

Unbiased screen for interactors of leucine-rich repeat kinase 2 supports a common pathway for sporadic and familial Parkinson disease

Alexandria Beilina^a, Iakov N. Rudenko^a, Alice Kaganovich^a, Laura Civiero^b, Hien Chau^c, Suneil K. Kalia^d, Lorraine V. Kalia^e, Evy Lobbestael^f, Ruth Chia^a, Kelechi Ndukwe^a, Jinhui Ding^g, Mike A. Nalls^h, International Parkinson's Disease Genomics Consortium¹, North American Brain Expression Consortium¹, Maciej Olszewskiⁱ, David N. Hauser^{a,j}, Ravindran Kumaran^a, Andres M. Lozano^d, Veerle Baekelandt^f, Lois E. Greeneⁱ, Jean-Marc Taymans^f, Elisa Greggio^b, and Mark R. Cookson^{a,2}

^aCell Biology and Gene Expression Section, ^gComputational Biology Core, Laboratory of Neurogenetics, and ^hMolecular Genetics Section, National Institute on Aging/National Institutes of Health, Bethesda, MD 20892; ^bDepartment of Biology, University of Padova, 35131 Padua, Italy; ^cDivision of Brain, Imaging, and Behaviour Systems Neuroscience, ^dDivision of Neurosurgery, Department of Surgery, and ^eDivision of Neurology, Department of Medicine, Toronto Western Research Institute, University Health Network, University of Toronto, Toronto, ON, Canada M5T 258; ^fLaboratory for Neurobiology and Gene Therapy, KU Leuven, 3000 Leuven, Belgium; ⁱLaboratory of Cellular Physiology, National Heart, Lung and Blood Institute/National Institutes of Health, Bethesda, MD 20892; and ⁱBrown University/National Institutes of Health Graduate Partnership Program, Department of Neuroscience, Brown University, Providence, RI 02912

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Mutations in leucine-rich repeat kinase 2 (*LRRK2*) cause inherited Parkinson disease (PD), and common variants around *LRRK2* are a risk factor for sporadic PD. Using protein–protein interaction arrays, we identified BCL2-associated athanogene 5, Rab7L1 (RAB7, member RAS oncogene family-like 1), and Cyclin-G–associated kinase as binding partners of LRRK2. The latter two genes are candidate genes for risk for sporadic PD identified by genome-wide association studies. These proteins form a complex that promotes clearance of Golgiderived vesicles through the autophagy–lysosome system both in vitro and in vivo. We propose that three different genes for PD have a common biological function. More generally, data integration from multiple unbiased screens can provide insight into human disease mechanisms.

BAG5 | GAK | trans-Golgi | autophagy

enetics contribute to the pathogenesis of Parkinson disease (PD) in two ways. Mutations in several genes can cause inherited PD (1), and risk factor variants contribute to the risk of sporadic PD (2). Some genes contribute to both mechanisms. These pleomorphic risk loci (3) include genes that encode α -synuclein and leucine-rich repeat kinase 2 (*LRRK2*) (4). However, risk factors for sporadic PD identified by genomewide association studies (GWASs) (5-9) actually nominate large genomic loci with multiple candidate genes (10). These loci may include variants that change amino acids or affect disease risk through gene expression (11). Also, whether all of the genes for PD converge on a small number of biological pathways is unknown (1). It is, therefore, important to develop unbiased approaches that would resolve whether genes for PD have similar biological functions and understand the mechanism(s) of disease risk. Here, we examine one genetic cause of PD (LRRK2) and show that identifying protein interaction partners can clarify disease mechanisms.

Results

LRRK2 contains several protein–protein interaction motifs (shown diagrammatically in Fig. 1*A*). To identify protein interactions, we probed protein–protein arrays with biotinylated Glutathione-S transferase (GST)-LRRK2_{970–2527}, which was kinase-active and therefore, likely folded (*SI Appendix*, Fig. S1). Biotinylated GST was used as a negative control to exclude false positives that might nonspecifically bind to many proteins. We also included arrays where LRRK2 was coincubated with GDP or GMPPcP. These experiments recovered several candidate interactors, including

three members of the Bcl2-associated athanogene (BAG) domain cochaperones (12) (*SI Appendix*, Table S1 and Datasets S1 and S2). Of these candidates, we chose to focus on BAG5, because it is expressed in neurons and enhances dopaminergic neuronal death (13).

Some known interactors of LRRK2, such as 14-3-3 proteins (14, 15), could not have been recovered with the truncated LRRK2 probe that lacks the S910/S935 14-3-3 binding sites. Therefore, we repeated the experiments with full-length Flag-tagged LRRK2 (16) with Flag-tagged green fluorescent protein (GFP) as a negative control. We confirmed interaction with the BAG cochaperones (*SI Appendix*, Table S2 and Dataset S3) and recovered 14-3-3 proteins as well as another well-characterized LRRK2 interactor, *STUB1*/CHIP (17). This experiment also yielded several additional candidates, including two candidates that are in risk loci for sporadic PD: *Rab7L1* (5, 6) and *GAK* (7, 8). Because of our interest in the genetic basis of PD, we chose to consider Rab7L1 (RAB7, member RAS oncogene family-like 1) and Cyclin-G-associated kinase (GAK) in subsequent experiments.

We first validated individual interactions between LRRK2 and each candidate. Flag-tagged BAG5 coimmunoprecipitated endogenous LRRK2 (Fig. 1*B*), and reciprocally, endogenous

Significance

Understanding loci nominated by genome-wide association studies (GWASs) is challenging. Here, we show, using the specific example of Parkinson disease, that identification of proteinprotein interactions can help determine the most likely candidate for several GWAS loci. This result illustrates a significant general principle that will likely apply across multiple diseases.

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¹Consortia members are listed in Supporting Information.

²To whom correspondence should be addressed. E-mail: Cookson@mail.nih.gov.

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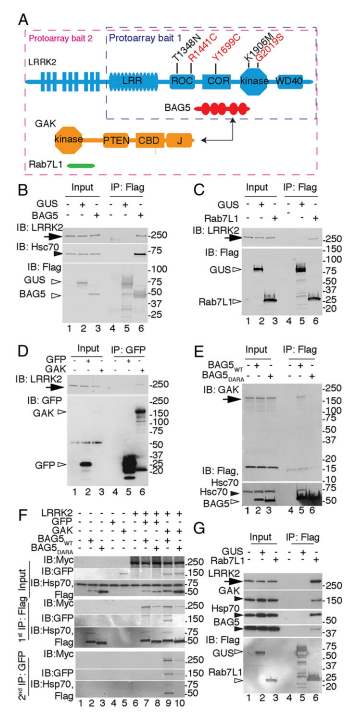


Fig. 1. A LRRK2 protein complex. (A) Ideograms of LRRK2 and candidate protein interactors. LRRK2 shows the location of the mutations used in this study, including pathogenic mutations (red) and T1348N or K1906M hypothesis-testing mutants (black). Domains include leucine-rich repeat (LRR), Ras of complex protein (ROC), and COR as well as kinase and WD40 domains. The protoarray baits used in each experiment are boxed. BAG5 has five BAG domains. GAK has PTEN-like domain (PTEN), Clathrin binding domain (CBD), and J domain. Rab7L1 has a single Ras-like domain. (B-E) Single-protein interactions. Coimmunoprecipitations using antibodies to (B and C) Flag or (D) GFP from cell lysates expressing (B) BAG5, (C) Rab7L1, or (D) GAK with (B) mock-transfected cells, (C) Flag-β-glucoronidase (GUS), or (D) GFP as negative controls probed for endogenous LRRK2 (arrows), Hsc70 (closed arrowhead), and tagged proteins (open arrowheads). IB, immunoblotting; IP, immunoprecipitation. (E) IP of BAG5 probed for endogenous GAK (arrowheads). (F and G) Cocomplex formation between LRRK2, GAK, BAG5, and Rab7L1. (F) Lysates were immunoprecipitated first with Flag for BAG5 and

BAG5 coimmunoprecipitated with LRRK2 in stable cell lines (*SI Appendix*, Fig. S24). Endogenous BAG5 coimmunoprecipitated with endogenous LRRK2 in the mouse brain (*SI Appendix*, Fig. S2B). Similarly, we found that endogenous LRRK2 coimmunoprecipitated Flag-tagged Rab7L1 (Fig. 1*C*) or GFP-tagged GAK (Fig. 1*D*). Collectively, these results show that LRRK2 can interact with three candidate proteins (BAG5, Rab7L1, and GAK), including endogenous protein in cells and the brain.

It is likely that different proteins have different sequence requirements for binding LRRK2. For example, when 14-3-3 proteins were dissociated from LRRK2 by treating with the LRRK2 kinase inhibitor LRRK2-IN1 (18), endogenous BAG5 remained bound (*SI Appendix*, Fig. S2C). To address the sequence requirements for BAG5-LRRK2 interactions further, we used a DARA mutant BAG5, where key aspartate and arginine residues in the BAG domains were replaced with alanine, diminishing heat shock cognate protein 70 (Hsc70) binding (13). DARA-BAG5 had lower binding to LRRK2 than WT BAG5 (*SI Appendix*, Fig. S2D) [mean intensity of BAG5 relative to LRRK2 (\pm SEM) of 0.49 \pm 0.023 for WT and 0.21 \pm 0.037 for DARA; P = 0.0097, two-tailed paired t test, n = 3 independent experiments], showing that recruitment of Hsc70/heat shock protein 70 (Hsp70) stabilizes the BAG5-LRRK2 complex.

Considering LRRK2, the C-terminal Ras of complex protein (COR) domain was sufficient for interaction with BAG5 (*SI Appendix*, Fig. S2E). LRRK1, with 46% similarity in the COR domain to LRRK2, also interacted with BAG5 (*SI Appendix*, Fig. S2F). These results show that the BAG5-Hsc70/Hsp70 complex is recruited to the COR domain of LRRK proteins. Adding recombinant BAG5 to LRRK2 did not affect kinase activity (*SI Appendix*, Fig. S2G).

To confirm the interactions using an additional independent technique, GST-BAG5 was used to pull down myc-LRRK2 from cells (*SI Appendix*, Fig. S3*A*). This technique also confirmed that the COR domain of LRRK2 was sufficient for BAG5 interaction (*SI Appendix*, Fig. S3*B*). Therefore, the LRRK2-BAG5 interactions were supported by protoarrays, coimmunoprecipitation, and GST pull downs.

Some LRRK2 interactors are also substrates for LRRK2 (19, 20). When testing whether BAG5 was a LRRK2 substrate, we recovered a copurifying autophosphorylated protein of ~150 kDa (*SI Appendix*, Fig. S4 *A* and *B*). Given the results in the second protoarray experiment with full-length LRRK2, we considered GAK (143 kDa) as a candidate for this unknown kinase. Supporting this hypothesis, we found that GAK coimmunoprecipitated with WT BAG5 but not DARA mutant, suggesting that this interaction is also Hsp70-mediated (Fig. 1*E*). Furthermore, endogenous BAG5 was coimmunoprecipitated with GAK in the mouse brain (*SI Appendix*, Fig. S4*C*). Deletion of the PTEN-like domain of GAK disrupted and expression of the PTEN-like domain was sufficient for interaction with both BAG5 and LRRK2 (*SI Appendix*, Fig. S4*D*).

These results do not discriminate whether BAG5 and LRRK2 bind independently to GAK or whether the three proteins are present in a cocomplex. To address these possibilities, we performed sequential coimmunoprecipitation experiments. GAK was recovered from BAG5 purification like in the single coimmunoprectipitation; then, LRRK2 was coimmunprecipitated from the GAK-BAG5 complexes (Fig. 1*F*). Furthermore, the cocomplex was more easily recovered with WT than DARA BAG5, again suggesting that Hsp70/Hsc70 stabilizes the overall assembly.

then with GFP for GAK, and they were probed for LRRK2 and interacting partners. (G) Flag-tagged Rab7L1 complexes probed for endogenous LRRK2, GAK, Hsc70, and Bag5. For all panels, markers on the right of the blots are in kilodaltons, and results are representative of at least three independent experiments.

To address whether the interactions require all components or can be formed in the absence of any one member, we knocked down each component in turn and performed coimmunoprecipitation experiments for the remaining partners. These results (*SI Appendix*, Fig. S4 *E*-*G*) showed that interactions remained even when BAG5, GAK, or LRRK2 was depleted from cells.

We next considered whether Rab7L1 might be part of the same protein complex. Rab7L1 was coimmunoprecipitated with LRRK2, and this binding was not significantly altered by mutations in LRRK2 (*SI Appendix*, Fig. S5 *A* and *B*). However, deletion of the N-terminal HEAT [Huntingtin, elongation factor 3 (EF3), protein phosphatase 2A (PP2A), TOR1] region of LRRK2 weakened interaction dramatically (*SI Appendix*, Fig. S5 *C* and *D*).

Rab7L1 is a single-domain protein, and therefore, we did not generate truncation constructs that would likely be misfolded. Instead, we asked whether LRRK2 interactions depended on the nucleotide state of Rab7L1. To this end, we generated T21N and Q67L mutants in Rab7L1 that were predicted to lack GTP binding and show impaired GTP hydrolysis based on structural modeling (*SI Appendix*, Fig. S6 *A*–*C*). As expected, T21N was deficient in GTP binding (*SI Appendix*, Fig. S6 *A*–*C*). As expected, T21N was deficient in GTP binding (*SI Appendix*, Fig. S6 *D*) [*F*(4,15), *P* = 4.5×10^{-13} , one-way ANOVA, *n* = 4 independent experiments]. Unexpectedly, Q67L had an ~10-fold higher GTP/GDP dissociation rate than WT Rab7L1 (*SI Appendix*, Fig. S6 *E* and *F*), and both mutants showed altered mobility on size exclusion chromatography (*SI Appendix*, Fig. S5 *G* and *H*). Therefore, both Rab7L1 mutants act as loss-of-function variants, likely because of failure to retain GTP. Both T21N and Q67L Rab7L1 showed

enhanced interaction with LRRK2 (*SI Appendix*, Fig. S6 *I* and *J*) [one-way ANOVA; F(4,10), $P = 4.71 \times 10^{-9}$, n = 3 independent experiments], suggesting that Rab7L1 binds LRRK2 in both the GTP- and GDP-bound states. However, GTP binding seemed to be important for some functions of Rab7L1, because both Rab7L1 mutants showed diminished (Q67L) or abolished (T21N) Golgi localization (*SI Appendix*, Fig. S6K). Endogenous Rab7L1 could simultaneously coimmunoprecipitate endogenous LRRK2, GAK, Hsp70, and BAG5 (Fig. 1*G*). Collectively, these results suggest that all five proteins are in a single complex.

We next asked whether the formation of a single-protein complex implied that these proteins have similar effects in cells. We used neurite shortening, which is exaggerated by pathogenic LRRK2 mutations (21-24). WT BAG5 caused neurite shortening, but this effect was diminished with the DARA mutant BAG5 (*SI Appendix*, Fig. S7 A–C) $[F(2,158) = 109, P < 2 \times 10^{-16}]$ by one-way ANOVA; n > 140 cells across three independent experiments]. Expression of full-length GAK, but not the deletion constructs incapable of binding LRRK2 (SI Appendix, Fig. S7D) $[F(3,215) = 5.97, P = 6.3 \times 10^{-4}$ by one-way ANOVA; n > 49 cells per construct across two independent experiments), and expression of WT Rab7L1, but not the GTP bindingdeficient Q67L variant, also decreased neurite length (SI Appendix, Fig. S7*E*) $[F(2,435) = 55, P < 2 \times 10^{-16}$ by one-way ANOVA; n > 125 cells across three independent experiments]. The effect of Rab7L1 was diminished in LRRK2 KO neurons (SI Appendix, Fig. S7F) [two-way ANOVA, F(1,281) = 21, P = 0.00076 for genotype; F(1,281) = 2.7, P = 0.06 for Rab7L1 variant; F(1,281) =21, $P = 2.04 \times 10^{-5}$ for interaction; n > 70 cells counted per

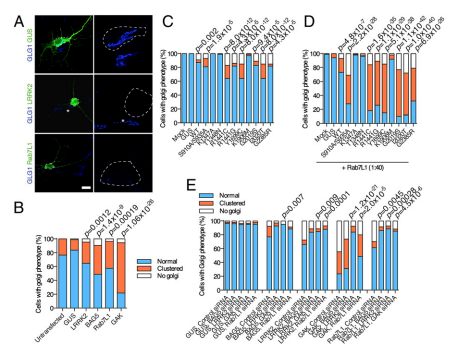


Fig. 2. (*A*) LRRK2 complex and LRRK2 mutations promote clearance of *trans*-Golgi-derived vesicles. Clustering and clearance of *trans*-Golgi by LRRK2 and interacting partners. Primary neurons were transfected with the indicated constructs and stained for transfection markers (green) and *trans*-Golgi (GLG1; blue). *Right* shows higher magnification of the circled neurons, whereas adjacent cells (asterisks) show normal Golgi morphology, including clustered Golgi in Rab7L1 (*Lower Right*) transfections and a LRRK2-transfected neuron with minimal residual Golgi staining (*Center Right*). (*B*) Blinded counts of cells with normal Golgi (blue), clustered Golgi (orange), or minimal or no apparent Golgi staining (white) as a proportion of all counted cells (n > 50 cells in three independent cultures per construct, *P* values are by Fisher exact test for proportions against GUS-transfected with indicated LRRK2 variants or GUS as a negative control and stained with antibodies to TGN46. Counting of different Golgi phenotypes after transfections with different LRRK2, BAG5 GAK, and Rab7L1. Cells were transfected with GUS (negative control for transfected with GUS (negative control for transfection), BAG5, LRRK2, GAK, or Rab7L1 and simultaneously treated with siRNA against the other components. Counting of different Golgi phenotypes after transfected with GUS (negative control for transfection), BAG5, LRRK2, GAK, or Rab7L1 and simultaneously treated with siRNA against the other components. Counting of different Golgi phenotypes after transfected with an is C and *D*.

condition), suggesting that the effects of Rab7L1 require expression of LRRK2.

We next asked where each protein might be in the cell. LRRK2 was present in microsomes as reported (25), and endogenous BAG5 and GAK were also present in the same subcellular fractions (*SI Appendix*, Fig. S84). To provide additional resolution, we expressed each interactor as tagged proteins in neurons, allowing us to use the same antibody for visualization. LRRK2 was cytosolic and vesicular (18, 26), and BAG5 was cytosolic and nuclear, whereas GAK and Rab7L1 were largely vesicular (*SI Appendix*, Fig. S8B). When coexpressed, LRRK2 and GAK showed vesicular colocalization. WT Rab7L1 promoted vesicular LRRK2 localization, but the Q67L Rab7L1 variant that was deficient in GTP binding prevented this effect (*SI Appendix*, Fig. S8C). These results show that a cocomplex of these proteins may form in vesicular compartments of the cell and that the effects require GTP binding by Rab7L1.

To further understand the possible function of the complex, we considered what type of vesicular compartments to which it might localize. Both Rab7L1 (27, 28) and GAK (29, 30) are reported to localize to the trans-Golgi network (TGN). WT Rab7L1 expressed alone (SI Appendix, Fig. S6K) or with LRRK2 (SI Appendix, Fig. S9 A and B) partially colocalized with markers of the TGN but not retromer (31) (SI Appendix, Fig. S9C). When we coexpressed both Rab7L1 and LRRK2, TGN staining was diminished and therefore, appeared clustered in the perinuclear area. Quantification showed that this phenotype was stronger for TGN (TGN46, trans-Golgi network integral membrane protein 2) than cis-Golgi (GM130, Golgin subfamily A member 2) markers (SI Appendix, Fig. S6 A and B) $[F(3,8) = 56.9, P = 9.7 \times 10^{-6}$ for TGN46; F(3,8) = 9.5, P = 0.0051 for GM130; one-way ANOVA). These results suggest that Rab7L1 directs LRRK2 to TGNderived vesicles.

In LRRK2-transfected neurons, the TGN was again clustered, or dramatically, only residual staining was visible (Fig. 24). Counting cells showed that there was a significant change in the proportions of cells with clustered or cleared Golgi after expression of LRRK2 or each interaction partner-BAG5, Rab7L1, or GAK (Fig. 2B, P values) (two-sided Fisher exact test corrected for the number of tests performed). Validating these results biochemically, HEK cells expressing LRRK2 had lower amounts of the TGN marker TGN46 (SI Appendix, Fig. S10 A and B) (P = 0.00038, t test with Welch correction, n = 5 samples) but not the *cis*-Golgi marker GM130 (*SI Appendix*, Fig. S10C) (P = 0.07). Considering the timing of these events, the recruitment of LRRK2 to TGN46-positive vesicles by Rab7L1 was complete by 24 h after transfection, but clearance of Golgi increased up to 72 h (SI Appendix, Fig. S11). Therefore, physical recruitment of LRRK2 to the TGN precedes removal of TGN-derived vesicles.

The clearance of the TGN potentially provides a cellular measure of the function of LRRK2 and binding partners. We next asked whether point mutations in LRRK2 affect Golgi clearance, thus allowing us to infer what functional roles of LRRK2 are important. All pathogenic mutations in LRRK2, including those mutations that increase kinase activity (G2019S) or decrease GTP turnover (R1441C and Y1699C), enhanced Golgi clearance (Fig. 2C, P values). However, hypothesis-testing mutations that diminish GTP binding (T1348N) or are kinase-inactive (K1906M) did not support Golgi clearance. The effects of LRRK2 mutations on Golgi clearance were enhanced by Rab7L1 (Fig. 2D, P values). These results suggest that the known enzymatic functions of LRRK2 (GTP binding and kinase activity) are important in the clearance of Golgi by a LRRK2 complex.

To define the minimum functional unit of the complex, we transfected cells with BAG5, LRRK2, GAK, or Rab7L1 and knocked down each of the other components in turn. We found that knockdown of any single component was able to rescue the effects of increased expression of other components on Golgi

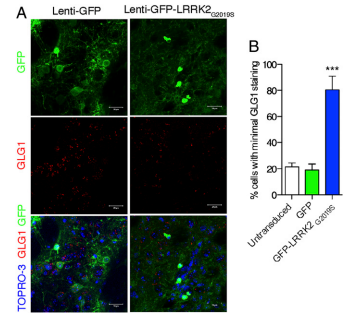


Fig. 3. Acute expression of mutant LRRK2 in adult mouse brain results in diminished staining of TGN markers in vivo. (*A*) Mouse striatum sections from animals 2 wk after injection with lentiviral vectors expressing either (*Left*) eGFP or (*Right*) G2019S mutant eGFP-LRRK2 stained for GFP (green), GLG1 (red), and TOPRO3 (blue in *Bottom*). (*B*) Counting of cells with minimal GLG1 staining in animals transduced with the vectors, with untransduced cells in the same sections also counted as controls. Error bars show SEM. ****P* < 0.001 (Tukey posthoc test from one-way ANOVA compared with eGFP control).

turnover (Fig. 2*E*, *P* values). Given that knockdown of each component did not disrupt protein interactions (*SI Appendix*, Fig. S4 *E*-*G*), these results show that, even if a complex forms, full function requires all components.

To confirm these results in vivo, we used lentiviral vectors to express enhanced green fluorescent protein (eGFP)-tagged G2019S LRRK2 in the striatum of adult mice. We found more cells with minimal staining for GLG1, a TGN marker suitable for immunostaining in rodent tissue, after expression of mutant LRRK2 compared with controls (Fig. 3) [F(2,12) = 131.4, $P = 6.93 \times 10^{-9}$; one-way ANOVA, n = 3 animals per group]. These results suggest that increased expression of LRRK2 has similar functional effects in the mouse brain as in cellular systems.

We next sought to understand the mechanisms involved in TGN turnover. The above results are reminiscent of the effects of expression and activity of another Parkinson-related protein, Parkin, that promotes removal of mitochondria by autophagy (32). LRRK2 has previously been reported to control autophagy in several systems (24, 33, 34). To test whether the autophagy-lysosome system contributed to LRRK2-dependent phenotypes, we treated cells with bafilomycin-A1, which prevents lysosomal acidification. Bafilomycin-A1 blocked the effect of LRRK2 (Fig. 4 A and B, P values) and also prevented the turnover of TGN46-positive vesicles when LRRK2 was cotransfected with GAK (*SI Appendix*, Fig. S124).

Asking whether the effects were specifically related to autophagy, we found that the effects of LRRK2 were diminished by siRNA against the autophagy protein ATG7 (Fig. 4*C*). Morphologically, cells expressing LRRK2 and treated with bafilomycin-A1 contained larger LRRK2-positive structures adjacent to lysosomes (*SI Appendix*, Fig. S12*B*). When cotransfected, both Rab7L1 and LRRK2 were found in swollen vesicles near lysosomes (*SI Appendix*, Fig. S12*C*). Some areas of TGN46 immunoreactivity

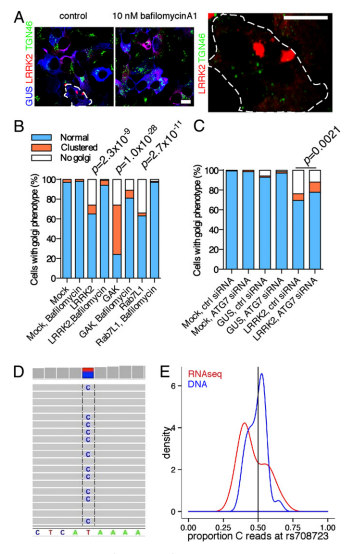


Fig. 4. Mechanism of turnover of Golgi-derived vesicles involves the autophagy–lysosomal system and is consistent with altered gene expression in the human brain. (A–C) Selective Golgi clearance involves the autophagy–lysosome system. (A) Cells coexpressing LRRK2 (red) and GUS (blue) stained for TGN46 (green) after treatment for 24 h with DMSO or 10 nM bafilomycin A1. A cell with only residual TGN46 is shown in *Right*. (B and C) Blinded counts of the proportions of Golgi phenotypes in cells treated with (B) bafilomycin A1 or (C) knockdown of Atg7. (D and E) Gene expression in the human brain. (D) Direct assay of rs708723 in Rab7L1 from the brain of an individual who was heterozygous at this marker using RNA-Seq. (E) Density plots for the proportion of reads with the C allele for RNA-Seq (red) and exome data for DNA (blue) in brains from individuals heterozygous for rs708723. There is departure from the expected proportion of 0.5 (black vertical line) for RNA-Seq data.

were enclosed by swollen LRRK2/Rab7L1-positive vesicles (*SI Appendix*, Fig. S12*D*). Collectively, these results support the hypothesis that turnover of TGN vesicles depends on the autophagy–lysosome system.

The above results make the assumption that increasing the expression of LRRK2 or its binding partners models the direction of effect of genetic risk in PD. To evaluate whether expression of these genes might explain the risk of PD in the human brain, we assayed a series of postmortem samples ($n \sim 60$) using exome sequencing and RNA-Seq. We examined the loci containing *Rab7L1* (*SI Appendix*, Fig. S13*A*), *GAK* (*SI Appendix*, Fig. S13*B*), or *LRRK2* (*SI Appendix*, Fig. S13*C*) for sequence

variants. Although we reliably detected expression of *LRRK2*, *GAK*, and *Rab7L1* in the human cerebral cortex, many pathogenic variants were too rare to be analyzed or in regions that did not code for RNA. However, rs708723 in the 3' UTR of *Rab7L1* could be identified directly from RNA-Seq data (Fig. 4*D*). There was lower expression of the *C allele* in RNA-Seq data in heterozygous carriers, and the proportions of *C alleles* in the aggregate counts were significantly different between exome and RNA-Seq data (Fig. 4*E*) (χ^2 test, $\chi^2 = 5.9$, *P* = 0.015). The *C allele* of rs708723 was associated with a lower risk of PD (odds ratio = 0.914, *P* = 0.000554 under an additive model). Therefore, higher expression of normal sequence variants is a plausible mechanism for risk of PD, at least for *Rab7L1*.

Discussion

Here, we used an unbiased survey of protein interactions of LRRK2 to identify binding partners and show that these proteins form a cocomplex that promotes TGN removal by autophagy-dependent mechanisms. TGN clearance also occurs in the brain in vivo, and expression levels of some of the protein components are a reasonable mechanism for risk of sporadic PD. Our results overall lead us to propose that several PD genes act in a common pathway (*SI Appendix*, Fig. S14).

Because LRRK2 has several proposed interaction domains, it may act as a scaffold for multiple interaction partners. Although we recovered known direct binding partners of LRRK2 (15, 35) and inferred indirect interaction with Hsp70 (17), not all proposed literature candidates were identified. This result may indicate a false-negative rate with arrays, which do not include all splice variants of all proteins in the human genome or all posttranslational modifications. The approach used here should be combined with other methods, which has been suggested more generally for understanding protein function (36).

With these caveats, our approach led us to show that LRRK2 promotes the relocalization to and clearance of *trans*-Golgi-derived vesicles. This effect is presumably related to the known function of LRRK2 in autophagy (18, 24, 33, 37, 38) but can be tied mechanistically to disruption of TGN. Interestingly, autophagic vesicles may derive from the *trans*-Golgi network (37); hence, altered turnover of Golgi-derived vesicles would impact autophagy function over time, consistent with effects of LRRK2 deficiency in animal models (34).

Critically, this activity is enhanced by all pathogenic mutations in LRRK2. However, LRRK2 mutants do not alter binding to its partner proteins. This observation suggests that the enzymatic activity of LRRK2, including GTP binding and kinase activity, may be critical for activity of the complex. Supporting this idea, hypothesis-testing mutations lacking GTP binding or kinase activity do not promote clearance of the TGN-derived structures. It is possible that, after being bound, protein interaction partners also influence either LRRK2 kinase activity or GTP affinity. Defining the stoichiometry of the complex and effects on LRRK2 is an important future goal.

The other proteins each seem to be required for LRRK2 to affect Golgi clearance by autophagy. We have shown that removal of BAG5, GAK, or Rab7L1 does not prevent LRRK2 binding with other partners, implying that formation of a partial complex is not sufficient for Golgi clearance. Within the complex, we speculate that each protein may play a specific role. It is of interest that Rab7L1 is localized to the *trans*-Golgi network, likely in its GTP-bound form, where it might recruit LRRK2 and other components to cooperatively cause TGN to be engulfed by the autophagosomes. The presence of BAG5 and Hsp70/Hsc70 suggests that this process requires remodeling of proteins or protein complexes. GAK is also known to cooperate to remodel clathrin complexes (39).

Our results have the broadest implications for mechanisms of understanding disease risk for loci identified by GWASs. Higher expression of Rab7L1 is associated with higher risk of PD in humans, expression quantitative trait locus identified by microarrays, thus providing a reasonable mechanism for the action of the WT allele. We cannot exclude that other genes at each GWAS locus may contribute to disease risk, because several of these loci are complex genetically (9). However, identification of GAK and Rab7L1 as binding partners of LRRK2 from an unbiased screen raises the probability that these genes explain some of the GWAS signal. This concept has important implications for other diseases that have been studied by GWASs, because it supports the idea that integrating datasets from multiple unbiased approaches leads to unexpected insights into human diseases.

Materials and Methods

Detailed materials and methods are included in *SI Appendix*. Briefly, identification of protein interactors was achieved by screening two batches of Protoarrays (Invitrogen) with either truncated GST-LRRK2₉₇₀₋₂₅₂₇ (Invitrogen) or full-length protein prepared as described (16). Protein interactions were validated using coimmunoprecipitation in HEK293FT cells or mouse brain

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and also GST pull downs in cells or brain. Subcellular fractionation was performed as described (25) using LRRK2-transfected HEK293FT cells. Protein localization and effects on turnover of TGN were performed using primary postnatal cortical neurons with calcium–phosphate transfection (40) or in HEK293FT cells. All counting experiments were performed by an operator blinded to the transfection construct used; details of blinding and statistical approaches are included in *SI Appendix*.

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