## TITLE

Regulation of mitophagy by the NSL complex underlies genetic risk for Parkinson's disease at Chr16q11.2 and on the MAPT H1 allele

## AUTHOR INFORMATION

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## RUNNING TITLE

New mitophagy Parkinson’s risk genes

ABSTRACT
Parkinson's disease (PD) is a common incurable neurodegenerative disease. The identification of genetic variants via genome-wide association studies (GWAS) has considerably advanced our understanding of the PD genetic risk. Understanding the functional significance of the risk loci is now a critical step towards translating these genetic advances into an enhanced biological understanding of the disease. Impaired mitophagy is a key causative pathway in familial PD, but its relevance to idiopathic PD is unclear. We used a mitophagy screening assay to evaluate the functional significance of risk genes identified through GWAS. We identified two new regulators of PINK1-mitophagy, KAT8 and KANSL1, previously shown to modulate lysine acetylation. These findings establish PINK1-mitophagy as a contributing factor to idiopathic PD. KANSL1 is located on chromosome 17q21 where the risk associated gene has long been considered to be MAPT. Our data provide evidence that this assignment is likely to be incorrect and that variability at KANSL1 underpins this association. Finally, these results enrich our understanding of physiological events regulating mitophagy and establish a novel pathway for drug targeting in neurodegeneration.

## KEY WORDS

GWAS / KANSL1 / KAT8 / mitophagy / Parkinson’s disease

## INTRODUCTION

Parkinson's disease (PD) is the most common movement disorder of old age and afflicts more than 125,000 in the UK ${ }^{1}$. Temporary symptomatic relief remains the cornerstone of current treatments, with no disease-modifying therapies yet available ${ }^{2}$. Until recently, the genetic basis for PD was limited to family-based linkage studies, favouring the identification of rare Mendelian genes of high penetrance and effect. However, genome-wide association studies (GWAS) have identified large numbers of common genetic variants linked to increased risk of developing the disease ${ }^{3,4}$. While these genetic discoveries have led to a rapid improvement in our understanding of the genetic architecture of $P D^{5}$, they have resulted in two major challenges for the research community. First, conclusively identifying the causal gene(s) for a given risk locus, and secondly dissecting their contribution to disease pathogenesis. Addressing these challenges is critical for moving beyond genetic insights to developing new disease-modifying strategies for PD.

Previous functional analyses of PINK1 and PRKN, two genes associated with autosomal recessive $P D$, have highlighted the selective degradation of damaged mitochondria (mitophagy) as a key contributor to disease pathogenesis. In mammalian cells, the mitochondrial kinase PINK1 selectively accumulates at the surface of damaged mitochondria, where it phosphorylates ubiquitin, leading to the recruitment of the E3 ubiquitin ligase Parkin. The recruitment of autophagy receptors leads to the engulfment of damaged mitochondria in autophagosomes, and ultimately fusion with lysosomes ${ }^{6-11}$. It has subsequently become clear that other PD-associated Mendelian genes, such as FBXO7, DJ-1 and VPS35 ${ }^{12}$, are implicated in the regulation of PINK1-mediated mitochondrial quality control. Based upon these data, we hypothesised that PD-GWAS candidate genes may also be involved in this process, providing a mechanistic link between these genes and the aetiology of idiopathic PD. In order to test that hypothesis, we used functional genomics to prioritise candidate genes on the PD GWAS loci, and we developed a biological screening assay as a tool to identify genes that regulate PINK1-mitophagy, and as such, are very likely to be genes that increase the risk of developing PD.

In this study, we show that KAT8 and KANSL1, two genes that were previously shown to be part of the same lysine acetylase complex partially located at the mitochondria ${ }^{13}$, are two new important regulators of PINK1-mediated mitochondrial quality control. These findings establish mitophagy as a contributing factor to idiopathic PD and provide a proof of principle for the value of screening approaches to identify causative genes in GWAS loci. Finally, these results suggest lysine acetylation as a potential new avenue for mitophagy modulation and therapeutic intervention.

## RESULTS

Genomic analyses of PD have identified over 80 loci associated with an increased lifetime risk for disease. In contrast to Mendelian PD genes, however, the assignment of a causative gene to a risk locus is often challenging. In order to identify new risk genes for PD, we undertook a triage of PD GWAS candidate genes using a combination of methods: i) Colocalization (Coloc) and transcriptome-wide association analysis (TWAS) ${ }^{14}$ using expression quantitative trait loci (eQTLs) information derived from Braineac ${ }^{15}$, GTEx and CommonMind resources ${ }^{16,17}$ to link PD risk variants with specific genes, ii) weighted protein-protein interaction (PPI) network analysis (WPPINA) ${ }^{18}$ based on Mendelian genes associated with PD, and iii) the prioritised gene set as described in PDGWAS ${ }^{3,19} .31$ open reading frames (ORFs) were nominated as putatively causal for associations at PD risk loci. 55\% of these genes were prioritised through multiple techniques (Fig. 1a), with three out of 31 genes (KAT8, CTSB and NCKIPSD) identified through all three prioritization methods (Fig. 1b, c). The 31 genes, together with 7 PD Mendelian genes and lysosomal storage disorder genes, previously shown to be enriched for rare, likely damaging variants in PD ${ }^{20}$, were then taken forward for functional analysis (Supplementary Table 1).

Based upon extensive data implicating impaired mitophagy in the aetiology of familial PD ${ }^{7}$, we hypothesized that additional PD-GWAS candidate genes, involved in the most common, idiopathic form of the disease, may play a role in this process. In order to test whether the 38 prioritised genes have a role in PINK1-mitophagy, we developed and optimized a high content screening (HCS) assay for phosphorylation of ubiquitin at serine 65 ( $\mathrm{pUb}(\mathrm{Ser} 65$ )), a PINK1-dependent mitophagy marker ${ }^{21}$, following mitochondrial depolarization (Extended Data Fig. 1a and 2). The 38 prioritised genes were individually knocked down (KD) using siRNA in Parkin over-expressing (POE)SHSY5Y human neuroblastoma cells. Increased mitochondrial clearance following mitochondrial depolarization induced by treatment with $10 \mu \mathrm{M}$ of oligomycin/antimycin $\mathrm{A}(\mathrm{O} / \mathrm{A})$ was validated as an endpoint for mitophagy (Extended Data Fig. 1b). Over 97\% of the pUb(Ser65) signal colocalised with the TOM20 mitochondrial marker in O/A treated cells (Extended Data Fig. 1c, d). siRNA KD efficiency was validated using both a pool of PINK1 siRNA, which decreased O/A induced pUb(Ser65) and subsequent TOM20 degradation (Extended Data Fig. 1e-g) without decreasing cell viability (Extended Data Fig. 3a, b), and a pool of Polo-like kinase 1 (PLK-1) siRNA that decreased cell viability by apoptosis (Extended Data Fig. 3a, b). The siRNA pools for the 38 candidate genes, together with controls, were screened in duplicate on each plate, across three replicate plates per run. Hits were identified as those wells where O/A-induced pUb(Ser65) was decreased or increased at greater than two standard deviations from the mean of the scramble (SCR) negative control siRNA.

KAT8 was selected based on reproducible downregulation of O/A-induced PINK1-dependent pUb(Ser65) across all three replicates (Fig. 2a and Extended Data Fig. 1h), without affecting cell
viability (Extended Data Fig. 3c). Notably, KAT8 was selected as a candidate gene on the basis of all three prioritization criteria - namely, proximity of the lead SNP to an ORF, colocalization of a brainderived eQTL signal with a PD GWAS association signal (Extended Data Fig. 4) and evidence of PPI with a known PD gene (Fig. 1). Furthermore, we find that colocalization and Transcriptome-wide Association Study (TWAS) ${ }^{22}$ analyses at this locus are consistent with the KD models in the HCS assay (Supplementary Tables 2 and 3) ${ }^{17}$. Both methods predict that the risk allele operates by reducing KAT8 expression in PD cases versus controls. The effect of KAT8 KD on pUb(Ser65) was further validated in POE SHSY5Y cells treated with either 1 or $10 \mu \mathrm{M} \mathrm{O/A}$, using both immunofluorescence (IF) and immunoblotting (IB) (Fig. 3a-d and Extended Data Fig. 5a-e). In order to assess whether other lysine acetyltransferases (KATs) could regulate PINK1-dependent mitophagy, the pUb(Ser65) screen was repeated in POE SHSY5Y cells silenced for 22 other KATs ${ }^{23,24}$. Only KAT8 KD led to a decreased pUb(Ser65) signal, emphasising the specificity of the KAT8 KD effect on pUb(Ser65) (Fig. 3e and Supplementary Table 4).

These functional data complement and support the Omic prioritization of KAT8 as a causative gene candidate for the chromosome 16q11.2 PD associated locus (Fig. 2b). To gain further insight into a possible role for KAT8 in the aetiology of PD, we explored the known functional interactions of this protein. KAT8 has previously been shown to partially localise to mitochondria as part of the NSL complex together with KANSL1, KANSL2, KANSL3, and MCRS1 ${ }^{13}$ (Fig. 4a). To test whether other components of the NSL complex also modulate mitophagy, the pUb(Ser65) screen was repeated in POE SHSY5Y cells silenced for each of the nine NSL components (HCFC1, KANSL1, KANSL2, KANSL3, KAT8, MCRS1, OGT, PHF20, WDR5) (Fig. 4a). Notably, reduction of KANSL1, KANSL2, KANSL3, MCRS1 and KAT8 expression led to decreased pUb(Ser65) after 1.5 or 3 h O/A treatment (Fig. 4b, c). Strikingly, KANSL1 is another PD GWAS candidate gene ${ }^{3}$. The effect of KANSL1 KD on pUb(Ser65) was further validated in POE SHSY5Y cells treated with $1 \mu \mathrm{M} \mathrm{O/A}$, using both IF and IB (Fig. 5a-d). A time course experiment was designed to compare the effect of KAT8 and KANSL1 KD on PINK1-dependent pUb(Ser65) (Fig. 6a,b and Supplementary table 5) and subsequent mitophagy (Fig. 7a,b). While individual KD of either KANSL1 or KAT8 affect phosphorylation of ubiquitin, KANSL1 KD decreased $\mathrm{pUb}(\mathrm{Ser} 65)$ and mitophagy, as assessed by measuring the puncta of the mitochondrial marker PMPCB ${ }^{25}$, more efficiently than KAT8 KD (Fig. 6a,b and Fig. 7a,b). To assess the role of KAT8/KANSL1 in neuronal function and survival in vivo, we used Drosophila as a simple model system. Notably, the NSL complex was originally discovered in Drosophila through the homologs of KAT8 and KANSL1 (mof and nsl1, respectively), but null mutants for these genes are associated with developmental lethal owing to profound transcriptional remodelling during development ${ }^{26}$. Therefore, we utilised inducible transgenic RNAi strains to target the KD of mof and
$n s / 1$ specifically in neuronal tissues. Using behavioural assays as a sensitive readout of neuronal function we found that pan-neuronal KD of mof or ns/1 caused progressive loss of motor (climbing) ability (Extended data Fig. 6a, b), and also significantly shortened lifespan (Extended data Fig. 6c, d). Interestingly, loss of $n s / 1$ had a notably stronger effect than loss of mof. Consistent with this, KD of ns/1 but not mof, in either all neurons or only in dopaminergic (DA) neurons, caused the loss of DA neurons (Extended data Fig. 6e, f).

KANSL1 is located within the extensively studied inversion polymorphism on chromosome 17q21 (Extended data Fig. 7a, b), which also contains MAPT - a gene frequently postulated to drive PD risk at this locus ${ }^{27}$. While the majority of individuals inherit this region in the direct orientation, up to $25 \%$ of individuals of European descent have a $\sim 1 \mathrm{mb}$ sequence in the opposite orientation ${ }^{28,29}$, inducing a larger $\sim 1.3-1.6 \mathrm{Mb}$ region of linkage disequilibrium (LD). Since this inversion polymorphism precludes recombination over a region of $\sim 1.3-1.6 \mathrm{Mb}$, haplotype-specific polymorphisms have arisen resulting in the generation of two major haplotype clades, termed H1 (common haplotype) and H 2 (inversion carriers), previously strongly linked to neurodegenerative disease ${ }^{30,31}$. Due to high LD, the genetics of this region have been hard to dissect, and robust eQTL analyses have been challenging due to the issue of polymorphisms within probe sequences in microarray-based analyses or mapping biases in RNA-seq-based analyses. Several variants (rs34579536, rs35833914 and rs34043286) are in high LD with the $\mathrm{H} 1 / \mathrm{H} 2$ haplotype and are located within KANSL1 (Fig. 8a,b), which could directly impact on KANSL1 protein function. In particular, one of the missense variants is a serine to proline change in KANSL1 protein sequence (S718P), and would therefore be predicted to alter the gross secondary structure of the KANSL1 protein (Fig. 8b). Furthermore, we used allele-specific expression (ASE) analysis to explore the possibility that PD risk might be mediated at this locus through an effect on KANSL1 expression. Using RNA sequencing data generated from 84 brain samples (substantia nigra $n=35$; putamen $n=49$ ), for which we had access to whole exome sequencing and SNP genotyping data thus enabling mapping to personalised genomes ${ }^{32}$, we quantified the variation in expression between the H 1 and H 2 haplotypes (Supplementary Table 6). While we identified ASE sites within MAPT (Extended Data Fig. 8 and Supplementary Table 7), we also identified 4 sites of allele-specific expression in KANSL1 (Fig. 8a), suggesting that the high PD risk H 1 allele is associated with lower KANSL1 expression, consistent with our functional assessment. Interestingly, sequence analysis of the human KANSL1 haplotype revealed that the high risk H1 haplotype is the more recent "mutant" specific to Homo sapiens, and that other primates and mammals share the rarer non-risk ancestral H 2 haplotype (Figure 8b).

To assess the specificity of the KANSL1 KD effect on PINK1-mitophagy, 32 open reading frames in linkage disequilibrium on the H 1 haplotype at the 17q21 locus (Extended data Fig. 7a, b
and Supplementary Table 8) were knocked down individually and their effect on $\mathrm{pUb}(\operatorname{Ser} 65)$ was assessed. While the effect of KANSL1 KD on pUb(Ser65) was confirmed, neither the KD of MAPT, nor the KD of each of the other 30 genes on this locus, led to a decreased in the pUb(Ser65) signal (Fig. 9). These data confirm the selectivity of our mitophagy screening assay and suggest that KANSL1 is likely to be a key PD risk gene at the 17q21 locus.

## DISCUSSION

Since the first PD GWAS study was performed in $2006{ }^{33}$ GWAS have identified about 90 independent loci for PD ${ }^{4}$. However, translating GWAS findings into a new molecular understanding of PDassociated pathways and new therapeutic targets has remained a major challenge for the scientific community. In order to screen for PD GWAS candidate genes that play a role in PINK1-mitophagy, and thus are likely to be genuine risk genes for PD, we have set up and optimised a HCS for pUb(Ser65), a marker of PINK1-dependent mitophagy, a key pathway in PD pathogenesis. This approach allowed the successful identification of two new genes associated with increased PD risk, that play a role in mitophagy. Interestingly, these two genes were previously shown to be part of the same complex, the NSL complex.

This study demonstrates the substantial potential of functional screens to exploit genetic data by providing orthogonal information that can confidently identify new risk genes. This is particularly important in genomic regions with uniformly high linkage disequilibrium, such as the $17 q 21$ inversion region which includes 32 ORFs of which many are highly expressed in brain and where existing finemapping and functional genomic analyses have been inconclusive. Interestingly, while MAPT has long been considered the risk associated gene at this locus, this has recently been questioned by Dong and colleagues, who also raised the significance of KANSL1 in driving PD risk at the locus ${ }^{34}$. Furthermore, functional screening can simultaneously provide mechanistic insights as exemplified in this case by the novel insights we provide into the molecular events regulating mitochondrial quality control and which support a role for mitophagy as a contributing factor to sporadic PD. While our data demonstrate an unequivocal role for KAT8 and KANSL1 in PINK1-dependent ubiquitin phosphorylation and subsequent mitophagy, the precise mechanism of regulation remains to be determined. Our results identify KAT8 and KANSL1 as modifiers of PINK1-dependent pUb(Ser65), suggesting that they may be regulating PINK1 kinase activity. It will be important to determine whether phosphorylation of other PINK1 substrates, such as Parkin and Rab8A ${ }^{9,35}$, are modulated by KAT8 and KANSL1. The KAT8 and KANSL1-containing NSL complex functions to promote histone acetylation and as such, is a master regulator of transcription ${ }^{36}$. Therefore, another possibility is that KAT8/KANSL1 may regulate PINK1 transcription, and subsequent translation. Interestingly, depletion of KAT8/KANSL1 was shown to cause significant downregulation of mitochondrial DNA transcription and translation, and ultimately impaired mitochondrial respiration ${ }^{13}$. Thus, it is also
possible that KAT8/KANSL1-dependent modulation of mitochondrial DNA indirectly regulates PINK1 mitochondrial accumulation and subsequent mitophagy. Finally, an intriguing possibility is that the KAT8/KANSL1 complex directly acetylates ubiquitin, which has previously been shown to be acetylated on six out of its seven lysines (K6, K11, K27, K33, K48, K63) ${ }^{37}$. It has been proposed that the KAT8/KANSL1 complex has targets in the mitochondria other than the mitochondrial DNA ${ }^{13}$. Further experiments are required to determine whether KAT8, which has been shown to partially localise at the mitochondrial outer membrane, where ubiquitin is phosphorylated ${ }^{13}$, may contribute to ubiquitin acetylation.

Important genetic discoveries in PD, in particular, the identification of the PINK1 ${ }^{38}$ and PRKN genes ${ }^{39}$, opened the field of selective mitophagy ${ }^{7}$. However, there is still a clear need for a better molecular understanding of mitochondrial quality control. Here we provide new insights into the mechanism by identifying two new molecular players, KAT8 and KANSL1. These new regulators of mitophagy provide the first direct evidence for a role of the PINK1-mitophagy pathway in idiopathic PD and the convergence between familial and idiopathic pathways in disease. Taken together, these findings open a novel avenue for the therapeutic modulation of mitophagy in PD, with potential implications across drug discovery in frontotemporal dementia and Alzheimer's disease, where mitophagy also plays an important role in disease pathogenesis ${ }^{40}$.

## METHODS

## Reagents

Oligomycin (mitochondrial complex V inhibitor) was purchased from Cayman Chemicals (11341) and from Sigma-Aldrich (O4876), and antimycin A (mitochondrial complex III inhibitor) was purchased from Sigma-Aldrich (A8674). All siRNAs were purchased as pre-designed siGENOME SMARTpools from Dharmacon: non-targeting (D-001206-13), PINK1 (M-004030-02), PLK1 (L-003290-00), KIF-11 (L-003317-00), KAT8 (M-014800-00), KANSL1 (M-031748-00), KANSL2 (M-020816-01), KANSL3 (M-016928-01), HCFC1 (M-019953-01), MCRS1 (M-018557-00), OGT (M-019111-00), PHF20 (M-01523402), WDR5 (M-013383-01). The following antibodies were used for immunocytochemistry: mouse anti TOM20 (Santa Cruz, sc-17764, 1:1000), rabbit anti phospho-ubiquitin (Ser65) (Cell Signaling, 37642, 1:1000), rabbit anti PMPCB (Proteintech 16064-1-AP, 1:1000), AlexaFluor 488 goat anti rabbit (Invitrogen, A11008, 1:2000), AlexaFluor 568 goat anti mouse (Invitrogen, A11004, 1:2000). The following antibodies were used for immunoblotting: mouse anti TIM23 (BD Biosciences, 611223, 1:1000), rabbit anti phospho-ubiquitin (Ser65) (Merck Millipore, ABS1513-I, 1:1000), mouse anti GAPDH (Abcam, ab110305, 1:1000), rabbit anti KAT8 (Abcam, ab200600, 1:1000), IRDye 680LT donkey anti mouse (LI-COR Biosciences, 925-68022, 1:20000), IRDye 800CW donkey anti rabbit (LICOR Biosciences, 925-32213, 1:20000).

## Selection of genes for High Content Screening

Candidates for High Content Screening were selected based on i) WPPINA; ii) complex prioritization; and, iii) coloc analysis. WPPINA analysis is reported in ${ }^{18}$ where the 2014 PD GWAS ${ }^{19}$ was analysed; candidate genes where selected among those prioritised and with an LD r2 $\geq 0.8$. The same pipeline has then been additionally applied to the 2017 PD GWAS ${ }^{3}$ to update the list of candidate genes. Briefly, a protein-protein interaction network has been created based on the Mendelian genes for PD (seeds) using data from databases within the IMEx consortium. The network has been topologically analysed to extract the core network (i.e. the most interconnected part of the network). The core network contains the proteins/genes that can connect $>60 \%$ of the initial seeds and are therefore considered relevant for sustaining communal processes and pathways, shared by the seeds. These processes have been evaluated by Gene Ontology Biological Processes enrichment analysis. The top SNPs of the 2017 PD GWAS have been used to extract open reading frames (ORFs) in cis-haplotypes defined by LD r2 $\geq 0.8$ (analysis performed in October 2017) ${ }^{3}$. These ORFs have been matched to the core network to identify overlapping proteins/genes in relevant/shared pathways. Results of complex prioritization (neurocentric prioritization strategy) were gathered from
${ }^{3}$ where this strategy was applied to the 2017 PD GWAS ${ }^{3}$. The coloc analysis was performed as reported in ${ }^{17}$, posterior probabilities for the hypothesis that both traits, the regulation of expression of a given gene and the risk for PD share a causal variant (PPH4), were calculated for each gene, and genes with PPH4 $\geq 0.75$ were considered to have strong evidence for colocalization. Summary statistics were obtained from the most recent PD GWAS ${ }^{4}$ and were used for regional association plotting using LocusZoom ${ }^{41}$.

## Cell Culture and siRNA transfection

POE SH-SY5Y cells are a kind gift from H. Ardley ${ }^{42}$. Cells were cultured in Dulbecco's Modified Eagle (DMEM, Gibco, 11995-065) and supplemented with 10\% heat-inactivated foetal bovine serum (FBS, Gibco) in a humidified chamber at $37^{\circ} \mathrm{C}$ with $5 \% \mathrm{CO}_{2}$. For siRNA transfection, cells were transfected using DharmaFECT1 transfection reagent (Dharmacon, T-2001-03) according to the manufacturer's instructions (for concentrations of siRNA, see sections below).

## ASEs

Sites of ASE were identified as described by Guelfi and colleagues ${ }^{32}$ by mapping RNA-seq data to personalised genomes, an approach specifically chosen because it aims to minimise the impact of mapping biases. RNA-seq data generated from 49 putamen and 35 substantia nigra tissue samples from the UK Brain Expression Consortium was used for this analysis. All samples were obtained from neuropathologically normal individuals of European descent and sites with greater than 15 reads in a sample were tested for ASE. Only sites present in non-overlapping genes were considered and data from both the tissues were considered together to increase power. Sites with minimum FDR $<5 \%$ across samples were marked as ASE sites. Plots were generated using Gviz3, with gene and transcript details obtained from Ensembl v92.

## High Content siRNA Screen

## Cell plating and siRNA transfection

siRNA was dispensed into Geltrex-coated 96-well CellCarrier Ultra plates (Perkin Elmer) at a final concentration of 30 nM using the Echo 555 acoustic liquid handler (Labcyte). For each well, $25 \mu \mathrm{l}$ of DMEM containing $4.8 \mu \mathrm{l} / \mathrm{ml}$ of DharmaFECT1 transfection reagent was added and incubated for 30 min before POE SH-SY5Y cells were seeded using the CyBio SELMA (Analytik Jena) at 15,000 cells per well, $100 \mu$ l per well in DMEM $+10 \%$ FBS. Cells were incubated for 72 h before treatment with 10 $\mu \mathrm{M}$ oligomycin $/ 10 \mu \mathrm{M}$ antimycin for 3 h to induce mitophagy.

IF and Image Capture and Analysis

Cells were fixed with 4\% PFA (Sigma-Aldrich, F8775), then blocked and permeabilised with 10\% FBS, $0.25 \%$ Triton $\mathrm{X}-100$ in PBS for 1 h , before immunostaining with $\mathrm{pUb}(\operatorname{Ser} 65)$ and TOM20 primary antibodies (in $10 \%$ FBS/PBS) for 2 h at room temperature. After 3 x PBS washes, AlexaFluor 568 antimouse and 488 anti-rabbit secondary antibodies and Hoechst 33342 (Thermo Scientific, 62249) were added (in 10\% FBS/PBS, 1:2000 dilution for all) and incubated for 1 h at room temperature.

Following a final $3 x$ PBS washes, plates were imaged using the Opera Phenix (Perkin Elmer). $5 x$ fields of view and $4 x 1 \mu \mathrm{~m}$ Z-planes were acquired per well, using the 40X water objective, NA1.1. Images were analysed in an automated way using the Columbus 2.8 analysis system (Perkin Elmer) to measure the integrated intensity of pUb(Ser65) within the whole cell (see Extended Data Fig. 1 and 2 for image analysis workflow and parameters).

## Screen quality control, data processing and candidate selection

Screen plates were quality controlled based on the efficacy of the PINK1 siRNA control and O/A treatment window (minimum 3-fold). Data were checked for edge effects using Dotmatics Vortex visualization software. Raw data was quality controlled using robust $Z$ prime $>0.5$. Data were processed using Python for $Z$ score calculation before visualization in Dotmatics Vortex. Candidates were considered a hit where $Z$ score was $\geq 2$ or $\leq-2$, and where replication of efficacy was seen both within and across plates.

## siRNA libraries

The siRNA libraries were purchased from Dharmacon as an ON-TARGETplus SMARTpool Cherry-pick siRNA library, 0.25 nmol in a 384 -well plate. siRNAs were resuspended in RNase-free water for a final concentration of $20 \mu \mathrm{M}$. SCR, PINK1 and PLK1 or KIF11 controls were added to the $384-$ well plate at a concentration of $20 \mu \mathrm{M}$ before dispensing into barcoded assay-ready plates.

## Mitochondrial enrichment and Western blotting

POE SH-SY5Y cells were transfected with 100 nM siRNA and incubated for 72 h . Cell lysates were fractionated into cytoplasmic and mitochondria-enriched samples, and run on SDS-PAGE before IB with the Odyssey ${ }^{\circledR}$ CLx Imager (LI-COR Biosciences). Mitochondrial enrichment and Western blotting protocols were described previously ${ }^{43}$.

## Immunofluorescence

POE SH-SY5Y cells were reverse transfected with 50 nM siRNA in 96 -well CellCarrier Ultra plates according to the manufacturer's instructions and incubated for 72 h . Cells were then treated, fixed and stained as per the screening protocol detailed above (for treatment concentrations and times, see figures). Image intensities were not modified or pre-processed. Images are presented as
maximum projections of the $\mathrm{pUb}(\operatorname{Ser} 65)$ or PMPCB-488 channel for one field of view. Insets show the Hoechst 33342 channel for the same field.

## Drosophila stocks and husbandry

Flies were raised under standard conditions in a humidified, temperature-controlled incubator with a $12 \mathrm{~h}: 12 \mathrm{~h}$ light:dark cycle at $25^{\circ} \mathrm{C}$, on food consisting of agar, cornmeal, molasses, propionic acid and yeast. The following strains were obtained from the Bloomington Drosophila Stock Center (RRID:SCR_006457): mof RNAi lines, P\{TRiP.JF01701\} (RRID:BDSC_31401); and P\{TRiP.HMS00537\} (RRID:BDSC_58281); ns/1 RNAi lines, P\{TRiP.HMJ22458\} (RRID:BDSC_58328); the pan-neuronal nSybGAL4 driver (RRID:BDSC_51941); and dopaminergic neuron driver (TH-GAL4; RRID:BDSC_8848); and control (/acZ) RNAi P\{GD936\}v51446) from the Vienna Drosophila Resource Center (RRID:SCR_013805). All experiments were conducted using male flies.

## Locomotor and lifespan assays

The startle induced negative geotaxis (climbing) assay was performed using a counter-current apparatus. Briefly, 20-23 males were placed into the first chamber, tapped to the bottom, and given 10 s to climb a 10 cm distance. This procedure was repeated five times (five chambers), and the number of flies that has remained into each chamber counted. The weighted performance of several group of flies for each genotype was normalized to the maximum possible score and expressed as Climbing index ${ }^{44}$.

For lifespan experiments, flies were grown under identical conditions at low-density. Progeny were collected under very light anaesthesia and kept in tubes of approximately 20 males each, around 50100 in total. Flies were transferred every 2-3 days to fresh medium and the number of dead flies recorded. Percent survival was calculated at the end of the experiment after correcting for any accidental loss.

## Immunohistochemistry and sample preparation

Drosophila brains were dissected from aged flies and immunostained as described previously ${ }^{45}$. Adult brains were dissected in PBS and fixed in $4 \%$ formaldehyde for 30 min on ice, permeabilized in $0.3 \%$ Triton X-100 for 3 times 20 min , and blocked with $0.3 \%$ Triton X-100 plus $4 \%$ goat serum in PBS for 4 h at RT. Tissues were incubated with anti-tyrosine hydroxylase (Immunostar Inc. \#22491), diluted in $0.3 \%$ Triton $\mathrm{X}-100$ plus $4 \%$ goat serum in PBS for 72 h at $4^{\circ} \mathrm{C}$, then rinsed 3 times 20 min with $0.3 \%$ Triton X-100 in PBS, and incubated with the appropriate fluorescent secondary antibodies overnight at $4^{\circ} \mathrm{C}$. The tissues were washed 2 times in PBS and mounted on slides using Prolong

Diamond Antifade mounting medium (Thermo Fisher Scientific). Brains were imaged with a Zeiss LMS 880 confocal. Tyrosine hydroxylase-positive neurons were counted under blinded conditions.

## Statistical Analysis

Intensity measurements from imaging experiments were normalised to SCR O/A for each experiment and presented as a percentage. N numbers are shown in figure legends and refer to the number of independent, replicate experiments. Within each experiment, the mean values of every condition were calculated from a minimum of 3 technical replicates. Intensity measurements from Western blot experiments were normalised to PINK1 O/A. GraphPad Prism 6 (La Jolla, California, USA) was used for statistical analyses and graph production. Data were subjected to either one-way or two-way ANOVA with Dunnett's post-hoc analysis for multiple comparisons, unless otherwise stated. All error bars indicate mean $\pm$ standard deviation (SD) from replicate experiments.

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## Author Contributions.

HPF, PL, JH, AW and PW conceived the idea. MS, DM, EA, AM, DT, MB, PW, JH, AW, MR, PL and HPF designed the experiments. MS, DM, EA, AM, NW, NW, KDS, SG, DZ, AP, DT, KP, CM, CB and HPF carried out analysis and experiments. MS, DM, EA, AM, PW, CM, AW, MR, PL and HPF wrote the manuscript, with input from all co-authors. HPF, PL and MR supervised the project.

Competing Interests: The authors declare that they have no conflict of interest

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## FIGURE LEGENDS

Figure 1 - Selection of the genes for the PINK1-dependent mitophagy screening.
A. The heat-map represents increasing evidence for gene prioritization (white, light blue, and dark blue: one, two, and three evidences, respectively). ColB = coloc analysis using Braineac, ColG = coloc analysis using GTEx, WPPINA = weighted protein interaction network; GWAS = genes prioritised in PD-GWAS ${ }^{3}$.
B. Genes prioritised by means of multiple prediction techniques, grouped based on the types of evidences.
C. Venn diagram highlighting the three genes prioritised by means of three prediction techniques.

Figure 2 - High content mitophagy screen identifies KAT8 as a modulator of pUb(Ser65) levels.
A. pUb(Ser65) Z-scores of one representative mitophagy screen plate.
B. Overview of the PD GWAS genetic signal at the KAT8 locus.

Figure 3 - KAT8 knockdown decreases pUb(Ser65).
A. Representative images of $p \mathrm{Ub}(\mathrm{Ser} 65)$ following treatment of SCR, PINK1 and KAT8 siRNA KD POE SH-SY5Y with $1 \mu \mathrm{M} \mathrm{O/A}$ for 3 h . Insets show the nuclei for the same fields. Scale bar: $20 \mu \mathrm{~m}$.
B. Quantification of $p U b(S e r 65)$ levels in $A(n=3$, two-way ANOVA with Dunnett's correction).
C. Representative IB of mitochondrial fractions from SCR, PINK1 and KAT8 KD POE SH-SY5Y treated with $1 \mu \mathrm{M} \mathrm{O} / \mathrm{A}$ for 1.5 or 3 h .
D. Quantification of $p \mathrm{Ub}(\mathrm{Ser} 65)$ in $\mathrm{C}(\mathrm{n}=5$, one-way ANOVA with Dunnett's correction).
E. pUb(Ser65) Z-scores of one representative KAT screen plate. See Supplementary Table 4 for the complete list of the genes screened.

Data are shown as mean $\pm$ SD.

Figure 4 - Knockdown of the mitochondrial components of the NSL complex reduces pUb(Ser65) levels.
A. Schematic representation of the NSL complex.

B, C. Quantification of pUb(Ser65) following treatment of SCR, PINK1 or NSL components siRNA KD POE SH-SY5Y cells with $1 \mu \mathrm{M}$ O/A for $1.5 \mathrm{~h}(\mathrm{~B})$ or $3 \mathrm{~h}(\mathrm{C})$. Data are shown as mean $\pm \mathrm{SD} ; \mathrm{n}=6$, one-way ANOVA with Dunnett's correction.

Figure 5 - KANSL1 knockdown decreases pUb(Ser65).
A. Representative images of $\mathrm{pUb}(\mathrm{Ser65}$ ) following treatment of SCR, PINK1 and KANSL1 KD POE SHSY5Y cells with $1 \mu \mathrm{M} \mathrm{O} / \mathrm{A}$ for 3 h . Insets show the nuclei for the same fields. Scale bar: $20 \mu \mathrm{~m}$.
B. Quantification of $\mathrm{pUb}(\mathrm{Ser} 65)$ in $\mathrm{A}(\mathrm{n}=3$, two-way ANOVA with Dunnett's correction).
C. Representative IB of mitochondrial fractions from SCR, PINK1 and KANSL1 KD POE SH-SY5Y treated with $1 \mu \mathrm{M} \mathrm{O/A}$ for 1.5 or 3 h .
D. Quantification of $p U b(S e r 65)$ in $C$ ( $n=5$, one-way ANOVA with Dunnett's correction). Data are shown as mean $\pm$ SD.

Figure 6 - KANSL1 and KAT8 knockdown decrease pUb(Ser65).
A. Representative images of $\mathrm{pUb}(\mathrm{Ser} 65)$ (green) following treatment of SCR, PINK1, KAT8 and KANSL1 KD POE SH-SY5Y cells with $1 \mu \mathrm{M} \mathrm{O/A} \mathrm{for} \mathrm{0-7} \mathrm{h} .\mathrm{Insets} \mathrm{show} \mathrm{the} \mathrm{nuclei} \mathrm{for} \mathrm{the} \mathrm{same} \mathrm{fields}$. Scale bar: $20 \mu \mathrm{~m}$.
B. Quantification of $p \mathrm{pb}(\mathrm{Ser} 65)$ in $\mathrm{A}(\mathrm{n}=6$, two-way ANOVA with Dunnett's correction). Data are shown as mean $\pm$ SD.

Figure 7 - KANSL1 and KAT8 knockdown decrease mitochondrial clearance.
A. Representative images of PMPCB (green) and nuclei (blue) following treatment of SCR, PINK1, KAT8 and KANSL1 KD POE SH-SY5Y cells with $1 \mu \mathrm{M}$ O/A for $0,3,6,9 \mathrm{~h}$. Scale bar: $10 \mu \mathrm{~m}$.
B. Quantification of the number of PMPCB spots in $A$ ( $n=5$, two-way ANOVA with Dunnett's correction).

Data are shown as mean $\pm$ SD .

Figure 8 - ASE sites in KANSL1 in LD with the H1/H2 SNP.
A. ASEs derived from putamen and substantia nigra in high linkage disequilibrium with the $\mathrm{H} 1 / \mathrm{H} 2$ tagging SNP, rs12185268 and their position along the KANSL1 gene. The missense variants track displays the variants annotated as missense by gnomAD v2.1.1 ${ }^{46}$. The valid track displays the heterozygous sites (orange $=$ missense) with an average read depth greater than 15 reads across all samples, which were examined for ASE. The topmost track displays the FDR-corrected minimum $\log 10 p$-value across samples for the sites that show an ASE in at least one sample.
B. Conservation of the KANSL1 protein across species. The four coding variants in the KANSL1 gene are in high LD ( $\mathrm{r} 2>0.8$ ) with the $\mathrm{H} 1 / \mathrm{H} 2$ haplotypes.

Figure 9. High content mitophagy screening of the ORFs on the 17 q21 locus identifies only KANSL1 as a modulator of $\mathrm{pUb}($ Ser65 ) levels.
pUb(Ser65) Z-scores of one representative 17q21 locus screen plate. See Supplementary Table 8 for the complete list of the genes screened.

## EXTENDED FIGURE LEGENDS

## Extended Data Figure 1. High Content siRNA Screen for modulators of pUb(Ser65).

A. Workflow of the high content screen for O/A-induced pUb(Ser65) levels.
B. Fold decrease in TOM20 levels following 1.5 and 3 h treatment with $0.1,1$ and $10 \mu \mathrm{MO} / \mathrm{A}$, compared to DMSO control.
C. Representative images of TOM20 and pUb(Ser65) following 3 h treatment of SCR KD POE SH-SY5Y cells with $10 \mu \mathrm{M}$ O/A. Scale bar: $20 \mu \mathrm{~m}$.
D. Quantification of the co-localization in $\mathbf{C}$ as \% of TOM20-positive pUb(Ser65) spots. Graph shows all replicates of non-transfected, SCR, PINK1 and PLK1 KD for 3 independent experiments.
E. Representative images of pUb(Ser65) following treatment of SCR and PINK1 KD POE SH-SY5Y cells with $10 \mu \mathrm{M} \mathrm{O} / \mathrm{A}$ for 3 h . Scale bar: $20 \mu \mathrm{~m}$.
F. Quantification of pUb(Ser65) in E ( $\mathrm{n}=6$, two-way ANOVA with Tukey's multiple comparisons test).
G. Representative analysis of integrated intensity of pUb(Ser65)and TOM20 for a single HCS plate.
H. pUb(Ser65) Z-scores of the two other replicate screen plates.

Data are shown as mean $\pm$ SD.

## Extended Data Figure 2. Image processing workflow of the high content screen for O/A induced pUb(Ser65).

Images were loaded as maximum projections. The Hoechst 33342 channel was used to find the nuclei and border nuclei were excluded. The cytoplasm was then found on the calculated Hoechst + TOM20-568 image. The pUb(Ser65)-488 spots were identified within the whole cell and their intensity measured. The outputs of the analysis were the number of nuclei selected and the mean integrated $\mathrm{pUb}($ Ser65 ) intensity, calculated as the area of the cell covered by $\mathrm{pUb}($ Ser65 ) spots x corrected intensity of the spots. Analysis parameters for each building block of the Columbus workflow are detailed in the boxes.

## Extended Data Figure 3. KAT8 knockdown has no effect on cell viability.

A. Representative images of nuclei following treatment of SCR, PINK1 and PLK1 siRNA KD POE SHSY5Y cells with $10 \mu \mathrm{M} \mathrm{O} / \mathrm{A}$ for 3 h . Scale bar: $20 \mu \mathrm{~m}$.
B. Quantification of the number of nuclei in A ( $n=6$, two-way ANOVA with Tukey's multiple comparisons test).
C. Z-scores of a representative screen plate showing that KAT8 or PINK1 siRNA KD don't affect cell viability, on the contrary to PLK-1 KD.

Data are shown as mean $\pm$ SD.

## Extended Data Figure 4. KAT8 eQTLs colocalise with SNPs associated with PD risk

The x-axis displays the physical position on chromosome 16 in megabases. The minus log $p$-values are plotted for every SNP present in both the PD GWAS ${ }^{3}$ and KAT8 eQTLs derived from the GTEx V7 caudate data. The $p$-values for the PD GWAS are plotted in yellow and $p$-values for KAT8 eQTLs are plotted in blue.

Extended Data Figure 5. KAT8 knockdown decreases pUb(Ser65) levels.
A. Representative images of pUb(Ser65) following treatment of SCR, PINK1 and KAT8 siRNA KD POE SH-SY5Y with $10 \mu \mathrm{M} \mathrm{O/A}$ for 3 h . Insets show nuclear staining for the same fields. Scale bar: $20 \mu \mathrm{~m}$. B. Quantification of $\mathrm{pUb}(\mathrm{Ser} 65)$ levels in $\mathrm{A}(\mathrm{n}=3$, two-way ANOVA with Dunnett's correction).
C. Representative IB of mitochondrial fractions from SCR, PINK1 and KAT8 KD POE SH-SY5Y treated with $10 \mu \mathrm{M} \mathrm{O/A}$ for 1.5 or 3 h .

D, E. Quantification of $p \mathrm{Ub}(\mathrm{Ser} 65)(\mathbf{D})$ and KAT8 levels $(\mathbf{E})$ in $C$ ( $n=5$, one-way ANOVA with Dunnett's correction).

Data are shown as mean $\pm$ SD.

Extended Data Figure 6. Neuronal loss of mof or nsl1 causes locomotor deficit, shortened lifespan and neurodegeneration.

A, B. Climbing ability of pan-neuronal (nSyb-GAL4) driven knockdown of mof (A) or ns/1 (B) measured at the indicated age of adults, compared to control RNAi (A: Kruskal-Wallis test, with Dunn's post-hoc multiple comparisons; B: Mann-Whitney test).

C, D. Lifespan of mof (C) or nsl1 (D) pan-neuronal knockdown (nSyb-GAL4) compared to control RNAi (Log-rank (Mantel-Cox) test).

E, F. Quantification of dopaminergic neurons (PPL1 cluster) after pan-neuronal or dopaminergic (DA) neuron (TH-GAL4) driven depletion of $\operatorname{mof}(\mathbf{E})$, nsl1 (F), or control RNAi. Representative images of PPL1 neurons (as bounded by the box) under depletion conditions are shown. Flies were aged 30 days, except for pan-neuronal ns/1 kd which are 16-days-old. Scale bar: $20 \mu \mathrm{~m}$; Mann-Whitney test. For all tests, n numbers are indicated in the graphs; $\mathrm{p}<0.0001={ }^{* * * * ; ~} \mathrm{p}<0.001={ }^{* * *}$.

649 Extended Data Figure 7. Overview of the PD GWAS genetic signal at the MAPT locus.
A. MAPT primary GWAS signal.
B. MAPT conditional GWAS signal.

## Extended Data Figure 8. ASE sites in MAPT in LD with the H1/H2 SNP.

ASEs derived from putamen and substantia nigra that are in LD with the $\mathrm{H} 1 / \mathrm{H} 2$ tagging SNP, rs12185268 and their position along the MAPT gene. The missense variants track displays the variants annotated as missense by gnomAD v2.1.1 ${ }^{46}$. The valid track displays the heterozygous sites (orange = missense) with an average read depth greater than 15 reads across all samples, in LD with $\mathrm{H} 1 / \mathrm{H} 2$, which were examined for ASE. The topmost track displays the -log10 scale for the minimum FDR across samples for the sites that show an ASE in at least one sample.

A

| Gene | Colb/G | WPPINA | GWAS |
| :---: | :---: | :---: | :---: |
| CAB39L | X |  |  |
| CCNT2 |  |  | X |
| CD38 | X |  | X |
| CTSB | X | X | X |
| DDRGK1 |  |  | X |
| DGKQ |  |  | X |
| GALC | X |  | X |
| GBA |  | X | X |
| GPNMB | X |  | X |
| HSD3B7 | X |  |  |
| INPP5F |  | X |  |
| KAT8 | X | X | X |
| KLHL7 | X |  | X |
| LRRK2 |  | X | X |
| LSM7 | X |  | X |
| MAPT |  | X | X |
| NCKIPSD | X | X | X |
| NSF |  | X |  |
| NUCKS1 | X |  | X |
| NUPL2 | X |  | X |
| PDLIM2 | X |  | X |
| PM20D1 | X |  |  |
| RAB7L1 | X | X |  |
| SH3GL2 |  | X | X |
| SLC41A1 | X |  | X |
| SNCA |  |  | X |
| SPPL2B | X |  |  |
| STK39 |  |  | X |
| VAMP4 | X |  |  |
| WDR6 | X |  |  |
| ZNF646 |  |  | X |

B

| Prediction Technique | Genes |
| :---: | :---: |
| CoIB/G; GWAS; WPPINA | CTSB |
|  | KAT8 |
|  | NCKIPSD |
|  | CD38 |
|  | GALC |
|  | GPNMB |
| CoIB/G; GWAS | KLHL7 |
|  | LSM7 |
|  | $N U C K S 1$ |
|  | NUPL2 |
|  | $P D L I M 2$ |
| CoIB/G; WPPINA | SLC41A1 |
|  | $R A B 7 L 1$ |
| GWAS; WPPINA | GBA |
|  | LRRK2 |
|  | $M A P T$ |
|  | $S H 3 G L 2$ |

C


Figure 1 - Selection of the genes for the PINK1-dependent mitophagy screening.

A


B

Figure 2 - High content mitophagy screen identifies KAT8 as a modulator of pUb(Ser65) levels.


Figure 3 - KAT8 knockdown decreases pUb(Ser65).

A


Figure 4 - Knockdown of the mitochondrial components of the NSL complex reduces pUb(Ser65) levels.


Figure 5 - KANSL1 knockdown decreases pUb(Ser65).


B


Figure 6 - KANSL1 and KAT8 knockdown decrease pUb(Ser65).

A


B


Figure 7 - KANSL1 and KAT8 knockdown decrease mitochondrial clearance.


Figure 8 - ASE sites in KANSL1 in LD with the H1/H2 SNP.


Figure 9 - High content mitophagy screening of the ORFs on the the 17q21 locus identifies only KANSL1 as a modulator of pUb(Ser65) levels.


Extended Data Figure 1. High Content siRNA Screen for modulators of pUb(Ser65).


Extended Data Figure 2. Image processing workflow of the high content screen for O/A induced pUb(Ser65).



Extended Data Figure 3. KAT8 knockdown has no effect on cell viability.


Extended Data Figure 4. KAT8 eQTLs colocalise with SNPs associated with PD risk.


Extended Data Figure 5. KAT8 knockdown decreases pUb(Ser65) levels.

A


10 days


C


$$
\text { - v. - } \mathrm{P}<0.0001
$$

- v. - $P<0.0001$

E


B


$D$


- v. - $\mathrm{P}<0.0001$

F pan-neuronal driver

DA neuron driver




A


B


Extended Data Figure 7. Overview of the PD GWAS genetic signal at the MAPT locus.


Extended Data Figure 8. ASE sites in MAPT in LD with the H1/H2 SNP.

| Gene | ColB | ColG | PPI | GWAS | MPD | MLS |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| ATP13A2 |  |  |  |  | X | X |
| CCNT2 |  |  |  | X |  |  |
| CD38 | X | X |  | X |  |  |
| CTSB | X | X | X |  |  |  |
| DDRGK1 |  |  |  | X |  |  |
| DGKQ |  |  |  | X |  |  |
| DJ1 |  |  |  |  | X |  |
| DNAJC13 |  |  |  |  | X |  |
| FBXO7 |  |  |  |  | X |  |
| GALC | X |  |  | X |  | X |
| GBA |  |  | X |  | X | X |
| GPNMB | X | X |  | X |  |  |
| HSD3B7 |  | X |  |  |  |  |
| IDUA |  |  |  |  |  | X |
| INPP5F |  |  | X |  |  |  |
| KAT8 |  | X | X | X |  |  |
| KLHL7 |  | X |  | X |  |  |
| LRRK2 |  |  | X | X | X |  |
| LSM7 | X | X |  | X |  |  |
| MAPT |  |  | X | X |  |  |
| NCKIPSD | X | X | X | X |  |  |
| NEK1 | X |  |  |  |  |  |
| NSF |  |  | X |  |  |  |
| NUCKS1 |  | X |  | X |  |  |
| NUPL2 | X | X |  | X |  |  |
| PDLIM2 |  | X |  | X |  |  |
| PM20D1 | X |  |  |  |  |  |
| PRKN |  |  |  |  | X |  |
| RAB7L1 | X | X | X |  |  |  |
| SH3GL2 |  |  | X |  |  |  |
| SLC41A1 |  | X |  | X |  |  |
| SNCA |  |  |  |  | X |  |
| SPPL2B | X |  |  |  |  |  |
| STK39 |  |  |  | X |  |  |
| VAMP4 | X | X |  |  |  |  |
| VPS35 |  |  |  |  | X |  |
| WDR6 | X | X |  |  |  |  |
| ZNF646 |  |  |  | X |  |  |

Supplementary Table 1. Complete list of the 38 genes screened in the high content screen. ColB = coloc analysis using Braineac, ColG = coloc analysis using GTEx, WPPINA $=$ weighted protein-protein interaction network; GWAS = genes prioritised in PD-GWAS
(Chang et al, 2017), MPD = Mendelian genes associated with PD, MLS = Mendelian genes associated with lysosomal storage disorders

| eQTL_ dataset | GTEx |
| :---: | :---: |
| sequencing_ method | RNA-seq |
| gene_symbol | KAT8 |
| tissue | Brain_Caudate_ basal_ganglia |
| braineac probe_id | NA |
| nsnps | 2499 |
| PPH0 | 4.89E-10 |
| PPH1 | $1.76 \mathrm{E}-07$ |
| PPH2 | $6.76 \mathrm{E}-04$ |
| PPH3 | 0.242293929 |
| PPH4 | 0.75702972 |
| PD_top_snp | chr16:31000809 |
| $\begin{aligned} & \text { coloc_top_ } \\ & \text { snp } \end{aligned}$ | chr16:31048079 |
| $\begin{aligned} & \text { coloc_SNP_ } \\ & \text { PPH4 } \end{aligned}$ | 0.08637809 |
| coloc_eQTL_ effect_allele | T |
| coloc_eQTL_ other_allele | C |
| $\begin{aligned} & \text { coloc_eQTL_ } \\ & \text { beta } \end{aligned}$ | -0.358743 |
| $\begin{aligned} & \text { coloc_eQTL_ } \\ & \text { SE } \end{aligned}$ | 0.0563187 |
| $\begin{aligned} & \text { coloc_eQTL_ } \\ & \text { Freq1 } \end{aligned}$ | NA |
| $\begin{aligned} & \text { coloc_eQTL_ } \\ & \text { p_val } \end{aligned}$ | 3.49E-09 |
| coloc_PD_Al1 | C |
| coloc_PD_Al2 | T |
| $\begin{aligned} & \text { coloc_PD_ } \\ & \text { beta } \end{aligned}$ | -0.0739 |
| $\begin{aligned} & \text { coloc_PD_ } \\ & \text { SE } \end{aligned}$ | 0.0115 |
| $\begin{aligned} & \text { coloc_PD_ } \\ & \text { Freq1 } \end{aligned}$ | 0.5972 |
| $\begin{aligned} & \text { coloc_PD_ } \\ & \text { p_val } \end{aligned}$ | $1.38 \mathrm{E}-10$ |

Supplementary Table 2. Results of the
Colocalization analysis for KAT8
PD_top_snp = lead SNP in the PD GWAS, coloc_top snp = Most likely SNP responsible for the colocalization signal, coloc_SNP_PPH4 = posterior probability of coloc top SNP being the true SNP responsible for the colocalization signal.


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| 0 |
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\omega\end{array}$
$\omega$
$\stackrel{+}{\sim} \underset{\sim}{\stackrel{\rightharpoonup}{\omega}} \underset{\sim}{\stackrel{\rightharpoonup}{\omega}}$
$\begin{array}{ll} & \overline{0} \\ & 0 \\ \infty & 0 \\ \infty \\ \sim & 0 \\ N & 0 \\ & 0 \\ & 0\end{array}$
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$\underset{\infty}{\text { 징 }}$
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$\begin{aligned} & \stackrel{\circ}{\stackrel{ }{\infty}} \\ & \stackrel{+}{\underset{~}{~}} \\ & \end{aligned}$
$\rightarrow \mathrm{N}$
$\stackrel{o}{\stackrel{\circ}{\circ}}$
$\stackrel{+}{\stackrel{\rightharpoonup}{N}}$
$\begin{aligned} & \circ \\ & \stackrel{\rightharpoonup}{N} \\ & \text { © } \\ & \text { N } \\ & \text { N }\end{aligned}$
$\begin{array}{ll} & 0 \\ \infty & 0 \\ \infty & 0 \\ \sim & 0 \\ & 0 \\ & 0 \\ & 0\end{array}$
$\xrightarrow[N]{\stackrel{\omega}{N}}$
$\stackrel{\rightharpoonup}{\text { a }}$
 KAT8
16
31127075
31142714
0.156
rs9938550
-6.82
rs12597511
0.19827
-4.61
6.178
240
1
lasso
0.17222
$2.56 \mathrm{E}-05$
-6.178
$6.49 \mathrm{E}-10$
$6.79 \mathrm{E}-08$
GTEx．Brain
Frontal＿Cortex
$-B A 9$

KAT8
16
31127075
31142714
0.207
rs9938550
-6.82
rs4527034
0.00998
-4.23
6.1724
240
9
lasso
0.04964
0.0139
-6.04195
$1.52 E-09$
$2.36 E-07$
GTEx．Brain
Cerebellum

| KAT <br> name | Alternative <br> name(s) |
| :--- | :--- |
| KAT1 | HAT1 |
| KAT2A | GCN5 |
| KAT2B | PCAF |
| KAT3A | CREBBP, CBP |
| KAT3B | EP300 |
| KAT4 | TAF1, TFII250 |
| KAT5 | TIP60 |
| KAT6A | MOZ, MYST3 |
| KAT6B | MORF, MYST4 |
| KAT7 | HBO1, MYST2 |
| KAT8 | MOF, MYST1 |
| KAT9 | ELP3 |
| KAT12 | GTF3C4, TFIIIC90 |
| KAT13A | NCOA1, SRC1 |
| KAT13B | NCOA3, SRC3, ACTR |
| KAT13C | NCOA2 |
| KAT13D | CLOCK |
|  | ACAT1 |
|  | ATAT1 |
|  | ATF2 |
|  | BLOC1S1 (GCN5L1) |
|  | NAT10 |

Supplementary Table 4. Complete list of the 22 KATs screened in the high content screen. (See Fig 3E)

Within each row, compare columns (simple effects within rows)

| Number of families | 8 |
| :--- | :--- |
| Number of comparisons per family | 3 |


| Dunnett's multiple comparisons test | Mean Diff. | 95.00\% CI of diff. | Significant? | Summary | Adjusted P Value |
| :---: | :---: | :---: | :---: | :---: | :---: |
| DMSO |  |  |  |  |  |
| SCR vs. KAT8 | 0 | -14.19 to 14.19 | No | ns | >0.9999 |
| SCR vs. KANSL1 | 0 | -14.19 to 14.19 | No | ns | >0.9999 |
| SCR vs. PINK1 | 0 | -14.19 to 14.19 | No | ns | >0.9999 |
| 1 h |  |  |  |  |  |
| SCR vs. KAT8 | 21 | 6.807 to 35.19 | Yes | ** | 0.0017 |
| SCR vs. KANSL1 | 30 | 15.81 to 44.19 | Yes | **** | <0.0001 |
| SCR vs. PINK1 | 38 | 23.81 to 52.19 | Yes | **** | <0.0001 |
| 2 h |  |  |  |  |  |
| SCR vs. KAT8 | 30 | 15.81 to 44.19 | Yes | **** | <0.0001 |
| SCR vs. KANSL1 | 53 | 38.81 to 67.19 | Yes | **** | <0.0001 |
| SCR vs. PINK1 | 71 | 56.81 to 85.19 | Yes | **** | <0.0001 |
| 3 h |  |  |  |  |  |
| SCR vs. KAT8 | 33 | 18.81 to 47.19 | Yes | **** | <0.0001 |
| SCR vs. KANSL1 | 51 | 36.81 to 65.19 | Yes | **** | <0.0001 |
| SCR vs. PINK1 | 73 | 58.81 to 87.19 | Yes | **** | <0.0001 |
| 4 h |  |  |  |  |  |
| SCR vs. KAT8 | 15 | 0.8065 to 29.19 | Yes | * | 0.0355 |
| SCR vs. KANSL1 | 28 | 13.81 to 42.19 | Yes | **** | <0.0001 |
| SCR vs. PINK1 | 49 | 34.81 to 63.19 | Yes | **** | <0.0001 |
| 5 h |  |  |  |  |  |
| SCR vs. KAT8 | 9 | -5.193 to 23.19 | No | ns | 0.307 |
| SCR vs. KANSL1 | 21 | 6.807 to 35.19 | Yes | ** | 0.0017 |
| SCR vs. PINK1 | 39 | 24.81 to 53.19 | Yes | **** | <0.0001 |
| 6 h |  |  |  |  |  |
| SCR vs. KAT8 | 14 | -0.1935 to 28.19 | No | ns | 0.0542 |
| SCR vs. KANSL1 | 24 | 9.807 to 38.19 | Yes | *** | 0.0003 |
| SCR vs. PINK1 | 42 | 27.81 to 56.19 | Yes | **** | <0.0001 |
| 7 h |  |  |  |  |  |
| SCR vs. KAT8 | 19 | 4.807 to 33.19 | Yes | ** | 0.0051 |
| SCR vs. KANSL1 | 23 | 8.807 to 37.19 | Yes | *** | 0.0005 |
| SCR vs. PINK1 | 40 | 25.81 to 54.19 | Yes | **** | <0.0001 |

Supplementary Table 5. p-values for Figure 6B.

| hetSNP | Sample ID | Read count | Allele | Ens.Alt | Individual ID | Tissue |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 17:44108355 | A653_043 | 47 | A | A | 004_06 | SNIG |
| 17:44108355 | A653_043 | 20 | G | A | 004_06 | SNIG |
| 17:44108355 | A653_441 | 69 | A | A | 032_09 | PUTM |
| 17:44108355 | A653_441 | 0 | G | A | 032_09 | PUTM |
| 17:44159849 | A653_031 | 2 | C | C | 024_09 | PUTM |
| 17:44159849 | A653_031 | 23 | T | C | 024_09 | PUTM |
| 17:44159849 | A653_719 | 58 | C | C | 035_09 | SNIG |
| 17:44159849 | A653_719 | 26 | T | C | 035_09 | SNIG |
| 17:44248769 | A653_093 | 36 | C | C | 004_08 | SNIG |
| 17:44248769 | A653_093 | 4 | T | C | 004_08 | SNIG |
| 17:44248769 | A653_184 | 30 | C | C | 013_09 | PUTM |
| 17:44248769 | A653_184 | 7 | T | C | 013_09 | PUTM |
| 17:44248769 | A653_326 | 26 | C | C | 017_09 | PUTM |
| 17:44248769 | A653_326 | 3 | T | C | 017_09 | PUTM |
| 17:44248769 | A653_617 | 22 | C | C | 029_09 | SNIG |
| 17:44248769 | A653_617 | 0 | T | C | 029_09 | SNIG |
| 17:44248769 | A653_679 | 23 | C | C | 030_06 | PUTM |
| 17:44248769 | A653_679 | 3 | T | C | 030_06 | PUTM |
| 17:44248769 | A653_753 | 43 | C | C | 029_09 | PUTM |
| 17:44248769 | A653_753 | 10 | T | C | 029_09 | PUTM |
| 17:44248769 | A653_794 | 24 | C | C | 036_09 | SNIG |
| 17:44248769 | A653_794 | 4 | T | C | 036_09 | SNIG |
| 17:44248814 | A653_043 | 10 | A | A | 004_06 | SNIG |
| 17:44248814 | A653_043 | 0 | G | A | 004_06 | SNIG |
| 17:44248814 | A653_326 | 19 | A | A | 017_09 | PUTM |
| 17:44248814 | A653_326 | 3 | G | A | 017_09 | PUTM |
| 17:44248814 | A653_617 | 12 | A | A | 029_09 | SNIG |
| 17:44248814 | A653_617 | 0 | G | A | 029_09 | SNIG |
| 17:44248814 | A653_679 | 19 | A | A | 030_06 | PUTM |
| 17:44248814 | A653_679 | 2 | G | A | 030_06 | PUTM |
| 17:44248814 | A653_753 | 25 | A | A | 029_09 | PUTM |
| 17:44248814 | A653_753 | 5 | G | A | 029_09 | PUTM |
| 17:44248814 | A653_950 | 28 | A | A | 006_10 | SNIG |
| 17:44248814 | A653_950 | 8 | G | A | 006_10 | SNIG |

Supplementary Table 6. Summary of the results of ASE analysis across the KANSL1 gene.

## Column name

hetSNP
Avg.Reads.all.samples
min.FDR
Allele1
Allele2
ens.alt
symbol
most.severe.consequence ASE

## Description

Heterozygous SNP with average read depth >15 across samples
Average read depth across all samples
Minimum false discovery rate across the samples
Allele 1 for the hetSNP
Allele 2 for the hetSNP
Ensembl alternate allele
Gene name obtained using the variant effect predictor tool (VEP)
Most severe of all observed consequence types reported for the hetSNP (VE Indicates ASE = ' $Y$ ' if $\min . F D R<0.05$, ' $N$ ' if min.FDR $>=0.05$

The following column details were from obtained from LDProxy (https://Idlink.nci.nih.gov/)
RSID Identifier for the hetSNP

Alleles
MAF
Dprime
R2

Identifier for the hetSNP
SNP alleles
Minor allele frequency
Indicator of allelic segregation for two genetic variants
Measure of correlation of alleles for two genetic variants.



| $\mathbf{1 7 q 2 1}$ ORFs |  |
| :--- | :--- |
| ACBD4 | HEXIM1 |
| ADAM11 | HIGD1B |
| ARHGAP27 | KANSL1 |
| ARL17A | KIF18B |
| ARL17B | LRRC37A |
| C1QL1 | LRRC37A2 |
| CCDC103 | MAP3K14 |
| CDC27 | MAPT |
| CRHR1 | MYL4 |
| DBF4B | NMT1 |
| DCAKD | NSF |
| EFTUD2 | PLEKHM1 |
| FMNL1 | RPRML |
| GFAP | SPPL2C |
| GJC1 | STH |
| GOSR2 | WNT3 |

## Supplementary Table 8. Complete list of the

 32 ORFs in the 17q21 locus screened in the high content screen. (See Fig 9)