faculty of
		Medicine, University of Malaya, Kuala Lumpur, Malaysia
Eng-King Tan	MD	Department of Neurology, National Neuroscience Institute,
		Duke NUS Medical School, Singapore General Hospital,
		Singapore, Singapore
Beisha Tang	MD	Department of Neurology, Xiangya Hospital, Central South
		University, Changsha, Hunan Province, China;
		Department of Gerontology, Xiangya Hospital, Central South
		University, Changsha, Hunan Province, China;
		National Clinical Research Center for Geritatric Disorders
		(Xiangya), Changsha, Hunan Province, China;
		Center for Medical Genetics, School of Life Sciences, Central
		South University, Changsha, Hunan Province, China
Nahid Tayebi	PhD	Medical Genetics Branch, National Human Genome
		Research Institute, NIH, Bethesda, Maryland, USA

Avner Thaler	MD,	Movement Disorders, Neurological Institute, Tel-Aviv				
	PhD	Medical Center, Tel-Aviv, Israel;				
		Sackler School of Medicine, Tel-Aviv University, Tel-Aviv,				
		Israel;				
		Sagol School of Neuroscience, Tel-Aviv University, Tel-Aviv,				
		Israel				
Astrid Thomas	MD,	Department of Neuroscience Imaging and clinical sciences,				
	PhD	Institute of Neurology, University Chieti Pescara, Chieti, Italy				
Tatsushi Toda	MD,	Department of Neurology, Graduate School of				
	PhD	Medicine, University of Tokyo, Bunkyo-ku, Tokyo, Japan				
Mathias Toft	MD,	Department of Neurology, Oslo University Hospital, Oslo,				
	PhD	Norway;				
		Institute of Clinical Medicine, University of Oslo, Oslo,				
		Norway				
Luis Torres	MD	Movement Disorders Unit, Instituto Nacional de Ciencias				
		Neurologicas, Lima, Peru				
Vitor Tumas	MD	Movement Disorders Unit, Universidade de São Paulo,				
		Ribeirão Preto, Brazil				
Enza Maria Valente	MD,	Neurogenetics Unit, IRCCS Fondazione Santa Lucia, Rome,				
	PhD	Italy;				
		Division of Medical Genetics, IRCCS-Casa Sollievo della				
		Sofferenza, San Giovanni Rotondo, Italy				
Christine Van	MD	Born Bunge Institute, Department of Neurology, University				
Broeckhoven		of Antwerp, Wilrijk, Belgium;				
		Center for Molecular Neurology, VIB, Wilrijk, Belgium;				
		Biomedical Sciences, University of Antwerp, Wilrijk, Belgium				
Laszlo Vecsei	MD	Department of Neurology, University of Szeged, Szeged,				
		Hungary;				
		Neuroscience Research Group, MIA-SZIE, Szeged, Hungary				
Carlos Velez-Pardo	MSC,	Department of Neurosciences, Universidad de Antioquia,				
	DSc	Medellin, Colombia				
Marie Vidailhet	MD	ICM, Institut du Cerveau et de la Moelle épinière, Paris,				
		France;				
		Sorbonne Universite, Paris, France;				
		Department of Neurology, Hopital Pitle-Salpetriere, Paris,				
		France				
Eva-Juliane	MD	Institute of Neurogenetics, University of Luebeck, Luebeck,				
Vollstedt		Germany				
Thomas T. Warner	MD	Queen Square Brain Bank for Neurological Disorders,				
		Department of Clinical and Movement Neurosciences, UCL				
		Queen Square Institute of Neurology, University College				
		London, London, United Kingdom;				
		Reta Liid Weston Institute, UCL Institute of Neurology,				
Caralias II		Department of Clinical Neurosciences, United Kingdom				
Caroline H.	IVIKCP,	Department of Clinical Neurosciences, University of				
williams-Gray	טחאן	Campriage, Campriage, United Kingdom				

Juliane	MD	Institute of Neurogenomics, Helmholtz Zentrum Muenchen,
Winkelmann		Neuherberg, Germany;
		Neurogenetics, Technische Universitaet Muenchen, Munich,
		Germany;
		Institute of Human Genetics, Klinikum rechts der Isar der
		TUM, Munich, Germany;
		Munich Cluster for Systems Neurology (SyNergy), Munich,
		Germany
Dirk Woitalla	MD	Department of Neurology, St.Josef Krankenhaus, Essen,
		Germany;
		Department of Neurology, Ruhr-Universitaet Bochum,
		Bochum, Germany
Nicholas W. Wood	MD	Queen Square Brain Bank for Neurological Disorders,
		Department of Clinical and Movement Neurosciences, UCL
		Queen Square Institute of Neurology, University College
		London, London, United Kingdom
Zbigniew K.	MD	Department of Neurology, Mayo Clinic, Jacksonville, Florida,
Wszolek		USA
Ruey-Meei Wu	MD	Department of Neurology, National Taiwan University
		Hospital, Taipei, Taiwan;
		Department of Neurology, National Taiwan University
		College of Medicine, Taipei, Taiwan
Yih-Ru Wu	MD	Department of Neurology, Chang Gung University, Chang
		Gung Memorial Hospital, Taipei, Taiwan
Tao Xie	MD,	Department of Neurology, University of Chicago, Chicago,
	PhD	Illinois, USA
Hiroyo Yoshino	PhD	Research Institute for Diseases of Old Age, Graduate School
		of Medicine, Juntendo University, Bunkyo, Tokyo, Japan
Baorong Zhang	MD	Department of Neurology, Second Affiliated Hospital,
		College of Medicine, Zhejiang University, Hangzhou,
		Zhejiang, China
Alexander Zimprich	MD	Department of Neurology, Medical University, Vienna,
		Austria

Supplement 2: Number of centers and reported genetic PD patients

Legend: In total, 103 centers shared numbers of available genetic PD patients (listed in the table), and additional 5 centers completed the survey but did not indicate numbers. Researchers were asked to report all PD patients with mutations in one of the listed genes that were registered at their center. Whether these patients are still available for future studies was not assessed, but given the fact that the genes of interest were all described within the last two decades, the proportion of historic cases included in these numbers should be relatively low.

*Mutations in GBA are considered risk variants for PD. The same applies to certain mutations in LRRK2 in the Asian population (e.g. G2385R), which we did not specifically assess in our survey, but which centers may have included in the reported numbers.

	centers	DJ1	GBA*	LRRK2*	Parkin	PINK1	SNCA	VPS35
Argentina	2	0	0	10	1	0	0	0
Australia	2	0	0	19	11	3	2	4
Austria	1	0	30	1	6	1	0	3
Belgium	2	0	12	43	1	0	1	0
Brazil	5	0	26	39	34	2	1	0
Canada	2	2	34	328	65	62	65	3
China	3	5	90	330	264	10	0	5
Colombia	1	0	13	3	12	0	0	0
Denmark	1	0	0	0	2	0	0	0
Ecuador	1	0	1	0	0	0	0	0
Estonia	1	0	15	25	5	0	1	5
Faroe Islands	1	0	0	1	0	0	0	0
Finland	3	0	15	2	2	0	2	0
France	1	2	124	189	145	19	27	5
Germany	4	1	119	20	23	12	2	2
Greece	2	0	38	1	3	0	77	0
Hungary	1	0	3	1	0	0	0	0
India	1	0	6	2	5	3	0	0
Ireland	1	1	21	2	25	1	0	0
Israel	2	1	240	180	1	0	0	1
Italy	9	3	227	147	117	28	22	0
Japan	4	0	65	20	293	23	19	4
Malaysia	1	0	0	200	0	0	0	0
Netherlands	1	1	15	1	3	0	1	0
Norway	2	0	35	40	5	4	0	0
Peru	1	0	20	1	1	0	0	0
Poland	2	0	0	1	42	2	2	0
Portugal	2	1	0	0	4	1	0	0
Republic of Korea	3	0	0	0	27	6	0	0

Russia	1	1	32	15	40	2	2	1
Saudi Arabia	1	1	0	0	15	4	0	0
Serbia	1	1	78	6	17	1	0	0
Singapore	1	0	20	0	0	0	0	0
South Africa	2	0	13	15	19	4	2	0
Spain	7	2	77	386	72	4	8	0
Sweden	2	2	37	14	2	0	6	0
Taiwan	3	1	27	12	30	11	0	2
Thailand	1	0	24	0	14	0	0	0
Tunisia	1	0	0	559	10	59	0	0
Turkey	3	3	2	2	36	2	2	0
United Kingdom	4	0	115	31	13	5	7	0
Uruguay	1	0	0	12	1	0	0	0
U.S.A.	15	1	1614	561	192	8	9	0