

FIGURE 4. Absence of post-phosphoryl modification in wild-type lung and testis. *A*, IMAC bead-binding assays for α -DG from wild-type tissues. α -DG-enriched samples from skeletal muscle (*skm*), brain, liver, lung, and testis of C57BL/6 mice were tested for binding to IMAC beads. The void (*v*) and bound (*b*) fractions were collected. The *arrow* indicates the IIH6-positive population of lung α -DG. The *arrowhead* indicates the IIH6-negative fraction of lung α -DG bound to beads. An *asterisk* indicates a background signal that is not specific for IIH6 antibody. *B*, chemical dephosphorylation of α -DG from wild-type tissues. α -DG-enriched samples from skeletal muscle (*skm*), brain, liver, lung and testis of C57BL/6 mice were treated with HFAq and then analyzed by Western blot using anti-DG core antibody.

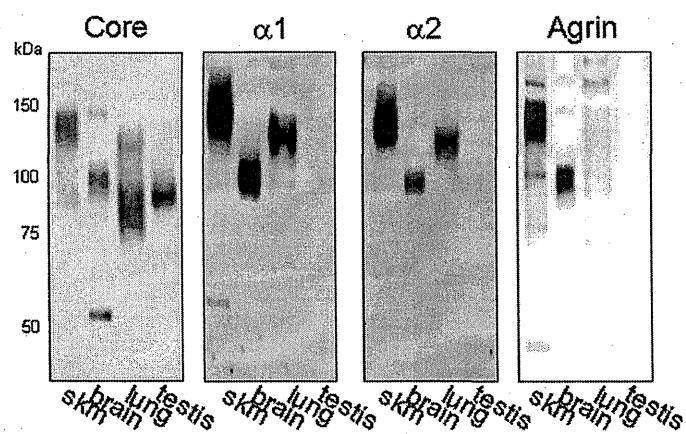


FIGURE 5. Ligand-binding assays for lung and testis α -DG. Ligand binding (laminin $\alpha 1$, $\alpha 2$, and agrin) was assessed in α -DG-enriched samples from skeletal muscle (*skm*), brain, lung, and testis using ligand overlay assays.

α -DG might have ligand proteins that do not require *O*-mannosyl modification; for example, a chondroitin sulfate proteoglycan biglycan has been shown to interact with protein core of the α -DG C-terminal domain in a glycosylation-independent

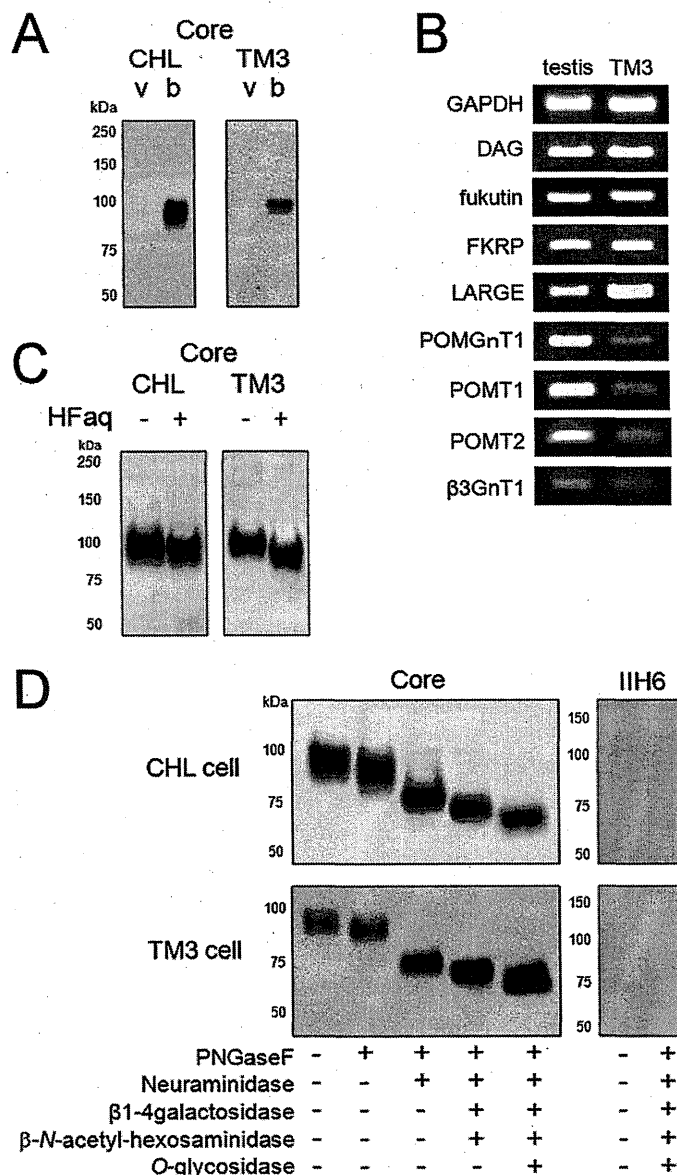


FIGURE 6. Absence of post-phosphoryl modification of α -DG in CHL and TM3 cells. *A*, IMAC bead-binding assays for α -DG from CHL cells and TM3 cells. α -DG-enriched samples from CHL and TM3 cell lysates were tested for binding to IMAC beads. The void (*v*) and bound (*b*) fractions were collected. *B*, RT-PCR analysis of TM3 cells. RT-PCR analysis was performed to detect transcripts encoding proteins implicated in α -DG glycosylation. *C*, chemical dephosphorylation of α -DG from wild-type tissues. α -DG-enriched samples from CHL and TM3 cell lysates were treated with HFAq. *D*, enzymatic deglycosylation of α -DG from CHL and TM3 cells. α -DG-enriched samples from CHL and TM3 cells were digested with glycosidase mixtures (peptide-*N*-glycosidase (PNGaseF), neuraminidase, $\beta 1-4$ galactosidase/ β -*N*-acetyl-hexosaminidase, and *O*-glycosidase). Following the IMAC bead-binding assay, HFAq treatment, and enzymatic deglycosylation, the samples were analyzed by Western blot, using antibodies against the α -DG core protein (Core) or the functionally glycosylated form (IIH6).

manner (38). DG is also thought to serve as a signaling molecule (39). For example, the cytoplasmic tail of β -DG interacts with several signaling molecules, including caveolin-3, Grb2, and mitogen-activated protein (MAP) kinase kinase 2 (40). Although the significance of these interactions is not well understood, it is possible that DG serves as a scaffold to position interacting proteins at their proper cellular location (9, 41). Taken together, these observations suggest that the presence of

α -Dystroglycan in FKRP Mutants

DG without post-phosphoryl modification could be functionally important in various tissue types.

Future work to determine the molecular structure of the post-phosphoryl moiety, and to identify genes involved in its biosynthesis, will contribute to understanding the biological basis of this unique post-translational modification and disease pathogenesis. Our present data contributes to the foundation for such research. Recently, it has been shown that LARGE can act as a bifunctional glycosyltransferase, with both xylosyltransferase and glucuronyltransferase activities (42). Involvement of these activities in the post-phosphoryl modification also should be clarified in the future.

Overall, our results indicate that phosphorylated O-mannose not only plays critical roles in the pathogenesis of dystroglycanopathy but also is a key determinant in the maturation of α -DG as a laminin receptor in normal tissues and cells.

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Comprehensive Research Synopsis and Systematic Meta-Analyses in Parkinson's Disease Genetics: The PDGene Database

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Abstract

More than 800 published genetic association studies have implicated dozens of potential risk loci in Parkinson's disease (PD). To facilitate the interpretation of these findings, we have created a dedicated online resource, PDGene, that comprehensively collects and meta-analyzes all published studies in the field. A systematic literature screen of ~27,000 articles yielded 828 eligible articles from which relevant data were extracted. In addition, individual-level data from three publicly available genome-wide association studies (GWAS) were obtained and subjected to genotype imputation and analysis. Overall, we performed meta-analyses on more than seven million polymorphisms originating either from GWAS datasets and/or from smaller scale PD association studies. Meta-analyses on 147 SNPs were supplemented by unpublished GWAS data from up to 16,452 PD cases and 48,810 controls. Eleven loci showed genome-wide significant ($P < 5 \times 10^{-8}$) association with disease risk: *BST1*, *CCDC62/HIP1R*, *DGKO/GAK*, *GBA*, *LRRK2*, *MAPT*, *MCCC1/LAMP3*, *PARK16*, *SNCA*, *STK39*, and *SYT11/RAB25*. In addition, we identified novel evidence for genome-wide significant association with a polymorphism in *JTG48* (rs7077361, OR 0.88, $P = 1.3 \times 10^{-8}$). All meta-analysis results are freely available on a dedicated online database (www.pdgene.org), which is cross-linked with a customized track on the UCSC Genome Browser. Our study provides an exhaustive and up-to-date summary of the status of PD genetics research that can be readily scaled to include the results of future large-scale genetics projects, including next-generation sequencing studies.

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¶ Memberships of the consortia are provided in Text S1.

Introduction

Parkinson's disease (PD) is the second most common neurodegenerative disease with a prevalence of ~1% over 60 years of age [1]. Approximately 5–10% of the patients show an autosomal dominant or recessive mode of inheritance, and several causative genes have been identified, e.g. *SNCA*, *LRRK2*, *PARK2*, and *PINK1* (for review see ref. [2]). Recently, two other novel autosomal dominant PD genes, *VPS35* and *EIF4G1* [3–5], have been identified, the former via application of next-generation sequencing techniques. It can be anticipated that causal mutations in additional genes will emerge within the next years. However, the vast majority of patients suffer from non-Mendelian forms of PD, which are likely caused by the combined effects of genetic and environmental factors. In order to decipher the genetic architecture underlying PD susceptibility, more than 800 genetic association studies have been performed over the past 20 years. While early candidate gene studies and subsequent meta-analyses provided conclusive evidence showing that polymorphisms in *SNCA* [6] (encoding alpha-synuclein), *LRRK2* [7] (leucine-rich repeat kinase 2), *MAPT* [8] (microtubule-associated protein tau), and *GBA* [9] (acid beta-glucosidase) significantly impact PD susceptibility, most association studies in the field provided inconclusive or even conflicting results.

During the last few years, genome-wide association studies (GWAS) [10–19] have postulated additional PD loci. While the early GWAS and a GWAS-meta-analysis [20] were of limited sample sizes and yielded mostly inconsistent results, more recent studies have identified a number of loci that were independently confirmed in follow-up studies (e.g. *GAK*, *BST1*, and *PARK16*, see Table 1 for all proposed GWAS findings across GWAS publications). Very recently, a GWAS meta-analysis [21] implicated several other new putative PD loci which currently await

further validation. Despite this progress, approximately 40% or more of the population-attributable risk probably remains unexplained by today's most promising PD loci [21]. To this end, genetic association studies remain one of the mainstays of PD genetics research. However, GWAS and other large-scale association studies typically only highlight the most promising results and often do not provide data on variants showing suggestive evidence for association, or previously implied variants that could not be confirmed in the GWAS setting. As a result, the cumulative genetic evidence in favor of or against association with certain variants in the PD field is becoming increasingly difficult to follow, evaluate and interpret. To address this problem, we have comprehensively collected, catalogued and systematically meta-analyzed the data from all genetic association studies published in the field of non-Mendelian PD, including GWAS, and made all results publicly available on a regularly updated online database, "PDGene" (<http://www.pdgene.org>).

Results

Database content

The results of this research synopsis are based on a freeze of the PDGene database content on March 31st 2011 (available upon request from the authors). At that time, PDGene included details on 828 individual studies across more than 50 different countries and six continents reporting on 3,382 polymorphisms in 890 genetic loci. Data for more than 2,000 SNPs were supplemented by results derived from up to three publicly available GWAS datasets [10,12,13] following extensive quality control and imputation. Ultimately, this procedure yielded a total of 867 polymorphisms across ~300 genetic loci that met our criteria for meta-analysis (see Methods). Additional independent GWAS data

Author Summary

The genetic basis of Parkinson's disease is complex, i.e. it is determined by a number of different disease-causing and disease-predisposing genes. Especially the latter have proven difficult to find, evidenced by more than 800 published genetic association studies, typically showing discrepant results. To facilitate the interpretation of this large and continuously increasing body of data, we have created a freely available online database ("PDGene": <http://www.pdgene.org>) which provides an exhaustive account of all published genetic association studies in PD. One particularly useful feature is the calculation and display of up-to-date summary statistics of published data for overlapping DNA sequence variants (polymorphisms). These meta-analyses revealed eleven gene loci that showed a statistically very significant ($P < 5 \times 10^{-8}$; a.k.a. genome-wide significance) association with risk for PD: *BST1*, *CCDC62/HIP1R*, *DGKQ/GAK*, *GBA*, *LRRK2*, *MAPT*, *MCCC1/LAMP3*, *PARK16*, *SNCA*, *STK39*, *SYT11/RAB25*. In addition and purely by data-mining, we identified one novel PD susceptibility locus in a gene called *ITGA8* (rs7077361; $P = 1.3 \times 10^{-8}$). We note that our continuously updated database represents the most comprehensive research synopsis of genetic association studies in PD to date. In addition to vastly facilitating the work of other PD geneticists, our approach may serve as a valuable example for other complex diseases.

for 147 SNPs yielding P values of ≤ 0.1 in these initial meta-analyses were provided by researchers of all remaining currently published Caucasian GWAS datasets [13,15–19,22]. Following the identification of genome-wide significant association with an intronic SNP (rs7077361) in *ITGA8* after addition of these data, we obtained additional data from the same GWAS datasets on ~1,400 SNPs in the chromosomal region encompassing *ITGA8* (chr10:15346353–15801533, hg18). Finally, independent replication data in Caucasian and Asian populations from the GEO-PD consortium [23] generated for ten recently described PD loci [21] were made available for inclusion. As a result, we were able to substantially increase the sample size (up to 16,452 PD cases and 48,810 controls) for a large number of some of the most promising PD loci. For instance, we were able to add data from up to 48,861 previously not analyzed combined cases and controls to meta-analyses of some of the recently proposed PD loci [21] (median sample size 14,896, see Table 2 and Table S1 for details). In addition to these focused analyses, PDGene displays meta-analysis results for more than seven million additional SNPs originating from up to three publicly available GWAS datasets [10,12,13]. The results are available online (e.g. as summarized in <http://www.pdgene.org/largescalemeta.asp>), where they are cross-linked to a customized and fully browsable track on the UCSC Genome Browser.

PDGene meta-analysis results

The PDGene meta-analyses of the 867 core polymorphisms were based on a median of 7,680 subjects (interquartile range 4,612–16,726). Additional meta-analyses were performed after stratification for Caucasian and Asian ancestry (for details on sample size and included ethnicities for individual meta-analyses see Table S1). In addition, we also performed random-effects meta-analyses across all three publicly available GWAS datasets [10,12,13] following genotype imputation using data from the International HapMap Consortium and 1000 Genomes Project. Ultimately this yielded

7,123,920 SNPs that could be meta-analyzed across at least two GWAS datasets (see Figure S1 for a quantile-to-quantile plot of the GWAS-only meta-analyses). All 867 core meta-analysis results are available online on PDGene as forest plots, summarizing the relative contributions of each dataset to the most current summary effect estimate, and in the form of cumulative plots, illustrating how summary ORs evolve over time. All meta-analysis results are plotted in Figure 1 (green dots) alongside the GWAS-only meta-analysis results (black and grey dots).

One-hundred-three meta-analyses across 12 genetic loci (*BST1*, *CCDC62/HIP1R*, *DGKQ/GAK*, *GBA*, *ITGA8*, *LRRK2*, *MAPT*, *MCCC1/LAMP3*, *PARK16*, *SNCA*, *STK39*, *SYT11/RAB25*) yielded summary ORs suggesting a genome-wide significant ($P \leq 5 \times 10^{-8}$) increase or decrease in PD risk in all ethnicities and/or after stratification for ethnic ancestry (Table 2, Table S1, and Figure S2 [forest plots]). None of these loci contained more than one SNP independently associated at genome-wide significance (as judged by pair-wise linkage disequilibrium assessments using 'SNAP' and r^2 -values of 0.2 as cut off <http://www.broadinstitute.org/mpg/snap/>). The majority of polymorphisms tested in the genome-wide significant loci do not show evidence for publication bias (Table S1). Finally, all genome-wide significant signals were robust against potential undetected sample overlap using a recently proposed procedure [24] (see Table S2 for more details). Combined sample sizes for all 12 loci were substantially larger here as compared to any previously published meta-analysis (Table S1), providing unequivocal evidence for an involvement of these loci in PD susceptibility. While power to detect genome-wide significance was excellent for most of these loci (>80% based on an OR of 1.15, and a minor allele frequency down to 0.05 using the Genetic Power Calculator, <http://pngu.mgh.harvard.edu/~purcell/gpc/>), power was less for a large number of other meta-analyses due to smaller sample sizes and allele frequencies (see Table S1 for details). Thus, no simple statistic can summarize the overall power of our study.

The above list includes an intronic polymorphism in *ITGA8* located on chromosome 10p13 for which we identified novel evidence for genome-wide association with PD risk (OR 0.88; $P = 1.3 \times 10^{-8}$, $I^2 = 0$, see Table 2, and Figure 2). This SNP had previously been proposed to be associated with PD risk at sub-genome-wide significance by Simon-Sanchez et al [13]. After obtaining and meta-analyzing GWAS data from ~1,400 additional SNPs in this region derived from all Caucasians GWAS datasets [10,12,13,15–19,21,22], rs7077361 remained the most significantly associated SNP in this region (Figure S3).

In addition to using random-effects models, we also performed exploratory fixed-effect meta-analyses on all eligible polymorphisms. These analyses did not reveal genome-wide significant effect sizes for any additional locus, except *ACMSD/TMEM163* (most significant SNP rs6723108, OR 0.91, $P = 1.3 \times 10^{-9}$, $I^2 = 46\%$ [95% CI 0–73%], Figure S4, panel 1) and *HLA* (most significant SNP chr6:32609909, OR 0.78, $P = 8.8 \times 10^{-15}$, $I^2 = 84\%$ [95% CI 70–91%], Figure S4, panel 2), both of which were reported to be associated with PD risk at genome-wide significance in previous work [16,21]. In both instances, the lack of genome-wide significance in the random-effects models (Table S1) was due to relatively pronounced heterogeneity of effect estimates across studies. However, the heterogeneity across the 11 datasets in the *ACMSD/TMEM163* meta-analysis is almost entirely due to variance of effect size estimates in the same direction (see Figure S4, panel 1), making it likely that *ACMSD/TMEM163* represents a genuine PD risk locus. For the SNP tested in the *HLA* locus (chr6:32609909, Figure S4, panel 2), heterogeneity is more pronounced and more complex owing to ORs on either side of

Table 1. Overview of genome-wide association studies (GWAS) published in PD until March 31, 2011.

GWAS	Design GWAS (Follow-up)	Population GWAS (Follow-up)	# SNPs	# PD GWAS (Follow-up)	# CTRL GWAS (Follow-up)	"Featured" genetic loci
Maraganore, 2005 (ref. 9)	Family-based (case-control)	USA-LEAPS (USA)	198,345	443 (332)	443 (332)	<i>CDCP2</i> , <i>GALNT3</i> , <i>GWA_2q36.3</i> , <i>GWA_4q28.1</i> , <i>GWA_4q28.3</i> , <i>GWA_5p15.32</i> , <i>GWA_7p14.2</i> , <i>GWA_10q21.1</i> , <i>PASD1</i> , <i>PRDM2</i> , <i>SEMA5A</i>
Fung, 2006 (ref. 10)	Case-control (-)	USA-NINDS	408,803	267 (-)	270 (-)	<i>BRDG</i> , <i>DLG2</i> , <i>GLT2SD2</i> , <i>GWA_4q35.2</i> , <i>GWA_7p12</i> , <i>GWA_10q11.21</i> , <i>GWA_11q11</i> , <i>GWA_16q23.1</i> , <i>GWA_22q13</i> , <i>NEGR1</i> , <i>ULK2</i> , <i>ZNF313</i>
Pankratz, 2009 (ref. 11)	Case-control (-)	USA-PROGENI/GenePD (-)	328,189	857 (-)	867 (-)	<i>DGKQ/GAK</i> , <i>GPRIN3</i> , <i>MAPT</i> , <i>SNCA</i>
Simon-Sanchez, 2009 (ref. 12)	Case-control (case-control)	USA-NINDS, Germany (USA, Germany, UK)	463,185	1,745 (3,452)	4,047 (4,756)	<i>LRRK2</i> , <i>MAPT</i> , <i>PARK16</i> , <i>SNCA</i>
Satake, 2009 (ref. 13)	Case-control (case-control)	Japan (Japan)	435,470	1,078 (993)	2,628 (15,753)	<i>BST1</i> , <i>LRRK2</i> , <i>PARK16</i> , <i>SNCA</i>
Edwards, 2010 (ref. 14)	Case-control (-)	USA-HIHG (-)	491,376	604 (-)	619 (-)	<i>MAPT</i> , <i>SNCA</i>
Hamza, 2010 (ref. 15)	Case-control (-)	USA-NGRC (-)	811,597	2,000 (-)	1,986 (-)	<i>GAK/DGKQ</i> , <i>HLA</i> locus, <i>MAPT</i> , <i>SNCA</i>
Spencer, 2011 (ref. 16)	Case-control (case-control)	UK-WTCCC2 (France)	1,733,533	1,705 (1,039)	5,175 (1,984)	<i>BST1</i> , <i>GAK/DGKQ</i> , <i>MAPT</i> , <i>PARK16</i> , <i>SNCA</i>
Saad, 2011 (ref. 17)	Case-control (case-control)	France (UK-WTCCC2, Australia)	492,929	1,039 (3,232)	1,984 (7,064)	<i>BST1</i> , <i>GWA_12q24</i> , <i>SNCA</i>
Simon-Sanchez, 2011 (ref. 18)	Case-control (case-control)	Netherlands	514,799	772 (-)	2024 (-)	<i>BST1</i> , <i>HLA</i> locus, <i>GAK/DGKQ</i> , <i>MAPT</i> , <i>SNCA</i>

The overview is based on content on the PDGene website (<http://www.pdgene.org>; current on March 31st, 2011). Studies are listed in order of publication date. '# PD GWAS' and '# CTRL GWAS' refers to sample sizes used in the initial GWAS datasets, whereas 'Follow-up' refers to the total number of replication samples where applicable. 'Featured genes' are those genes/loci that were declared as 'associated' in the original publication; note that criteria for declaring association varies across studies. Genetic loci in bold font denote genes showing genome-wide significant results ($P < 5 \times 10^{-8}$) in the PDGene meta-analyses. doi:10.1371/journal.pgen.1002548.t001

1. This could be due to a number of reasons, e.g. subtle and uncorrected population substructure and/or different LD patterns between the analyzed SNP and the actual functional variant(s) [16]. Thus, although the evidence is currently not as conclusive as for *ACMSD/TMEM163* it still appears quite possible that there is one or more PD association signals in the *HLA* region. Regardless of these considerations, additional data are needed to more firmly assess the role of both loci in contributing to PD susceptibility.

Ethnicity-specific meta-analysis results

SNCA, *LRRK2*, *BST1*, and *PARK16* show evidence for genome-wide significance in meta-analyses restricted to Caucasian and Asian populations (Table 2). Furthermore, data obtained from the GEO-PD consortium [23] suggest that the effect estimates for some of the recently discovered PD loci (i.e. *CCDC62/HPI1R*, *MCCI*, and *STK39*) [21] may be comparable in Caucasian and Asian populations (Table S1), although additional datasets are needed to establish genome-wide significance in populations of Asian-descent for these loci. Conversely, only insufficient data are currently available to assess the effect sizes of *GAK* and *SYT11/RAB25* on PD risk in Asians: *GAK* rs6599388 violated Hardy-

Weinberg equilibrium in Asian datasets from the GEO-PD consortium and was thus excluded from further analyses on that ethnic group [23]. *SYT11/RAB25* chr1:154105678 was excluded from all analyses due to technical reasons in the study by the GEO-PD consortium [23]. Moreover, none of the reported *SYT11/RAB25* and *GAK* SNPs from the recent GWAS meta-analysis [21] were captured directly or by proxy (with an $r^2 \geq 0.8$) in the Japanese GWAS dataset [14,23]. Finally, Asian-descent populations cannot be appropriately assessed for PD association with the *MAPT*-H1/H2 haplotype, rs10928513 in *ACMSD*, and rs7077361 in *ITGA8* owing to monomorphicity at these sites [14,23].

Evaluating the credibility of significant associations

To estimate the epidemiologic credibility of associations with polymorphisms showing sub-genome-wide significant association with PD ($P > 5 \times 10^{-8}$), we applied two "credibility" measures for each such result. First, we calculated Bayes factors (BF, expressed here as \log_{10} -values, "logBF") assuming an average non-null odds ratio of 1.15, as approximation of a typical "complex disease effect size", and a spike and smear prior distribution of effects [25]. Our

Table 2. Genome-wide significant summary meta-analysis results of the PDGene database in populations of Caucasian and Asian decent.

Caucasian ethnicity												
Locus	Polymorphism	Location (hg18)	MAF	Allele contrast	N datasets	N samples	OR (95% CI)	P-value	I ² (95% CI)	HuGENet	BF	
<i>GBA</i>	N370S	chr1:153451576	0.01	G vs. A	15	44,851	3.51 (2.55–4.83)	1.44 × 10 ⁻¹⁴	38 (0–66)	A	6.6	
<i>SYT11/RAB25</i>	chr1:154105678	chr1:154105678	0.02	T vs. C	6	17,300	1.73 (1.48–2.02)	2.35 × 10 ⁻¹²	0 (0–52)	B*	8.2	
<i>PARK16</i>	rs947211	chr1:204019288	0.23	A vs. G	12	69,262	0.91 (0.88–0.94)	8.00 × 10 ⁻¹⁰	0 (0–66)	A	6.8	
<i>STK39</i>	rs2390669	chr2:168800188	0.13	C vs. A	14	35,159	1.19 (1.12–1.25)	1.37 × 10 ⁻⁰⁹	18 (0–56)	A	4.9*	
<i>MCCC1/LAMP3</i>	rs11711441	chr3:184303969	0.14	A vs. G	25	46,502	0.86 (0.82–0.91)	9.20 × 10 ⁻¹⁰	18 (0–50)	A	6.8	
<i>DGKQ</i>	rs11248060	chr4:954359	0.12	T vs. C	10	57,716	1.21 (1.15–1.27)	3.04 × 10 ⁻¹²	11 (0–52)	A	9.2	
<i>BST1</i>	rs11724635	chr4:15346199	0.43	C vs. A	26	46,586	0.88 (0.84–0.91)	1.87 × 10 ⁻¹⁰	43 (10–64)	A	7.5	
<i>SNCA</i>	rs356219	chr4:90856624	0.41	G vs. A	31	79,494	1.29 (1.25–1.33)	6.06 × 10 ⁻⁶⁵	16 (0–46)	A	61.0	
<i>ITGA8</i>	rs7077361	chr10:15601549	0.12	C vs. T	11	61,036	0.88 (0.84–0.92)	1.51 × 10 ⁻⁰⁸	0 (0–55)	A	5.7	
<i>LRRK2</i>	rs1491942	chr12:38907075	0.21	G vs. C	21	34,123	1.17 (1.13–1.22)	6.44 × 10 ⁻¹⁵	0 (0–38)	A	11.8	
<i>CCDC62/HIP1R</i>	rs10847864	chr12:121892551	0.39	T vs. G	23	38,367	1.15 (1.11–1.18)	4.37 × 10 ⁻¹⁷	0 (0–35)	A	14.4	
<i>MAPT/STH</i>	H1H2	chr17:42131818 41149582	0.20	H2 vs. H1	37	50,389	0.78 (0.75–0.80)	7.97 × 10 ⁻⁵²	0 (0–29)	A	48.1	
Asian ethnicity												
Locus	Polymorphism	Location (hg18)	MAF	Allele contrast	N datasets	N samples	OR (95% CI)	P-value	I ² (95% CI)	HuGENet	BF	
<i>PARK16</i>	rs823156	chr1:204031263	0.17	G vs. A	5	22,870	0.74 (0.68–0.81)	2.09 × 10 ⁻¹²	0 (0–58)	A	9.2	
<i>BST1</i>	rs4538475	chr4:15347035	0.38	G vs. A	3	20,393	0.80 (0.75–0.86)	9.53 × 10 ⁻¹⁰	0 (-)	A	6.8	
<i>SNCA</i>	rs6532194	chr4:90999925	0.40	T vs. C	5	22,844	1.29 (1.20–1.39)	4.91 × 10 ⁻¹¹	31 (0–74)	A	8.0	
<i>LRRK2</i>	rs34778348	chr12:39043595	0.04	A vs. G	13	10,441	2.23 (1.89–2.63)	2.97 × 10 ⁻²¹	0 (0–53)	B*	15.2	

Whenever multiple polymorphisms showed genome-wide significant association in the same locus, only the variant with the smallest P-value is listed here. Note that, overall, 103 PDGene meta-analyses results across the 12 loci listed above yield genome-wide significant evidence for association with PD. For a complete list of these as well as the non-genome-wide significant meta-analysis results performed for the datafreeze, see Table S1. MAF = minor allele frequency in cases and controls combined; N = Number, OR = Odds Ratio; CI = confidence interval; I² = estimate of percentage of between-study heterogeneity that is beyond chance. BF = Bayes factor. *Note that additional polymorphisms in these loci showing genome-wide significant association with PD yield are graded with “strong epidemiologic credibility” (grade A) according to the HuGENet criteria [26,27], and a Bayes Factor >5 [25], respectively (see Table S1 for more details). doi:10.1371/journal.pgen.1002548.t002

second assessment was based on the Human Genome Epidemiology Network’s (HuGENet) interim criteria for the assessment of cumulative epidemiologic evidence in genetic association studies [26,27]. The results of these analyses are summarized in Table S1.

There was strong epidemiologic support in both assessments for all loci showing genome-wide significant association. This included several additional polymorphisms in these same loci that only showed sub-genome-wide significant association. However, there was no additional sub-genome-wide significantly associated locus that received unequivocally strong support from both credibility assessments (Table S1). In this list, the strongest support was assigned to SNP chr6:32588205 in the *HLA* locus receiving the best possible grade in the HuGENet criteria (grade A), but more moderate support in the Bayesian analyses (logBF = 4.4). However, the relevance of this assessment needs to be evaluated as the underlying analysis was only based on four GWAS datasets.

Discussion

The PDGene database represents a comprehensive, regularly updated and freely available online research synopsis of genetic association studies in PD. Detailed summaries of the most compelling findings are provided within an easy-to-use, dedicated online framework, displaying forest plots, cumulative meta-analyses, and an up-to-date ranking of “Top Results”. To allow

comparison of PDGene results with association findings from other complex diseases and to facilitate their interpretation with respect to functional genetics data, all meta-analysis results have been ported as a customized track onto the UCSC Genome Browser. This will also allow for an integration and visualization [28] of association results from large-scale resequencing data (e.g. from whole-exome or whole-genome studies) into PDGene once these become available.

To the best of our knowledge, our study represents the most comprehensive research synopsis in the field of PD genetics. In addition, it represents the first disease-specific genetic database that allows a systematic and exhaustive inclusion of GWAS data, and may serve as a model for similar databases in other complex genetic diseases. Owing to our multi-pronged data retrieval and analysis protocol we were able to perform meta-analyses on the vast majority of PD risk-gene candidates, including those “featured” as top association results in all published GWAS. In particular, this includes the five novel loci recently featured in the recent GWAS meta-analysis [21]. Through collaboration with other PD genetics laboratories we obtained independent summary data for these and 142 additional SNPs, substantially extending the hitherto available evidence. Taken together, our analyses provide unequivocal evidence that *BST1*, *CCDC62/HIP1R*, *DGKQ/GAK*, *GBA*, *ITGA8*, *LRRK2*, *MAPT*, *MCCC1/LAMP3*, *PARK16*, *SNCA*, *STK39*, *SYT11/RAB25* represent genuine PD risk loci, while the role of several other loci (e.g. *ACMSD*/

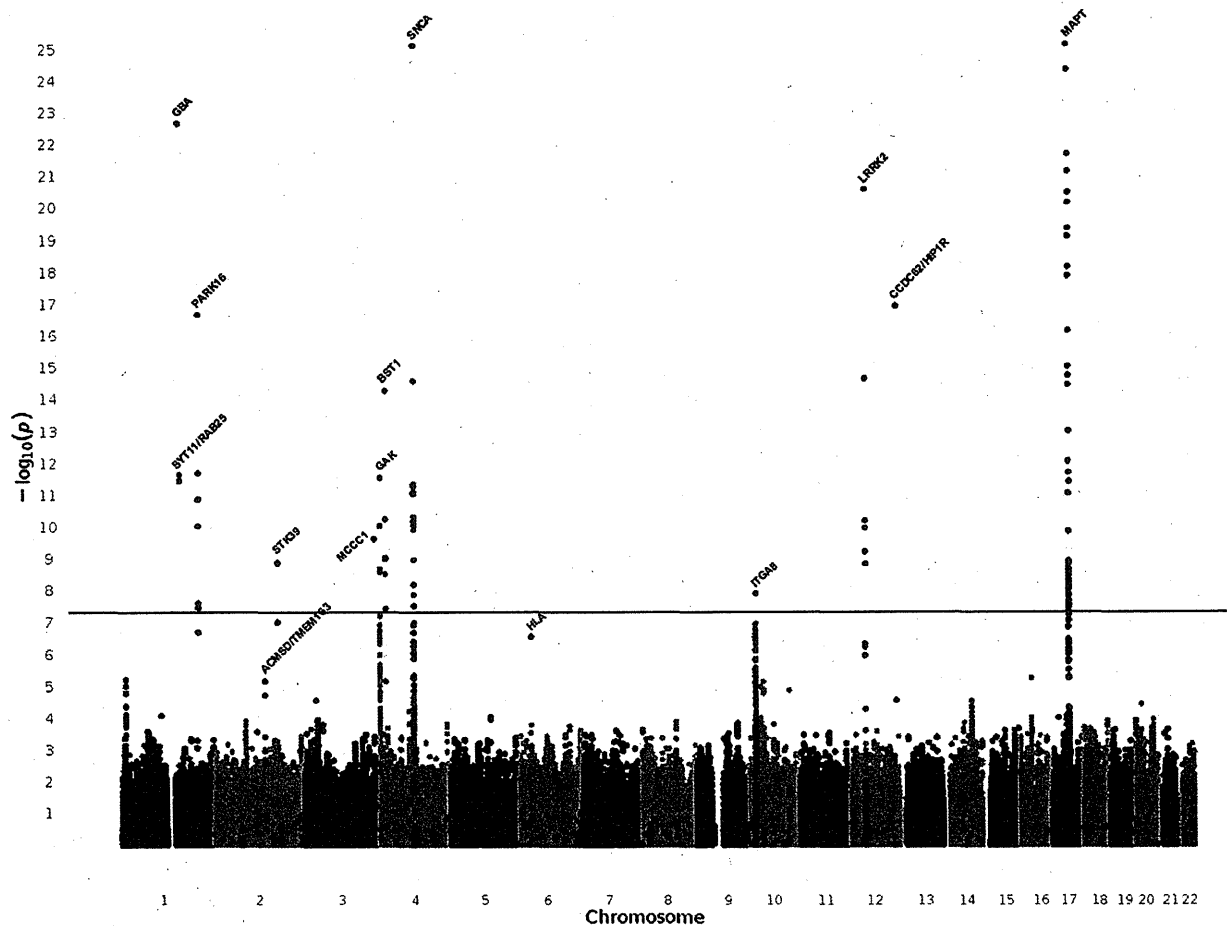


Figure 1. Manhattan plot of all meta-analysis results performed in PDGene. This summary combines association results from 7,123,986 random-effects meta-analyses based on the March 31st 2011 datafreeze of the PDGene database. Results are plotted as $-\log_{10} P$ -values (y-axis) against physical chromosomal location (x-axis). Black and grey dots indicate results originating exclusively from the three fully publicly available GWAS datasets [10,12,13] (see Methods), while green dots are based on a combination of smaller scale studies, supplemented by GWAS datasets (where applicable). Gene annotations are provided for genes highlighted in the main text. doi:10.1371/journal.pgen.1002548.g001

TMEM163, and the *HLA* locus) remains to be determined. The unpublished data aggregated here from various PD genetics groups for selected candidate genes represents the first step towards a systematic meta-analysis across the full GWAS datasets

from the same populations. Once completed, the results of this “mega” meta-analysis will be posted on the PDGene database, allowing users to browse the complete results via the customized genome browser track already in place.

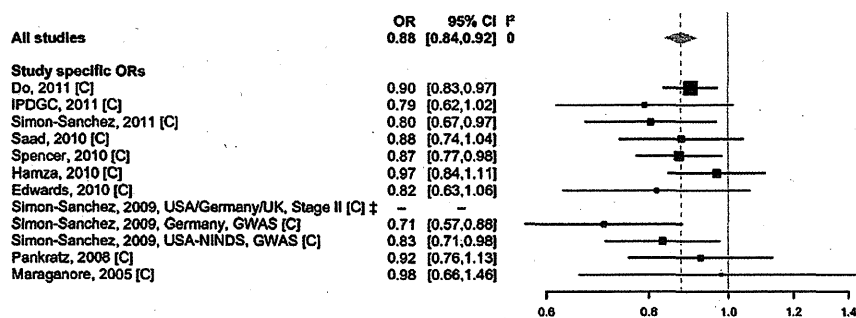


Figure 2. Forest plot of the meta-analysis of rs7077361 in *ITGA8*. Study-specific allelic odds ratios (ORs, black squares) and 95% confidence intervals (CIs, lines) were calculated for each included dataset. The summary OR and CI was calculated using the DerSimonian Laird random-effects model (grey diamond) [31]. C= Caucasian ancestry. doi:10.1371/journal.pgen.1002548.g002

Of particular interest are loci with unusually large effect sizes. While most loci in PDGene have only small effects on PD risk (with ORs ranging from 1.10 to 1.35, which are typical for complex diseases), for some loci much larger ORs were estimated (i.e. *GBA* [OR 3.51 in Caucasians], *LRRK2* [OR 2.23 in Asians], and *SYT11/RAB25* [OR 1.73 in Caucasians], see Table 2). The risk-allele frequencies at these polymorphisms are typically rather small (i.e. below 0.05), resulting in low population attributable risks for these loci (for the above mentioned loci individually less than 2%).

Interestingly, the meta-analysis results of *GBA* N370S as well as the *LRRK2* rs34778348 are solely based on candidate-gene approaches since these SNPs are not on any of the current GWAS arrays or imputation reference panels. Thus, even in the “GWAS era” smaller-scale, non-GWAS but “focused” genetic studies, will likely continue to play an important role. This is also true when it comes to providing independent replication of proposed disease associations and/or when validating imputation-derived results by direct genotyping in sufficiently sized datasets. PDGene systematically concatenates all these different types of data into one database framework, vastly facilitating an assessment of the overall evidence for any given SNP or locus.

The strength of our approach is further exemplified by the identification of genome-wide significant association between disease risk and a SNP in *ITGA8*, which was not featured as a relevant PD gene in any previous study. *ITGA8* (encoding integrin alpha 8, a type-I transmembrane protein) is functionally interesting as it is expressed in brain [29], mediates cell-cell interactions and regulates neurite outgrowth of sensory and motor neurons [30]. Additional studies are needed to further assess the potential role of this gene in PD pathogenesis. Furthermore, PDGene shows that two additional loci, not highlighted by the recent GWAS meta-analysis [21], yield genome-wide significant results in the PDGene meta-analyses, i.e. PARK16, originally implicated as a PD susceptibility locus in an Asian GWAS [14] but not highlighted in the recent GWAS meta-analysis on Caucasian samples [21] and *GBA*, a gene that was found solely by candidate-gene approaches. Another strength of our study is that it combines genetic data from currently more than 50 different countries allowing a systematic assessment of genetic associations across populations of different ethnic descent. For instance, these analyses suggest that variants in *BST1*, *LRRK2*, the PARK16 locus, and *SNCA* show genome-wide significant association with PD risk in both Caucasian and Asian-descent samples. Furthermore, the recently described Caucasian GWAS loci *CCDC62/HIP1R*, *MCCI*, and *STK39* [21] also show similar effect size estimates in populations of Asian-descent [23]. PD association data originating from other ethnic groups are still relatively scarce. However, they could easily be added to the already existing data on the respective polymorphisms available on PDGene.

In summary, we have created a continuously updated online resource for genetic association studies in the field of PD. Synthesizing essentially all available data in the field led to the identification of *ITGA8* as a novel potential PD risk locus. Our quantitative approach to data integration across a multitude of different study designs can be readily scaled to include large-scale resequencing efforts that will emerge over the coming years, making the complex field of PD genetics accessible to a broad range of investigators.

Methods

Note that the following section only provides a brief summary of the methods applied to our study. A much more detailed description can be found in Text S1.

Literature searches

Inclusion criteria. For inclusion in PDGene, a study has to meet three criteria: 1) It must evaluate the association between a bi-allelic genetic polymorphism (minor allele frequency ≥ 0.01 in the healthy control population of at least one study) and Parkinson’s disease (PD) risk in datasets comprised of both affected (defined as clinically and/or neuropathologically diagnosed “Parkinson’s disease”) and unaffected individuals; 2) it must be published in a peer-reviewed journal; 3) it must be published in English. For this manuscript, we also included data on ten SNPs generated in the GEO-PD Consortium datasets [14,23] and obtained data for the newly identified SNP rs7077361 in *ITGA8* from the Japanese GWAS dataset [14].

Exclusion criteria. In brief, genetic association data of the following studies were excluded from the meta-analyses (see Text S1 for details): family-based studies without available subject-level data (however, unrelated case-control data enriched for familial cases were not excluded), studies investigating only disease controls, multi-allelic polymorphisms, and studies of polymorphisms in mitochondrial DNA. We also excluded genetic data of apparently “poor” quality if discrepancies could not be resolved after contacting the study authors (e.g. inadequate genotyping/sequencing protocols or discrepancies in terms of allele names or frequencies when compared with public databases; more details can be found in Text S1).

Search strategies. Our literature searches until March 31st, 2011, yielded 27,210 articles, which were screened for eligibility using the title, abstract, or full-papers, as necessary. Additional screening of bibliographies in reviews, published meta-analyses, and original genetic association studies were also performed. Overall, full text versions of 1,534 articles were obtained. Following the inclusion and exclusion criteria outlined above, 828 articles were included in PDGene until March 31st 2011 (also see Figure 3).

Statistical analyses

Meta-analyses. Random-effects allelic meta-analyses [31] were performed if a minimum of four independent datasets existed per polymorphism. Summary odds ratios [ORs] and 95% confidence intervals [CIs] were calculated irrespective of ethnic descent as well as for distinct ethnic groups (i.e. Caucasians, and Asians) if sufficient data were available. In addition, we performed a number of sensitivity analyses (excluding the initial studies and datasets in which HWE was violated in control individuals), systematically assessed between-study heterogeneity (via I^2), and assessed the credibility of each at least nominally significant meta-analysis result by calculating Bayes factors (BF; here expressed as $\log_{10}(\text{BF}) = “\log\text{BF}”$) [25] and by determining a grading score developed by the Human Genome Epidemiology Network (HuGENet) [26,27].

Assessment of small-study bias/publication bias. This is of particular importance in meta-analyses of published association data and was carefully addressed here: First, we added *publicly* available GWAS data [10,12,13] to the vast majority of SNPs. Since these data are typically unbiased, this should decrease the potential for small-study bias/publication bias. Secondly, for 147 SNPs of the core PDGene meta-analyses that showed statistically suggestive results ($P \leq 0.1$), we obtained additional data from all currently published, but *not publicly* available GWAS datasets, further decreasing a potential impact of small-study bias/publication bias. Thirdly, we directly assessed the evidence for small study bias by applying a recently proposed regression test [32] on all nominally significant ($P < 0.05$) meta-analysis results. The results of these analyses are fully displayed in Table S1.

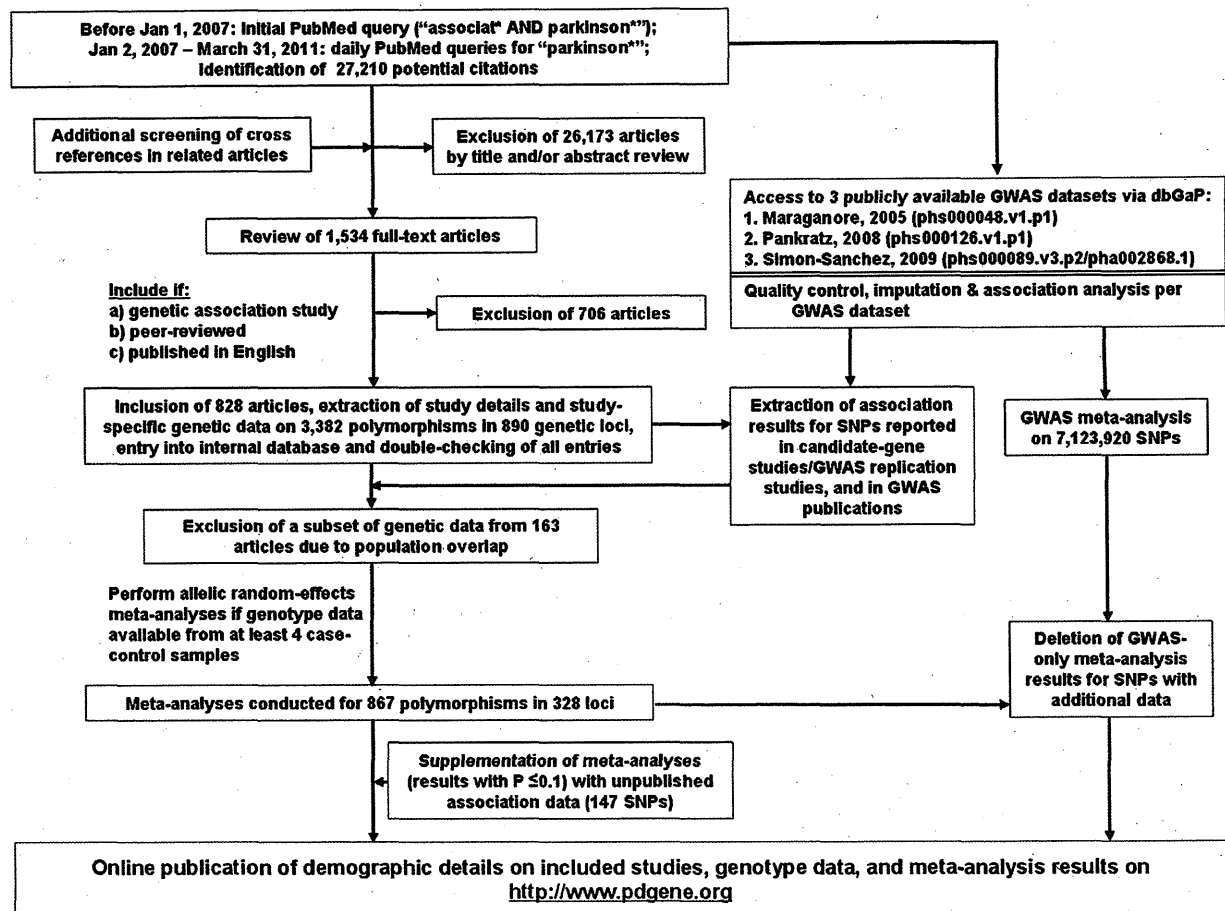


Figure 3. Flowchart of literature search, data extraction, and analysis strategies applied for PDGene.
doi:10.1371/journal.pgen.1002548.g003

GWAS-only meta-analyses. We obtained individual-level genotype data for all publicly available PD GWAS datasets from NCBI's "dbGAP" database (a total of three [10,12,13] at the time of the datafreeze, March 31st, 2011). Genotype data were cleaned using standard procedures, followed by imputation of untested genotypes (using reference panels from HapMap and the 1000 Genomes Project), and association analyses incorporating imputation uncertainty (case-control datasets only), age, sex, and population stratification. Overall, this procedure led to a total of 7,723,931 unique SNPs, 7,123,920 of which were present in at least two, and 711,271 in at least three datasets. Meta-analyses (either combining test-statistics and standard errors using random-effects models, or by combining P-values weighted by sample size, see Text S1 for more details) were performed on the 7,123,920 SNPs present in at least two of the GWAS datasets.

Online database

After completion of all data-management and analysis steps, all study-specific variables, genotype data (except for GWAS), and meta-analysis plots are posted on a dedicated, publicly available, online adaptation of the PDGene database using the same software and code as our databases for Alzheimer's disease [33] and schizophrenia [34]. The online database is hosted by the "Alzheimer Research Forum" and can be accessed via its own designated URL (<http://www.pdgene.org>).

Database code

The database software can easily be ported to other genetically complex diseases and will be made available on a collaborative basis to interested researchers upon request.

Supporting Information

Figure S1 QQ plots showing the distribution of expected versus observed P-values for the GWAS-only meta-analysis results. Analyses were performed using the METAL software (ref. [21] in Text S1). The excess of observed P-values (Figure S1, panel 1) is entirely due to association signals in the *SNCA*, *MAPT*, *LRKK2*, and *DGKQ/GAK* loci as can be seen in Figure S1, panel 2 that showcases the P-value distributions after removal of 18,622 SNPs in these regions ($\lambda = 1.007$). (TIF)

Figure S2 Forest plots of allelic meta-analyses for SNPs showing genome-wide significant association ($P < 5 \times 10^{-8}$) with PD susceptibility in the March 31st 2011 datafreeze. Study-specific allelic odds ratios (ORs, black squares) and 95% confidence intervals (CIs, lines) were calculated for each included dataset. The summary OR and CI was calculated using random-effects models (grey diamond). Whenever multiple polymorphisms showed genome-wide significant association in the same locus, only the variant with the smallest P-value is listed here for meta-

analysis results after stratification for Caucasian and Asian ancestries. For a complete list of meta-analyses performed for the datafreeze, see Table S1. Figure S1, panel 1-S1, panel 12 and S1, panel 13-S1, panel 16 display the SNP showing the most significant genome-wide association in datasets of Caucasian ancestry and Asian ancestry, respectively. Details and references of all included studies displayed here can be found on the PDGene database (<http://www.pdgene.org>). I₂ = estimate of percentage of between-study heterogeneity that is beyond chance, “excl initial” = summary OR and 95% CI after meta-analysis after exclusion of the initial study, C = Caucasian ancestry, A = Asian ancestry, H = Hispanic descent, D = African descent, “•” = initial study (applies to candidate-gene studies), “†” = no data provided or data was not eligible for inclusion in meta-analysis, “‡” = study excluded due to overlap, “#” = HWE violation in controls ($P < 0.05$, not applicable to quality-controlled GWAS datasets, see Text S1), “i” = SNP monomorphic in the respective dataset, “ø” = meta-analysis after excluding initial study not applicable.

(PDF)

Figure S3 Locus plot of the *ITGA8* region on chromosome 10p13 (15346353–15801533 bp, hg18). The figure displays association results for ~1,400 SNPs in the *ITGA8* region including at least four independent datasets. SNPs are color-coded based on linkage disequilibrium (r^2) estimates from the CEU 1000G dataset (release June 2010). All LD estimates refer to the most significantly associated SNP rs7077361. SNPs color-coded in grey indicate missing LD estimates in the CEU dataset. Recombination rates were estimated based on the CEU dataset, and are displayed as blue line in the background. Gene annotations are based on RefSeq and the UCSC Genome browser. Locus plots were generated using the LocusZoom Stand-alone package (http://genome.sph.umich.edu/wiki/LocusZoom_Standalone).

(TIF)

Figure S4 Forest plots of fixed-effect meta-analyses for SNP rs6723108 in the *ACMSD/TMEM163* locus and chr6:32609909 in the HLA locus. Symbols are the same as for Figure S2 (see above).

(TIF)

Table S1 Overview of all 867 polymorphisms meta-analyzed in the March 31st 2011 datafreeze using random-effects allelic models. Random-effects allelic meta-analyses were performed on polymorphisms for which four or more independent datasets were available. Meta-analyses after stratification for different ethnic descent were performed if at least three independent datasets were available in the respective stratum (applicable only to samples of European and Asian descent). Each nominally significant meta-analysis result ($P < 0.05$) was graded according to the HuGENet interim criteria. For details on how these criteria are applied, see Text S1. Meta-analysis results in this table are ordered by genomic location. OR = Odds Ratio, CI = confidence interval, N minor = number of minor alleles, Ethnicities: C = Caucasian, A = Asian, D = African Descent, H = Hispanic, O = Other/Mixed, Low OR = $OR < 1.15$ or ≥ 0.87 , respectively, F = loss of significance in the respective meta-analysis after exclusion of the first study, HWE = loss of significance after excluding studies violating HWE ($P < 0.05$), Regr = evidence for small-study/publication bias using a modified regression test (see Text S1), A = Grade A (‘strong’ epidemiologic credibility), B = Grade B (‘modest’ epidemiologic credibility), C = Grade C (‘weak’ epidemiologic credibility), logBF = Bayes Factor (see Text S1). “*” denotes SNPs that have been supplemented by additional data after the datafreeze (in total

this applies to 147 SNPs, see Text S1 for the description of included datasets).

(XLS)

Table S2 Investigation of the extent of statistical inflation assuming sample overlaps of 1%, 5%, and 10% across cases and controls in datasets originating from the same countries. Hypothetical sample overlap across datasets was assumed between different candidate-gene/replication studies and between candidate-gene/replication studies and GWAS datasets if they originated from the same country. These analyses were performed applying random-effects models and adding the sum of weighted co-variances of overlapping datasets to the overall study variance (see ref. [24] in the main text). Note that the assumption of undetected overlapping samples does not apply (and was therefore not modeled here) to overlap between individual GWAS as duplicate samples in these datasets were removed prior to meta-analysis. It also does not apply to independent datasets used in the same publication where duplicate samples had been removed by the authors prior to analysis and publication. We emphasize that this table describes hypothetical scenarios, because the geographical origin of each study had been investigated extensively and potentially overlapping datasets had been excluded as part of PDGene’s data inclusion protocol. Thus, the extent of overlap across geographically distinct datasets within the same countries is reduced to accidental recruitment of the same subjects more than once in different datasets throughout the respective countries, and can be expected to be less than ~1%. This estimate is based on data of the GEO-PD consortium, where sufficient data were centrally available of 6,072 subjects from 20 geographically distinct sites in 13 countries that had been investigated for potentially duplicate samples across sites, but no duplicate subjects (neither between nor within countries) were identified when matching on ethnicity, birth, sex, and genotype. The investigation of overlap was not applicable here for Asian datasets, as they originated from different countries and/or were cleaned by the respective authors prior to publication.

(DOC)

Text S1 Supplementary material. This file includes supplementary methods and references as well as the list of members of the GWAS consortia, the GEO-PD Consortium, and consortia-specific acknowledgements.

(PDF)

Acknowledgments

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Author Contributions

Conceived and designed the experiments: CM Lill, MB McQueen, JPA Ioannidis, L Bertram. Performed the experiments: CM Lill, JT Roehr, S Bagade, B-M Schjeide, E Meissner, U Zauft, NC Allen, KJ Anderson, G Beecham, D Berg, JM Biernacka, A Brice, AL DeStefano, CB Do, N

Eriksson, SA Factor, MJ Farrer, T Foroud, T Gasser, T Hamza, JA Hardy, P Heutink, C Klein, JC Latourelle, DM Maraganore, ER Martin, M Martinez, RH Myers, H Payami, WK Scott, M Sharma, AB Singleton, K Stefansson, T Toda, JY Tung, J Vance, NW Wood, CP Zabetian, 23andMe, GEO-PD, IPDGC, Parkinson's Disease GWAS, WTCC2. Analyzed the data: CM Lill, JT Roehr, MB McQueen, FK Kavvoura, L Bertram. Wrote the paper: CM Lill, JPA Ioannidis, L Bertram. Helped write the manuscript: E Meissner, MJ Farrer, T Foroud, T Gasser, C

Klein, DM Maraganore, H Payami, AB Singleton, M Sharma, F Zipp, H Lehrach. Helped analyze the data: S Bagade, T Liu, M Schilling, CB Do, N Eriksson, T Hamza, EM Hill-Burns, MA Nalls, N Pankratz, W Satake, M Sharma. Interpretation of results: CM Lill, JPA Ioannidis, L Bertram. Study coordination: CM Lill, T Foroud, JA Hardy, H Payami, AB Singleton, P Young, RE Tanzi, MJ Khoury, F Zipp, H Lehrach, JPA Ioannidis, L Bertram. Literature searches and data entry: CM Lill, S Bagade, B-M Schjeide, E Meissner, U Zauft, N Allen.

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国際共同研究における Genome-Wide Association Study (GWAS)

Genome-wide association study (GWAS) by international collaboration

戸田 達史

Key Words : GWAS, メタ解析, SNP, HapMap計画

■ Abstract ■

dbSNPやHapMap計画などの情報基盤や数十万種のSNPを解析できる技術基盤の整備により、ゲノムワイド関連解析 (GWAS) が2007年頃より実用的な戦略となり、疾患感受性遺伝子の発見が相次いだ。すでに165の疾患について、ゲノムワイド有意水準 $P < 5 \times 10^{-8}$ をクリアする904個の論文が発表された。国際共同研究によりGWASメタ解析を行い、よりeffect sizeの小さなものも同定するという試みが行われだしており、「第2世代のGWAS」とも言われている。

■ GWASの現状

2006年以降、common diseaseの感受性遺伝子の探索研究は新しい段階を迎えた。これをもたらしたおもな要因は二つの基盤整備である。まず情報基盤として、dbSNPやHapMap計画で代表されるように、ヒトゲノム全域にわたる膨大な多様性情報が集積されてきた。次に技術基盤として、数十万種のSNP (Single Nucleotide Polymorphism) を数千もの個体について並列解析できるプラットフォームが市販化された。

まずHapMap計画によると、日本人と白人は約25-30万個のタグSNP (これを調べれば連鎖不平衡で結ばれた近傍の多くのSNPの代表になる) で、ほぼ全ゲノムの遺伝子がカバーされる。つまり約30万個のタグSNPを患者と対照で調べれば、ほぼ全ゲノムの遺伝子を調べたことになる。そこで具体的には例えば患者1,000人、対照1,000人、計2,000人各人の50万個のSNPの遺伝子型を決定する。すなわちSNPチップとして2,000枚の実験を行う。それぞれSNP-1,,,,, SNP-500000ずつ、患者、対照に

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におけるそれぞれのアレルの出現頻度を合計し、偏りがないかどうかの検定を行うのである。

これらを活用することによって、ゲノムワイド関連解析 (GWAS: Genome-Wide Association Study) が実用的な戦略となり、2007年にはNature、Science誌などに立て続けに成果が発表されることとなった。その後のGWASによる疾患感受性遺伝子の発見ラッシュには目を見張るものがあり、すでに165の疾患について、ゲノムワイド有意水準 $P < 5 \times 10^{-8}$ をクリアする904個の論文が発表されている (図)。

■ GWASによる疾患感受性遺伝子発見

神経疾患のパーキンソン病を例にとると、我々のグループは患者の95%を占める孤発性PDのリスク遺伝子を同定するため、GWASを行い、PD発症に関わる2つの新しい遺伝子座PARK16、BST1を同定した。また、常染色体優性遺伝性PDの原因遺伝子 α -synuclein、LRRK2の孤発性PDへの関与を証明した。国際共同研究として次に、我々と併行してヨーロッパ起源の集団のPDのGWAS研究を行っていたグループと、データを交換した。彼らの研究では、 α -synuclein領域、Tau領域のみに、強い関連が検出されていた。そこで、我々の発見したPARK16・BST1・LRRK2の人数を増やして再現研究をしたところ、PARK16とLRRK2の関連は強く再現されたが、BST1は再現されなかった。逆に、我々は、彼らの検出したTauの関連の再現を試みた。我々の検体セットでは、再現されなかった。よって、 α -synuclein・PARK16・LRRK2は2人種に共通のPDリスクであり、Tau・BST1はのリスク多型の影響は人種特異的であると考えた¹⁾。

さらにパーキンソン病においては、さらなる国際共同研究として欧米の5つのグループがそれぞれ独立に行っていたGWASを合わせてメタ解析を行い（計患者5,333, 対照12,019）、有意なSNPをさらに患者7,053, 対照9,007で再現実験を行い、ゲノムワイド有意水準 $P < 5 \times 10^{-8}$ を超える遺伝子を従来の6個の他に、*ACMSD*, *STK39*, *LAMP3*, *SYT11*, *CCDC62*の5個を同定した²⁾。

すべては網羅できないが他の例では、クローン病においては、6個のGWASを合わせてメタ解析を行い（計患者6,333, 対照15,056）、さらに有意なSNPを患者15,694, 対照14,026で再現実験を行い、ゲノムワイド有意水準を超える遺伝子を新規に30個同定し、計71個となった³⁾。

また2型糖尿病においては、8個のGWASを合わせてメタ解析を行い（計患者8,130, 対照38,987）、さらに有意なSNPを患者34,412, 対照59,925で再現実験を行い、ゲノムワイド有意水準を超える遺伝子を新規に12個同定した⁴⁾。

さらに驚くべきことに、GIANT Consortiumは国際共同研究として46個のGWASを合わせてメタ解析を行い（計133,653人）、50,074人で再現実験を行い、ゲノムワイド有意水準を超える身長に関わる遺伝子を少なくとも180個同定した⁵⁾。

GWASによって多数の疾患感受性遺伝子が同定されたものの、それらは遺伝要因全体の一部しか説明できないことから（missing heritability）、このような圧倒的な数の試料を各地から集めてゲノムワイドメタ解析を行いよりeffect sizeの小さなものも同定するという試みが行われだしており、「第2世代のGWAS」とも言われている。

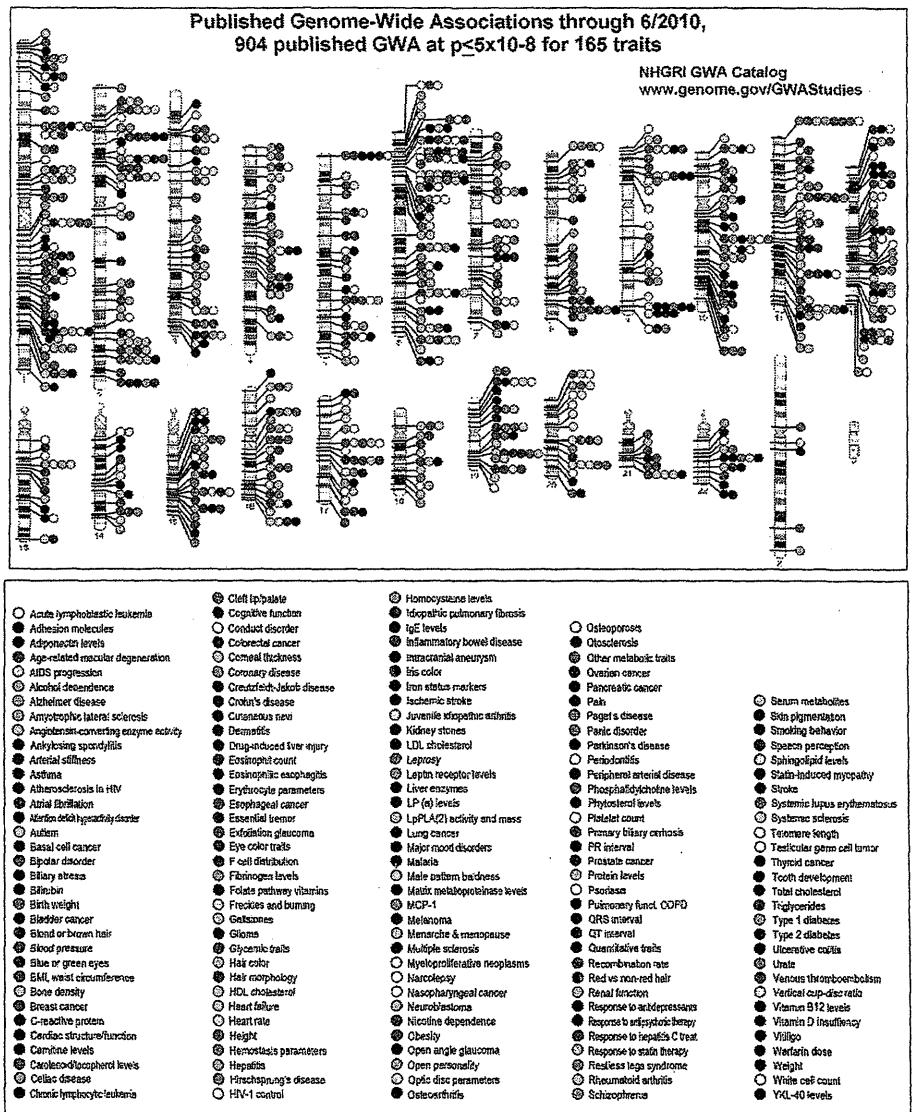


図 ゲノムワイド有意水準 $P < 5 \times 10^{-8}$ をクリアする多因子疾患の遺伝子座

一方でCommon Disease-Multiple Rare Variant仮説による、頻度は低いが発症へのEffect sizeが大きいrare variantも、missing heritabilityとして重要な位置を占める。

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6 パーキンソン病の分子遺伝学—ゲノム関連解析研究—

とだ たつし さたけわたる
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 1985年東京大学医学部卒業，神経内科学入局。94年東京大学人類遺伝学助手。96年東京大学医科学研究所助教授。2000年大阪大学臨床遺伝学教授。2008年神戸大学神経内科学／分子脳科学教授。日本人類遺伝学会賞，日本神経学会賞，朝日賞，文部科学大臣表彰。研究テーマは神経内科学，神経疾患の分子遺伝学・治療。趣味はワイン，グルメ。

Key words : パーキンソン病, α シヌクレイン, PARK16, GWAS

Abstract

患者の95%を占める孤発性PDは多因子疾患である。孤発性PDのリスク遺伝子を同定するため，ゲノムワイド関連解析を行い，PD発症に関わる2つの新しい遺伝子座PARK16, BST1を同定した。また，常染色体優性遺伝性PDの原因遺伝子SNCA, LRRK2の孤発性PDへの関与を証明した。さらに，人種差が，PDの遺伝的不均一性に，寄与していることを示唆した。国際共同研究によりGWASメタ解析を行いより多くの感受性遺伝子を同定することが行われだしており，「第2世代のGWAS」とも言われている。さらなる遺伝子の解明が期待される。

はじめに

パーキンソン病 (PD) 症例の90%以上は孤発性発症であるが、5-10%は家族性（その一部はメンデル遺伝性）に発症する。メンデル遺伝性パーキンソン病 (PD) 家系の連鎖解析などから6つのメンデル遺伝性PD原因遺伝子 (α シヌクレイン、パーキン、LRRK2遺伝子など) が明らかにされた。孤発性PD、メンデル遺伝性PDとも、一部共通の発症メカニズムが存在していると考えられ、それらを切り口にして孤発性PDの病態解明が進んでおり、ミトコンドリア障害、酸化ストレス障害の病態

への関与に加え、新たにユビキチン・プロテアソーム系の機能低下，つまり蛋白分解異常からドパミン細胞死に至る経路の重要性が示されたり。

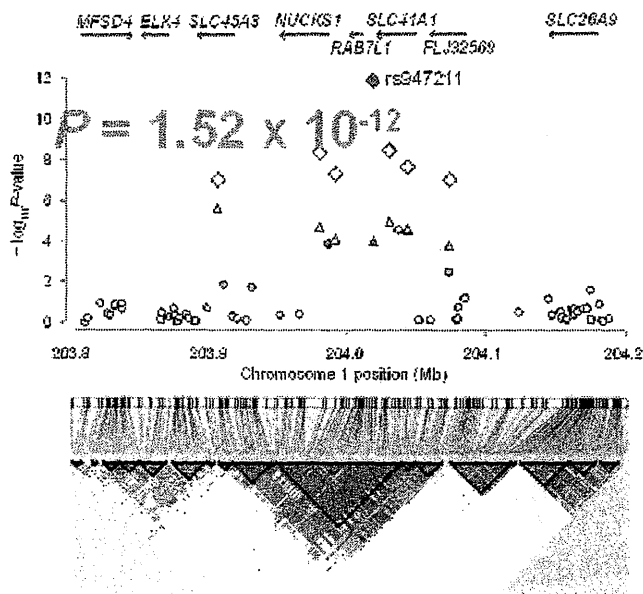
一方dbSNPやHapMap計画などの情報基盤や数十万種のSNPを解析できる技術基盤の整備により，ゲノムワイド関連解析 (GWAS) が2007年頃より実用的な戦略となり，現在多数の疾患でGWAS (ゲノムワイド関連解析) が行われており，疾患感受性遺伝子の発見がいくつかある。我々はパーキンソン病の大規模なGWASを行い，4つの感受性遺伝子を発見した¹⁾。本稿ではパーキンソン病のGWASについて述べる。

1. パーキンソン病は多因子遺伝性疾患

症例的には大多数 (95%) の孤発性PDの原因は，現時点では不明であるが，環境因子と1つ1つは影響力の弱い遺伝因子 (おそらく数10個) によってなり，その総和が，ある閾値を超えたとき発症するという多因子疾患であると考えられている。アイスランド国民を対象とした大規模な疫学的調査の結果が発表され，同胞再発危険率は6.7で，PD発症には遺伝因子が影響していることが示された。

Molecular genetics for Parkinson's disease -genomewide association study- : Tatsushi Toda, Wataru Satake, Division of Neurology, Kobe University Graduate School of Medicine, 7-5-1 Kusunoki-chou, Chuo-ku, Kobe 650-0017, Japan

a *PARK16* (1q32)



b *α-synuclein* (4q22)

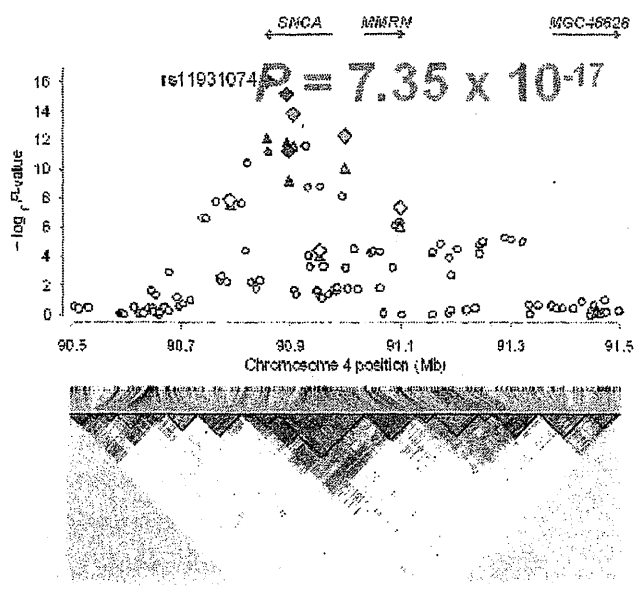


図1 日本人パーキンソン患者2011検体，対照18381検体のゲノムワイド関連解析より同定された4つのパーキンソン病遺伝子座の抜粋²⁾

PDの感受性遺伝子を発見するため，ここ10年間，多くの研究がなされてきたが⁵⁾，アルツハイマー病におけるApoE4多型のような確実に発症リスクを高める遺伝因子はなかなか確認されていなかった。ゲノムワイド有意水準 ($P < 5 \times 10^{-8}$) を満たす確実なものは， α シヌクレインの3' 非翻訳領域SNP (Gasserら³⁾，筆者ら⁴⁾が同定) とゴーシェ病遺伝子*GBA*のrare variant⁵⁾⁶⁾の2つの遺伝子のみであった。

2. パーキンソン病の GWAS (ゲノムワイド関連解析)

パーキンソン病に関しても30~50万SNPチップのGWASが数個論文報告されたが⁷⁾，小~中人数の解析であり，これらの論文では確実なことは言えなかった。我々は，大規模の患者対照集団と，56万個のSNPを搭載したイルミナHap550アレイを用いて，GWASと，2つの独立な再現研究を行い，PDの遺伝リスク因

子を明らかにしようとした。患者検体は，11施設より提供され，総数としては，患者2,011検体，対照18,381検体を用いた。まず，GWASステージとして，患者1,078検体，対照2,628検体について，それぞれ56万個のSNP型を決定した²⁾。

SNPや検体のジェノタイプ成功率などのよい患者988検体，対照2,521検体の，435,470 SNPのジェノタイプをもちいて，GWASステージのtrend検定では， $P < 5 \times 10^{-8}$ のゲノムワイド有意水準を超える遺伝子は α -synucleinのみであったものの，その一方で，Quantile-Quantile解析からは， α -synuclein以外の領域のSNPにおいても，帰無仮説のもとで予想される χ^2 統計量分布からの，有意，かつ強力なインフレーションを観察した。このことは，これらインフレートしたSNPの中に，真なる感受性座が存在していることを意味している。

そこで，GWASステージで関連を認めたp値

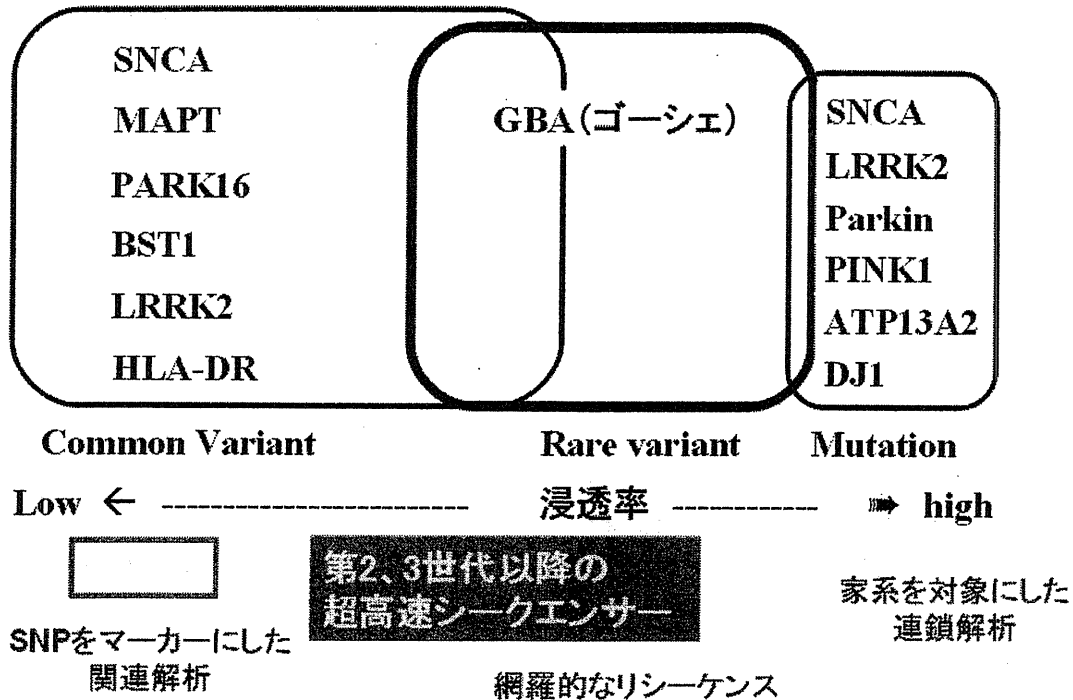


図2 パーキンソン病の遺伝メンデル遺伝性変異以外に、SNPとしてシヌクレイン (SNCA), PARK16, BST1, LRRK2, rare variantとしてゴースェ遺伝子 (GBA) が重要。

上位から337 SNPそれぞれが、2つの独立した患者・対照検体セットでも有意か、を検証する再現研究を行った。さらに全検体 (PD 2,011検体, 対照18,381検体) のジェノタイプデータを用いてメタ解析を行ったところ、絶対的な有意水準 $P < 5 \times 10^{-8}$ をクリアする4つのPD感受性遺伝子座を見いだした²⁾。

まず2つの新しいPD感受性座を、1q32 (*PARK16*と命名, $P = 1.52 \times 10^{-12}$) と4p15 ($P = 3.94 \times 10^{-9}$) に発見した。1q32領域は、3つの遺伝子 (*NUCKS1*, *RAB7L1*, *SLC41A1*) を含む連鎖不平衡ブロックであるが、発現量の形質座(eQTL)解析から、*NUCKS1*が、最も有力な責任遺伝子であると考えた (図1a)。また、4p15領域は、*BST1*のみを含んでいた。さらに、常染色体優性遺伝性PDの原因遺伝子である、 α -synuclein (4q22, $P = 7.35 \times 10^{-17}$, 図1b) と*LRRK2* (12q12, $P = 2.72 \times 10^{-8}$) の領域を同定した。 α -synucleinは、我々も以前報告した確

実なPD感受性遺伝子である。また、本研究で初めて、*LRRK2*領域のありふれた多型が、ゲノムワイド水準でPDと関連することが示された。

次にヨーロッパ起源の集団のPDのGWAS研究を行っていたグループと、データを交換した。彼らの研究では、 α -synuclein領域、*Tau*領域のみに、強い関連が検出されていた³⁾。そこで、我々の発見した*PARK16*・*BST1*・*LRRK2*の再現研究を依頼したところ、*PARK16*と*LRRK2*の関連は強く再現されたが、*BST1*は再現されなかった。逆に、我々は、彼らの検出した*Tau*の関連の再現を試みたが、我々の検体セットでは、再現されなかった。よって、 α -synuclein・*PARK16*・*LRRK2*は2人種に共通のPDリスクであり、*Tau*・*BST1*はのリスク多型の影響は人種特異的であると考えた³⁾。

また孤発性PDの発症に、常染色体優性遺伝性PDの原因遺伝子が、密に関係していること

表 日本人集団における各遺伝リスクの大きさ (寄与度)

遺伝リスクマーカー	日本人集団における リスクアレル頻度	オッズ比	人口寄与危険度 (もしこのリスクが、 日本人集団になかったら、 何%の患者が減るか)	
PARK16	rs947211	0.50	1.30	13.0
BST1	rs11931532	0.42	1.24	9.1
SNCA	rs11931074	0.58	1.37	17.6
LRRK2	rs1994090	0.08	1.39	3.0
Tau	rs393152	1.00	1.30	23.0
GBA	RecNcil	0.0018	7.21	1.1 (GBA内の11個の変異を 全てたし合わせると、4.6)

人口リスク寄与度 (もしこのリスクが日本人集団になかったら、何%の患者が減るかの指標) を示す。SNPはオッズ比は低いけど人口リスク寄与度は高い。GBAなどのrare variantはその逆である。

を明示した。原因変異により常染色体優性遺伝性PDを引き起こす遺伝子が、そのSNPを通して、孤発性PDの遺伝リスクとなったことは興味深い。このような関係は、PDを超えて、ほかの疾患に対しても、あてはまると思われる²⁾。

PARK16領域には、3つの遺伝子が存在するが、発現量的形質座解析から、**NUCKS1**が、最も有力な責任遺伝子であると考えた (図1a)。**NUCKS1**は、リン酸化部位をふくむ核タンパクであるが、神経系における機能は、未知であり、新たなPD発症のパスウェイを開拓する可能性がある³⁾。

BST1は、細胞内Ca²⁺貯蔵からのCa²⁺放出を誘発するサイクリックADPリボースの形成を触媒する酵素であり⁴⁾、最近提唱されている、ドパミン細胞死のCa²⁺ストレス説を想起させ、興味深い。これらは従来のPD病態説からは全

く新規な遺伝子であり、従来にとらわれず、新規なものを同定できるところにGWASの強みがある。

さらに最近アメリカの別グループのGWASから、HLA-DR抗原の領域との関連が報告された ($P = 1.9 \times 10^{-10}$)。PD脳ではDR抗原陽性のミクログリアが検出されること、NSAIDsは疫学的にPDのリスクを減少させることなどもあり、PDと炎症の関係を示しており興味深い¹⁰⁾。

3. 第二世代のGWAS

さらにパーキンソン病においては、さらなる国際共同研究として欧米の5つのグループがそれぞれ独立に行っていたGWASを合わせてメタ解析を行い (計患者5333, 対照12019)、有意なSNPをさらに患者7053, 対照9007で再現実験を行い、ゲノムワイド有意水準 $P < 5 \times$

10⁻⁸を超える遺伝子を従来の6個の他に、*ACMSD*, *STK39*, *LAMP3*, *SYT11*, *CCDC62*の5個を同定した¹¹⁾。

GWASによって多数の疾患感受性遺伝子が同定されたものの、それらは遺伝要因全体の一部しか説明できないことから (missing heritability), SNPとしては、このような圧倒的な数の試料を各地から集めてゲノムワイドメタ解析を行いより多くの感受性遺伝子を同定することが行われたとしており、「第2世代のGWAS」とも言われている。

おわりに

おそらく数10個あるパーキンソン病の疾患感受性遺伝子としては、 α シヌクレイン, GBA, 今回のPARK16, BST1, LRRK2など以外に確立されたものは少ないのが現状であり、今後のGWASからさらなる遺伝子の解明、そこから新たな疾患パスウェイとそこからの治療薬開発が期待される (図2)。これら4座位 (1q32, 4p15, 4q22, 12q12) の、人口リスク寄与度 (もしこのリスクが日本人集団になかったら、何%の患者が減るかの指標) を示すが、それぞれ13%, 8%, 18%, 3%と見積もられ、SNPはオッズ比は低いが人口リスク寄与度は高い。GBAなどのrare variantはその逆であり、どちらも重要である (表)²⁾。

しかし現在のSNPチップによるGWASでは、アレル頻度の低いものは同定不可能であり (搭載されていない), rare variantは見過ごされてしまう。エクソンキャプチャー, 次世代シーケンサーによるエキソームリシーケンシングが行われており、期待される。

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会長: 大内憲明 (東北大学病院がんセンター長 東北大学腫瘍外科学分野教授)

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