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SHORT REPORTS

An Al-guided screen identifies probucol as an enhancer of mitophagy through modulation of lipid droplets

Natalia Moskal¹, Naomi P. Visanji², Olena Gorbenko¹, Vijay Narasimhan³, Hannah Tyrrell¹, Jess Nash¹, Peter N. Lewis¹, G. Angus McQuibban^{1*}

1 Department of Biochemistry, University of Toronto, Toronto, Canada, 2 Edmund J Safra Program in Parkinson's Disease and Morton and Gloria Shulman Movement Disorders Centre, Toronto Western Hospital, Toronto, Canada, 3 Zebrafish Centre for Advanced Drug Discovery and Keenan Research Centre for Biomedical Science, Li Ka Shing Knowledge Institute, St. Michael's Hospital and Department of Medicine and Physiology, University of Toronto, Toronto, Canada

* angus.mcquibban@utoronto.ca

Abstract

Failures in mitophagy, a process by which damaged mitochondria are cleared, results in neurodegeneration, while enhancing mitophagy promotes the survival of dopaminergic neurons. Using an artificial intelligence platform, we employed a natural language processing approach to evaluate the semantic similarity of candidate molecules to a set of well-established mitophagy enhancers. Top candidates were screened in a cell-based mitochondrial clearance assay. Probucol, a lipid-lowering drug, was validated across several orthogonal mitophagy assays. In vivo, probucol improved survival, locomotor function, and dopaminergic neuron loss in zebrafish and fly models of mitochondrial damage. Probucol functioned independently of PINK1/Parkin, but its effects on mitophagy and in vivo depended on ABCA1, which negatively regulated mitophagy following mitochondrial damage. Autophagosome and lysosomal markers were elevated by probucol treatment in addition to increased contact between lipid droplets (LDs) and mitochondria. Conversely, LD expansion, which occurs following mitochondrial damage, was suppressed by probucol and probucol-mediated mitophagy enhancement required LDs. Probucol-mediated LD dynamics changes may prime the cell for a more efficient mitophagic response to mitochondrial damage.

Introduction

Clearance of damaged mitochondria is important for the survival of dopaminergic neurons, the loss of which is responsible for the classical motor symptoms of Parkinson disease (PD). PD is a progressive neurodegenerative disease characterized by rigidity, akinesia, and bradykinesia as well as wide ranging nonmotor symptoms such as anxiety, depression, sleep disturbances, and loss of smell [1]. Current treatment options address symptoms but fail to disrupt the progression of the disease, thus a disease-modifying treatment remains a major unmet need. Mitochondrial dysfunction is clearly implicated in PD pathogenesis, so enhancing the

Abbreviations: ABCA1, ATP binding cassette transporter A1; AI, artificial intelligence; AUC, area under the curve; CCCP, carbonyl cyanide mchlorophenyl hydrazone; DGAT, diacylglycerol acyltransferase; LD, lipid droplet; MPP⁺, 1-methyl-4-phenylpyridinium; mtDNA, mitochondrial DNA; PARP, poly ADP-ribose polymerase; PD, Parkinson disease; PINK1, PTEN-induced putative kinase 1; ROC, receiver operating characteristic; ROCK2, Rho-associated protein kinase 2; VDAC1, voltagedependent anion-selective channel protein 1; vDC, ventral diencephalon. removal of damaged mitochondria (mitophagy) might have potential as a therapeutic intervention [1].

Evidence for the importance of mitophagy in PD pathogenesis comes from both sporadic and genetic cases. Several disease-causing, loss-of-function mutations in genes encoding proteins that mediate mitophagy have been identified, including PINK1 and PRKN [2,3]. Additionally, several disease-causing mutations in genes not directly associated with mitophagy have secondary negative effects on mitochondrial health or on the retrograde transport of damaged mitochondria to axons, such as SNCA, GBA, and LRRK2 [4–6]. Besides genetic causes of PD, mitochondrial damage and mitophagy impairment have also been widely implicated in sporadic disease, including the inactivation of proteins that mediate mitophagy [7–9]. The clear connection between this pathway and the health of dopaminergic neurons emboldened us to search for novel mitophagy-enhancing compounds as potential disease-modifying therapeutics for PD.

A review by Georgopoulos and colleagues describes several compounds currently known to stimulate and potentiate mitophagy [10]. However, most of these compounds also induce mitochondrial damage or apoptosis. While these are bona fide mitophagy enhancers, induction of mitochondrial damage or apoptosis would likely preclude their clinical use as this may worsen PD pathogenesis. To address the unmet need for mitophagy enhancers with therapeutic potential, we employed a computational approach using artificial intelligence (AI) to identify previously uncharacterized mitophagy enhancers. We have previously successfully used this strategy that detects patterns and associations across several large datasets to identify drugs with disease modifying potential for PD that target aggregation of alpha synuclein [11–13].

Here, using a similar approach, we screened a candidate list of molecules from the Drug-Bank database for similarity to known mitophagy enhancers. Many of the drugs have already been safely administered to humans. Repurposing drugs from other indications offers the opportunity to accelerate the clinical trials pipeline, given the presence of preexisting pharmacological and toxicological information about candidate compounds [14].

In a previous screen, we focused on compounds that accelerated the transition of the E3 ubiquitin ligase, Parkin, from the cytosol to the mitochondria-a step, which is integral to this mitophagy pathway [15]. While this step is highly amenable to microscopy-based phenotypic screening, the limitations of this approach include possible omission of hits that function downstream of Parkin or that target other mitophagy pathways [16]. In this present screening iteration, we evaluated the clearance of damaged mitochondria from cells. Ultimately, if this downstream step is improved, then the negative consequences of mitochondrial damage in the dopaminergic neurons may be mitigated [17]. This effort led to the identification of a compound that enhances mitophagy following mitochondrial damage and led us to elucidate its mechanism of action, resulting in the identification of a putative newfound role for the ATP binding cassette transporter A1 (ABCA1) in mitophagy, through its effects on lipid droplets (LDs) dynamics.

Results

Artificial intelligence (AI) simplifies mitochondrial clearance screen for mitophagy enhancers

We employed a computational approach using AI (IBM Watson for Drug Discovery) to identify drugs amenable to repurposing as PD therapeutics. Specifically, an in silico screen was performed to identify potential mitophagy enhancers from the DrugBank database (www. drugbank.ca), based on their similarity to positive control mitophagy enhancers. Our training set was based on a review article about mitophagy modulators, and it comprised the following 7 drugs: PTEN-induced putative kinase 1 (PINK1) activator kinetin; poly ADP-ribose polymerase (PARP) inhibitor olaparib, p53 inhibitor pifithrin-alpha; nicotinamide (NAD⁺ accumulation); and sirtuin1 activators resveratrol, fisetin, and SRT1720 [10] (S1C Fig). Extensive model validation was performed prior to commencing the screen in cells (S1 Fig; see Methods for details.)

Ultimately, we assessed 3,231 candidate drugs from the DrugBank database for semantic similarity to the training drugs (S1C Fig). IBM Watson for Drug Discovery creates text fingerprints for all the training set molecules, in addition to 3,231 candidate molecules from the DrugBank database. We computed a similarity score (to the training set) score for each candidate entity, and they ranked drugs from highest to lowest accordingly (S1 Dataset).

The top 79 most similar candidates identified by IBM Watson for Drug Discovery were screened in HeLa cells stably expressing GFP Parkin and mito-DsRed and subjected to prolonged treatment with carbonyl cyanide m-chlorophenyl hydrazone (CCCP), which depolarizes the mitochondrial membrane potential. Loss of mito-DsRed signal occurs in mitophagic cells, which comprise a large percentage of the population (S2C and S3B Figs) [18]. In cells pretreated with DMSO instead of small molecules, $78 \pm 2.83\%$ of cells have low/no mito-DsRed signal (S3B Fig). MAD z-scores were calculated, as previously described for each of the 79 compounds (S2A Fig and S3 Dataset) [19]. Three compounds were selected for retesting based on MAD z-score values (S2A Fig), thereby producing a hit rate of 3.8%.

Probucol augments later stages of mitophagy

Upstream of the degradation of damaged mitochondria is the targeting of mitochondria to lysosomes. The mitoQC assay can be used in both cells and in vivo to probe this precise step in mitophagy. Briefly, cells were transfected with Cerulean-Parkin and RG-OMP25, a plasmid containing mCherry and GFP in tandem with the mitochondrial targeting sequence of OMP25 (Fig 1A). Upon localization to acidic lysosomes, GFP signal is quenched, which distinguished mitochondria localized to lysosomes because they appear as puncta with red-only signal. Less than 5 red-only puncta are present in cells under normal conditions, but the number of puncta increases following induction of mitochondrial damage [20]. Cells with greater than 5 red-only puncta corresponding to mito-lysosomes were classified as "mitophagic" (Fig 1A). Following 6-hour CCCP treatment, $40.25 \pm 6.3\%$ cells are mitophagic compared to 13.32 ± 4.83% under normal conditions. A higher percentage of probucol-treated cells were mitophagic ($66.29 \pm 5.6\%$; Fig 1B), compared to treatments with the other candidate compounds.

Compounds were retested in an orthogonal mitochondrial clearance assay; probucol increased the extent to which mitochondrial substrates were degraded following induction of mitochondrial damage. Immunoblotting was performed to visualize levels of the outer mitochondrial membrane protein voltage-dependent anion-selective channel protein 1 (VDAC1) and the inner mitochondrial membrane substrate ATP synthase F1 subunit alpha (ATP5A) (S2D, S2E and S4 Figs). Predictably, VDAC1 levels declined over the CCCP time course. While probucol treatment does not increase VDAC1 degradation under basal conditions, it promoted enhanced degradation of VDAC1 over the CCCP time course (S2D and S2E Fig).

We also probed steps further upstream in the mitophagy cascade, including PINK1-mediated phosphorylation of ubiquitin at S65 and recruitment of Parkin to damaged mitochondria (S5 Fig). Phosphorylation of mitochondrial ubiquitin S65 following mitochondrial damage did not further increase following probucol treatment based on immunostaining and immunoblotting experiments (S5A, S5B and S5D Fig). Parkin recruitment to damaged



Fig 1. Probucol increased the targeting of mitochondria to lysosomes through ABCA1. (A) The mitoQC reporter was expressed in HeLa cells along with Cerulean-Parkin. Mitophagy was stimulated by treating cells with CCCP for 6 hours along with either DMSO or candidate molecules from the screen. Mitochondria appear as red-only puncta when localized to acidic cellular compartments. (B) Cells with >5 red-only puncta were defined as mitophagic, and the mean percentage of mitophagic cells in each treatment condition is displayed. (C) Seven-day old flies expressing the mitoQC reporter in dopaminergic neurons were fed food supplemented with probucol in combination with either paraquat or water. (D) Immunostaining using antibody against TH segmented and defined dopaminergic neurons. The mean percentage of red-only mitochondrial area in each dopaminergic neuron is displayed. (E) Control flies and flies coexpressing the human ABCA1 transgene and the mitoQC reporter were administered probucol in the presence and absence of paraquat. The mean percentage of red-only mitochondrial area in ELA cells stably expressing mitoQC and Cerulean-Parkin, which were transfected with either pLKO1 vector or with shABCA1. The mean percentage of red-only mitochondrial area percentage of red-only mitochondrial area present the either pLKO1 vector or with shABCA1. The mean percentage of red-only mitochondrial area percentage of red-only mitochondrial area present the either pLKO1 vector or with shABCA1. The mean percentage of red-only mitochondrial area percentage of red-only mitochondrial area percentage of red-only mitochondrial area percentage of red-only was assessed in HELA cells stably expressing mitoQC and Cerulean-Parkin, which were transfected with either pLKO1 vector or with shABCA1. The mean percentage of red-only mitochondrial area percentage of red-only mitochondrial area percentage of a least 3 biological replicates, each indicated by data points in B, D, E, and F. Bars represent mean values and error bars represent SEM. *, ***,

cells were assessed for B, D, E, and F, respectively. Dopaminergic neurons from at least 2 fly brains were analyzed for each treatment with at least 10 ROIs per brain. Statistical analysis was performed using ANOVA and Dunnett's multiple comparison correction. The data underlying the graphs shown in the figure can be found in <u>S1 Data</u>. ABCA1, ATP binding cassette transporter A1; CCCP, carbonyl cyanide m-chlorophenyl hydrazone; ROI, region of interest; TH, tyrosine hydroxylase.

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mitochondria likewise was not increased by probucol treatment (<u>S5C Fig</u>), indicating that this mitophagy enhancer likely exerts its effect on steps further downstream of Parkin recruitment or through other Parkin-independent mitophagy pathways [16].

To further substantiate the findings in cells, the mitoQC reporter was expressed in the dopaminergic neurons of flies fed food supplemented with DMSO or probucol alone or in combination with paraquat, a mitochondrial toxin that causes PD in humans and PD-related phenotypes such as loss of dopaminergic neurons and locomotor impairments in model organisms such as flies [21]. Paraquat induces a mitophagy response characterized by an increase in mito-lysosomes in dopaminergic neurons [22]. Mitophagy further increased in flies coadministered probucol along with paraquat (Fig 1C and 1D). Basal mitophagy did not differ in the dopaminergic neurons of flies fed either DMSO- or probucol-supplemented food.

Probucol improves mitochondrial damage-induced phenotypes across several animal models of PD

Drosophila and *Danio rerio* serve as PD model systems, as they replicate much of human PD pathogenesis and display phenotypes, which reflect human disease presentation. Flies and zebrafish exhibit loss of dopaminergic neurons resulting in impaired locomotion following exposure to PD-causing toxins such as 1-methyl-4-phenylpyridinium (MPP⁺) and paraquat [21,23]. After replicating these features in the two model organisms, we tested probucol's disease-modifying potential. The climbing ability and survival of flies declined following paraquat administration, but cotreatment with probucol improved locomotor function and survival (Fig 2A and 2B).

Zebrafish embryos were incubated in MPP⁺ in combination with probucol or DMSO. Following 6 days of incubation, the movement of adult zebrafish was captured using ZebraBox. The distance travelled by the zebrafish exposed to DMSO in combination with MPP⁺ was visibly reduced compared to zebrafish incubated in DMSO alone, but the addition of probucol to the MPP⁺ increased the distance travelled by the zebrafish (Fig 2C and 2D). SR3677, a chemical inhibitor of Rho-associated protein kinase 2 (ROCK2) was employed as a positive control. We previously characterized this compound as a mitophagy enhancer and found that it improved locomotor decline in flies fed paraquat [22].

In addition to toxin-based models of PD, we also tested probucol's effect in a genetic model of mitochondrial dysfunction. Specifically, in heteroplasmic flies with approximately 90% of mitochondrial DNA (mtDNA) that contained a temperature-sensitive mutation in mitochondrial cytochrome c oxidase subunit I (*mt:Coll^{T3001}*). Shifting these flies to a nonpermissive temperature causes mitochondrial dysfunction resulting in climbing defects and significant reduction of life span [22,24,25]. Climbing and life span were improved in heteroplasmic *mt: Coll^{T3001}* flies fed food containing probucol rather than DMSO (S6 Fig). Much like paraquat, the phenotypes displayed by these flies arise from mitochondrial dysfunction, and probucol reduced their severity, possibly by promoting mitophagy in this context as well, given that mitophagy has previously been shown to remove mitochondria bearing deleterious mutations [26].

To directly probe the cell type impacted in PD, Tg(dat:EGFP) zebrafish embryos were coincubated with MPP⁺ and probucol or vehicle control. Dopaminergic neurons can easily be



Fig 2. Probucol alleviated PD-related phenotypes, which arose from mitochondrial dysfunction in vivo. (A) Effect of probucol administration on paraquat-induced climbing defect. The percentage of flies to climb across a height of 12.5 cm. (B) Effect of probucol administration on survival, in the presence and absence of paraquat cotreatments. (C) Single particle tracking traces to visualize distance travelled by zebrafish in wells following administration of DMSO, probucol, and MPP⁺, as indicated. (D) Distance travelled by zebrafish in each treatment group in C, in addition to following treatment with positive control compound SR3677. (E) Tg(*dat:EGFP*) zebrafish brains were imaged following treatment with DMSO, probucol, and MPP⁺. Dashed lines highlight the vDC region of interest. (F) The number of vDC neurons in each treatment group in E was quantified. Data information: At least 3 independent biological replicates were performed for each experiment, with individual replicates depicted by data points in F. Bars and error bars represent mean and SEM values, respectively. Unpaired Student *t* tests were performed to analyze data in A, log-rank tests were used to analyze survival data, and one-way ANOVA analysis was performed to analyze data in D and F along with Dunnett's multiple comparison correction. *, ***, and **** indicate *p*-values <0.05, <0.01, <0.005, and <0.001, respectively. The data underlying the graphs shown in the figure can be found in S1 Data. MPP+, 1-methyl-4-phenylpyridinium; PD, Parkinson disease; vDC, ventral diencephalon.

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visualized and counted in this transgenic model²⁷. Following 1 day of incubation in MPP⁺, the number of dopaminergic neurons in the ventral diencephalon (vDC), which is analogous to the human nigrostriatal region, was reduced. In contrast, probucol-fed zebrafish retained more of their dopaminergic vDC neurons (Fig 2E and 2F). This reduced loss of dopaminergic neurons likely led to the probucol-mediated improvements to PD-related phenotypes.

ABCA transporter and its effects on lipid droplets mediate probucol's mitophagy enhancement

To determine whether probucol's canonical target, ABCA1, is involved in its effects on mitophagy, we performed several genetic manipulations to change ABCA1 levels in cell culture or in the dopaminergic neurons of flies. The same mitophagy assay used to characterize probucol's effects was also performed in this context.

Human ABCA1 and mitoQC transgenes were coexpressed, or mitoQC was expressed alone in the dopaminergic neurons of flies (Figs 1E, S7A and S7D). Once again, paraquat treatment increased the percentage of red-only mitochondrial puncta, which represent mitochondria localized to lysosomes. Basal mitophagy was unaffected by overexpression of the transgene. However, in flies fed paraquat, ABCA1 reduced the percentage of mitochondrial area localized to lysosomes (Fig 1E). Immunoblotting was performed to confirm expression of human ABCA1 (S7A Fig).

We also reduced ABCA1 levels with shRNA (S7B Fig) in cells expressing mitoQC and Cerulean-Parkin and treated cells as in Fig 1A. While probucol increased the percentage of redonly mitochondrial area indicative of mito-lysosomes in cells transfected with the control pLKO1 vector, probucol no longer increased mitophagy in cells transfected with shABCA1 (Figs 1F and S7C).

The extent of crosstalk between lipid homeostasis and autophagy reported in other studies and ABCA1's role in lipid efflux led us to test whether probucol altered LDs. LDs have been found to form in response to starvation-induced autophagy and deferiprone-induced mitophagy [28,29]. LDs similarly increased significantly in cells treated with CCCP for 24 hours (Figs 3C and S8). Surprisingly, probucol treatment reduced the LD area expansion that occurred following mitophagy induction but had no effect on LD area under basal conditions (Figs 3C and S8). LD area also increased in the dopaminergic neurons when flies consumed paraquat in their food. Probucol supplementation decreased the LD area expansion to levels comparable to those under basal conditions (Fig 3A and 3B).

LD-mitochondria contacts have been observed in several studies. Peridroplet mitochondria localized to contact sites exhibit altered bioenergetic properties and serve as a source of ATP for LD expansion [30]. However, under conditions of high autophagic flux, LDs are thought to interact with mitochondria to buffer the lipotoxic species that are produced as byproducts of autophagy [28]. Interestingly, a greater proportion of LDs colocalized with mitochondria in probucol-treated cells under basal conditions where mitochondria appeared intact and visibly elongated (Figs 3D and S8). Under the conditions in which this increased contact is observed, mitophagic flux is not elevated (Fig 1C and 1D), so it is unlikely that the contacts facilitate buffering of lipotoxic byproducts like in starvation-induced autophagy [28].

Finally, to evaluate the importance of LDs on the mitophagy enhancement conferred by probucol treatment, diacylglycerol acyltransferases (DGAT) 1 and DGAT2 inhibitor treatment was added to suppress the LD expansion, which occurred following prolonged mitochondrial stress (Figs <u>3C</u> and <u>S8</u>). The addition of DGAT inhibitors to probucol treatment in the context of paraquat-induced mitochondrial dysfunction in flies attenuated probucol's enhancement of mitophagy in the dopaminergic neurons of flies (Fig <u>3E</u>). Mitophagy was no longer elevated by



Fig 3. ABCA-mediated effects of probucol on mitophagy depended on LDs, which increased proximity to mitochondria upon probucol treatment. (A) TH-positive dopaminergic neurons were segmented in BODIPY-stained brains from flies fed food supplemented with the indicated combinations of probucol and paraquat. (B) The percentage of LD area over the total TH-positive cell area is quantified from A. (C) The percentage of LD area over total cell area in HeLa cells treated with either DMSO or probucol alone or in the presence of CCCP with and without DGAT inhibitors. (D) The overlap between BODIPY-stained LDs and mitochondria was assessed using the Manders Correlation Coefficient in HeLa cells treated as described in C). (E) Probucol treatment was coadministered along with paraquat in the presence and absence of DGAT inhibitors. The percentage of red-only mitochondrial area was measured. (F) HeLa cells were transfected with pLKO1 and shABCA1 plasmids and treated with the indicated combination of probucol and CCCP for 24 hours. BODIPY staining was used to quantify the percentage of cell area occupied by LDs. Data information: Results are represent the SEM. * and *** represent *p*-values <0.05 and 0.005, respectively. Statistical analysis was performed using ANOVA analysis, with multiple comparison correction for E and F. Unpaired Student *t* test analysis was performed to compare DMSO and probucol groups in B, C, and D. The data underlying the graphs shown in the figure can be found in S1 Data. CCCP, carbonyl cyanide m-chlorophenyl hydrazone; DGAT, diacylglycerol acyltransferase; LD, lipid droplet; TH, tyrosine hydroxylase.

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probucol in the context of paraquat, when DGAT inhibitors are present. Since DGAT inhibitors reduced LD area and attenuated probucol's effect on mitophagy, this suggests probucol's effect on LDs may be responsible for its subsequent effects on mitophagy.

Since probucol no longer exerted effects on mitophagy without LDs and when ABCA1 levels are reduced, we probed whether ABCA1 affected probucol-mediated LD expansion. This critical step in probucol's mechanism of action on mitophagy was no longer evident when ABCA1 levels were reduced with shRNA (Figs <u>3F</u> and <u>S7B</u>).

Probucol increases LC3 lipidation and lysosome abundance

Since early mitophagy steps were unaltered by probucol, downstream steps were assessed next. Immunoblotting can be used to evaluate the lipidation status of LC3. The lower molecular weight, lipidated form of LC3 (LC3-II), correlates with autophagosome levels and therefore increases as autophagy proceeds. However, it is important to note that increased LC3-II levels alone, in the absence of cotreatments with lysosome inhibitors, cannot directly indicate increased autophagic flux [31]. LC3-II levels increased following probucol treatment under basal conditions (S9 Fig) in HEK293 cells with endogenous Parkin present at low levels. LC3-II levels were predictably higher upon mitophagy induction with CCCP treatment compared to basal conditions, as other studies show. However, the difference in LC3-II between DMSO- and probucol-treated cells did not persist under these conditions.

A recent report found that autophagy must be tuned to provide sufficient dynamic range to resolve differences in LC3 lipidation following manipulations such as drug treatments [32]. This can be accomplished by employing bafilomycin, an inhibitor of lysosome-autophago-some fusion at low doses so the effects of manipulations would be apparent. Experiments were repeated in HeLa cells, which lack endogenous Parkin with the addition of low-dose bafilomycin treatment. While the difference between DMSO- and probucol-treated groups under basal conditions was not as robust as in HEK293 cells, the addition of bafilomycin revealed a difference in LC3 lipidation between DMSO- and probucol-treated cells in the CCCP group (Fig 4A and 4B).

An endogenously mCherry-tagged *Lamp* reporter fly line was employed to assess lysosomes in vivo. Lamp is a lysosomal protein, which increases in abundance following paraquat treatment [33]. Dopaminergic neurons were segmented with TH immunostaining, and the area occupied by Lamp-positive lysosomes within dopaminergic neurons was measured. As expected, paraquat addition to fly food increased abundance of Lamp-positive puncta (Fig 4C). Probucol treatment increased the area of lysosomes in the dopaminergic neurons, under basal conditions where no exogenous stimulus was added to trigger lysosome accumulation (Fig 4C and 4D).

To determine whether probucol's effect on mitophagy were responsible for its ability to improve paraquat-induced climbing impairment, climbing assays were performed as described in Fig 2A, but the dopaminergic neuron-specific TH-GAL4 driver was used to drive expression of RNAi targeting candidate genes. In this manner, we dissected which factors were dispensable for probucol-mediated climbing improvements. Consistently with experiments in cells that showed that probucol had no effect on Parkin subcellular distribution (S5 Fig), *parkin* RNAi did not abrogate the effects of probucol-mediated improvement of climbing impairment, which was evident in both the mCherry^{RNAi} and *parkin*^{RNAi}-expressing flies (Fig 4E).

However, probucol treatment no longer improved climbing when *ABCA* RNAi was expressed in dopaminergic neurons (Fig 4E). These findings show that probucol treatment increased downstream autophagy steps such as autophagosome and lysosome biogenesis, seemingly priming the cells for a more rapid and efficient mitophagy response when damage



Fig 4. LC3 lipidation and lysosome area increased following probucol treatment and ABCA was necessary for probucolmediated climbing improvements. (**A**) HeLa cells were either incubated in DMEM, DMEM with CCCP, or HBSS media for 6 hours in the presence or absence of bafilomycin. Lysates were separated by SDS-PAGE and immunoblotting was performed using antibodies that recognize LC3 and tubulin, as a loading control. (**B**) Densitometry was performed to measure the levels of lipidated LC3-II, which were normalized to the tubulin loading control. (**C**) Endogenously tagged *Lamp*-3xmCherry flies were fed food supplemented with DMSO or probucol in the presence and absence of paraquat. (**D**) The Lamp area in each cell was measured as a percentage of total cell area, as defined by segmentation of TH-positive dopaminergic neurons. (**E**) RNAi targeting either *parkin* or *ABCA* was expressed in the dopaminergic neurons with the TH-Gal4 driver. The effect of probucol on paraquat-induced climbing impairment was assessed in flies from the various genotypes. Climbing as a percentage of flies to cross 12.5 cm height is displayed. (**F**) Hypothetical mechanism of mitophagy enhancement by probucol may involve up-regulation of LDs adjacent to mitochondria. LD expansion occurs following mitochondrial damage, but in the presence of probucol, the abundance of LDs is reduced, resulting in increased abundance of mito-lysosomes. Data information: Bars represent mean values and error bars correspond to SEM from at least 3 independent biological replicates, which are indicated with data points in graphs B and D. Statistical analysis to assess

differences between DMSO and probucol within the same treatment condition or genotype group was performed using one-way ANOVA tests with multiple comparison correction in B and E, while unpaired *t* tests were used to compare DMSO and probucol in D. *, *, and *** indicate p-value <0.05, <0.01, and <0.005, respectively. The data underlying the graphs shown in the figure can be found in S1 Data. CCCP, carbonyl cyanide m-chlorophenyl hydrazone; LD, lipid droplet; TH, tyrosine hydroxylase.

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strikes. ABCA, probucol's target, likely facilitates this effect through alterations to LD dynamics (Fig 4F).

Discussion

Using our screening approach, several previously characterized mitophagy enhancers were recovered both in silico and in cells. In silico screening identified staurosporine among the top hits (17/3,231, top 0.5%), which is a well-characterized activator of mitophagy, but it was excluded from further experiments as it induces apoptosis [34]. In our cell-based screen, dichlorocopper ranked as 1/79 in our screen. Copper binds to and increases the kinase activity of autophagy regulatory kinases ULK1 and ULK2 [35,36]. ULK1/2 mediate autophagosome formation downstream of PINK1/Parkin [37,38]. Dichlorocopper would not have been identified had our mitophagy screen focused on Parkin recruitment, but nevertheless enhanced mitophagy, so the design of this screen may be superior to our previous screening approach [22]. The recovery of compounds with established mitophagy-promoting effects gave us confidence in the predictive power and efficacy of our dual screen. By filtering out compounds that induce mitochondrial damage or apoptosis, our effort was focused on identifying new compounds and mechanisms leading to mitophagy enhancement.

Interestingly, the set of compounds used to train our in silico model to identify mitophagy enhancers largely consisted of SIRT1 agonists (S1C Fig). SIRT1 affects mitophagy by up-regulating the mitophagy receptor BNIP3 in aged mouse kidney [39]. Several compounds function by increasing the cellular NAD⁺ pool, which is a cofactor for SIRT1. This bias led us to speculate that our dual screen would identify several more mitophagy enhancers that function through this common mechanism shared among the training set. Despite the bias, only 1 of the 3 final hits (3-methoxybenzamide) was a SIRT1 agonist. Interestingly, a recent Phase I clinical trial has demonstrated efficacy for nicotinamide, an NAD⁺ precursor, in PD, so the mechanism of action encompassed in the training set is likely nevertheless a relevant disease target [40].

Ultimately, the screen identified probucol, a drug used to treat hypercholesterolemia prior to the advent of statins. The target, which probucol inhibits, is the ATP-binding cassette transporter ABCA1, and diverse assays in cells and flies, which involved genetically reducing ABCA levels, abolished probucol's effects on mitophagy in cells and climbing in flies, while expressing human ABCA1 reduced mitophagy in dopaminergic neurons [41]. Experiments probing distinct steps in the mitophagy pathway found that probucol impacted mitophagy and in vivo phenotypes independent of PINK1/Parkin but required LDs, as pharmacological inhibition of LD biosynthesis abrogated probucol's mitophagy enhancing effect.

Under normal conditions, probucol had several relevant effects: It (1) increased LD-mitochondria contacts; (2) increased late endosomes/lysosomes; and (3) increased autophagosome lipidation. Importantly, probucol did not increase mitolysosome abundance under normal conditions but does so following mitochondrial damage. LDs adjacent to mitochondria can supply fatty acids during nutrient stress [42] and can interact with and transfer lipids and proteins to both lysosomes and autophagosomes [43,44]. LD mobilization by lipase PNPLA5 is required to facilitate the formation of autophagic membranes, including in the context of mitochondrial damage [45]. Given that we observed increased abundance of late endosomes/ lysosomes and mature autophagosomes under basal conditions, we speculate that the latter is true and may occur adjacent to mitochondria under basal conditions, but further investigation is required. Ultimately, the increased abundance of two components, which can subsequently fuse to form mito-lysosomes when mitophagy is induced, likely primes the cell for a more efficient and protective degradative response.

Cells and flies were subjected to prolonged mitochondrial damage in the form of CCCP treatment or paraquat feeding. Under these treatment conditions, we observed an increase in LD abundance. This is consistent with studies by Nguyen and colleagues and by Long and colleagues, which demonstrate increase LD levels upon starvation-induced autophagy and deferiprone-induced mitophagy [28,29]. The two studies attribute different roles for LDs in these contexts. LDs buffer lipotoxic species, which are generated as a byproduct of autophagic degradation and facilitate the subcellular transition endolysosomes undergo from the peripheries of the cell towards damaged perinuclear mitochondria, respectively. We did not investigate the reason for LD expansion occurs upon mitochondrial damage, but we did find that LDs were necessary for probucol's effects on mitophagy.

Lysosome position is critical for effective macroautophagy and found to be disrupted by inhibition of LD biosynthesis in deferiprone-induced mitophagy [29,46]. The increased abundance of LDs at mitochondria may facilitate the positioning of lysosomes away from the peripheries where they become active [46,47]. In vivo, clear overlap between LDs and lysosomes was visible both in dopaminergic neurons and in other cells of the fly brain, supporting this possibility.

Interestingly, LD expansion following mitochondrial damage was reduced in probucoltreated cells and flies. This feature of probucol's mechanism may be particularly relevant, given recent studies that found increased LD accumulation in the dopaminergic neurons of PD patients [48]. Likewise, the reduced accumulation supports the idea that LDs may be mobilized by probucol treatment to facilitate mitophagy [45]. Probucol mitigated both LD accumulation and mitochondrial damage—two features of PD pathogenesis. Our data points to probucol's canonical target, ABCA1, are required for both probucol's effects on mitophagy and on LD expansion. Targeting a point of crosstalk between these two pathogenesis pathways may be advantageous.

ABCA1 R219K gene polymorphisms impacts PD progression, as measured using the Hoehn and Yahr scale [49]. The K allele, which is associated with slower PD progression, also affects the lipid profile of those who carry the genotype. Compared to individuals carrying ABCA1 R219K RR or RK, ABCA1 R219K K genotype carriers have elevated high-density lipoprotein cholesterol and lower triglyceride levels [50]. It remains to be determined whether the differences in lipid profiles are responsible for clinical differences between these groups.

Across toxin-based and genetic models of mitochondrial damage and PD in two different species, probucol improved survival, locomotor function, and reduced the loss of dopaminergic neurons. A prior paper also found probucol-induced improvements to phenotypes caused by increased free radical production in *C. elegans*. In this study, how probucol facilitates these improvements was not examined but was postulated to occur via its antioxidant properties [51]. Given probucol's promising effects in several preclinical animal models, it might be fruitful to mine human epidemiologic data for any associations between probucol treatment and reduced risk of PD, since it is still in use in Japan and China. Statins also increase mitophagy, but in a Parkin-dependent manner, unlike probucol [52]. Whether statins impact LD expansion following mitochondrial damage may represent an interesting avenue for future inquiry.

In conclusion, our study showcased a dual in silico/cell-based screening methodology, which identified known and new mechanisms leading to mitophagy enhancement. ABCA1, which localizes to endolysosomes in cells and regulates lipid homeostasis, likely acts as a

mediator of crosstalk between LD dynamics and mitophagy since LDs are required for the mitophagy enhancement conferred by probucol [53].

Methods

Cells and tissue culture

All cell lines and sources are compiled in the Materials table (Table 1). Cells were cultured in Dulbecco's Modified Eagle Media supplemented with 10% fetal bovine serum. Cells were routinely tested for mycoplasma using the e-myco VALID mycoplasma testing kit (FroggaBio, 25239). Cells were maintained at 37° C temperature and 5% CO₂ in a humidified atmosphere.

Immunoblotting

Lysates were harvested with lysis buffer (0.1 M Tris–HCl, 0.01% SDS (pH 9)) containing protease inhibitor cocktail (BioShop, PIC002.1) followed by 20 minutes of boiling and vortexing at 95°C. Lysates were then pelleted by high-speed centrifugation for 20 minutes at 4°C, and supernatant was transferred into a new microcentrifuge tube for BCA assays to standardize protein loading across each experiment (Pierce, 23227).

Immunoblotting for VDAC1, ATP5A, ABCA1 was performed with 10% SDS-PAGE gels to separate proteins followed by transfer onto PVDF membrane (Immobilon, IPVH00010) at 110 V for 80 minutes or 8-hour 38 V transfer in the cold room. Samples to be probed for LC3 were separated on 15% SDS-PAGE gels instead. The transfer apparatus setup included an ice pack and a stir bar.

After Ponceau staining and imaging to assess overall protein loading, membranes were washed with TBST and blocked in 5% skim milk diluted in TBST for 30 minutes at room temperature. Incubation in primary antibodies diluted in 2.5% skim milk was performed overnight at 4°C. Three 5-minute TBST washes were performed prior to incubation of blots in secondary antibody diluted in 2.5% milk for 2 hours. Three final 10-minute TBST washes were followed by visualization of proteins with ECL (BioRad, 11705062). Densitometry analysis was performed using ImageLab 6.0 software (BioRad), and the protein of interest was normalized to either Ponceau staining or loading controls such as GAPDH or tubulin. Antibodies used in this study are compiled in Materials table (Table 1).

Immunofluorescence

Cells grown on coverslips (1.5 H thickness) were fixed with 4% PFA for 15 minutes, followed by 3 PBS washes. Permeabilization was performed with 0.1% Tx-100 incubation for 15 minutes, followed by 3 more PBS washes. Blocking with 10% goat serum diluted in PBS was performed for 30 minutes at room temperature, or overnight at 4°C. Coverslips were then incubated in primary antibody diluted in 1% goat serum (1:500) overnight at 4°C. The next day, following 3 PBS washes, coverslips were incubated in secondary antibody diluted in 1% goat serum (1,500) for 1 to 2 hours at room temperature. Three 10-minute PBS washes were followed by mounting onto slides with Fluoromount containing DAPI (Invitrogen 00495952).

Immunofluorescence of fly brains began with 20-minute fixation with 4% PFA, followed by 3× washes with 0.1% PBST (Tween-20 diluted in PBS) washes. Permeabilization was performed with 1% PBST (Triton-X-100 diluted in PBS) for 1 hour, followed by 3× washes in PBS. Blocking of fly brains in 10% goat serum for 1 hour was followed with overnight incubation at 4°C in primary antibody diluted in 1% goat serum. The next day, 10-minute 0.1% PBST (Tween-20 diluted in PBS) washes were followed by incubation in secondary antibody diluted in 1% goat serum for 2 hours. Three more 10-minute 0.1% PBST washes and a final 10-minute

Table 1. Materials.

MATERIAL	SOURCE	IDENTIFIER
Antibodies		
mouse monoclonal anti-ATP5A	Abcam	14748
mouse monocloncal anti-ABCA1	Abcam	18180
mouse monoclonal anti-VDAC1	Abcam	14734
rabbit polyclonal anti-phospho ubiquitin S65	Sigma Aldrich	1513-I
mouse monoclonal anti-tubulin	Santa Cruz	sc-5286
mouse monoclonal anti-actin	Abcam	ab8226
rabbit polyclonal anti-LC3B	Cell Signaling	2775
mouse monoclonal anti-GAPDH	Invitrogen	398600
polyclonal rabbit horseradish peroxidase-conjugated secondary antibody	Jackson Immunoresearch Laboratories	111035144
polyclonal mouse horseradish peroxidase-conjugated secondary antibody	Jackson Immunoresearch Laboratories	115035003
Alexa Fluor 647 donkey anti-rabbit	Invitrogen	A31573
Alexa Fluor 594 goat anti-mouse	Invitrogen	A11004
Chemicals		
probucol (cells: 1 µM, flies: 0.25 mM, zebrafish: 2 µM)	Tocris	2775
СССР (10 µМ)	Sigma Aldrich	2759
Τ863 (20 μM)	Cayman Chemical Co.	258007
PF-06424439 (10 μM)	Tocris	6348
leupeptin (0.5 mM)	Bioshop	LEU001
E-64 (2 µM)	Bioshop	EEL640
Paraquat (0.5 mM)	Sigma Aldrich	36541
SR-3677 (2 μM)	Tocris	3677
BODIPY (1 µg/mL)	Invitrogen	D3922
Bafilomycin (2.5 nM)	Cell Signaling Technology	54645
MPP ⁺ (1 mM)	Sigma Aldrich	D048
Cell lines		
Human: HeLa	Peter Kim lab	N/A
Human: HeLa cells stably expressing GFP Parkin and mito-DsRed	Richard Youle lab	N/A
Human: HEK293	Peter Kim lab	N/A
Organisms		
Drosophila: TH-Gal4	BDSC	8848
Drosophila: UAS-park shRNA	BDSC	31259
Drosophila: heteroplasmic mt:Coll ^{T300I}	Thomas Hurd lab	N/A
Drosophila: Canton(S)	Thomas Hurd lab	N/A
Drosophila: UAS-hABCA1/CvO	BDSC	84758
Drosophila: UAS-ABCA shRNA	BDSC	38353
Drosophila: UAS-mitoOCattP2	Alex Whitworth lab	N/A
Drosophila: UAS-mitoOCattP16	Alex Whitworth lab	N/A
Drosophila: UAS-mCherry shRNA	BDSC	35785
Drosophila: dLamp-3xmCherry	Gabor Juhasz lab	N/A
Zebrafish: Tg(dat:EGFP)	Wen lab	N/A
Zebrafish: WT Tuebingen	Zebrafish International Resource Center	ZI 57
Recombinant DNA		
Plasmid: pLKO1	Sigma Aldrich	SHC001
Plasmid: shABCA1	Sigma Aldrich	TRCN000029093
Plasmid: Cerulean-Parkin	Peter Kim, Sick Kide Hospital	N/A
Plasmid: mitoOC	Peter Kim, Sick Kids Hospital	N/A
	- ever min, oren mus mospitui	

(Continued)

Table 1. (Continued)

MATERIAL	SOURCE	IDENTIFIER
Software and algorithms		
ImageJ	NIH	https://imagej.net/software/fiji/
Cell Profiler	Carpenter-Singh lab, Broad Institute	https://cellprofiler.org/
Cell Profiler Analyst	Carpenter-Singh lab, Broad Institute	https://cellprofileranalyst.org/
Watson Drug Discovery	IBM	N/A
GraphPad Prism v9		https://www.graphpad.com/
BioRender		https://biorender.com/

https://doi.org/10.1371/journal.pbio.3001977.t001

PBS wash were performed prior to overnight incubation of fly brains in Fluoromount (Invitrogen 00495802) and mounting the next day onto slides. Coverslips (1.5 H thickness) were sealed to the slide with clear nail polish. Antibodies used in this study are compiled in Materials table (Table 1).

Fly husbandry

Fly lines used in this study are compiled in Materials table (Table 1). Drosophila were maintained at 25 °C and at 70% relative humidity in 12-hour light/dark cycles and were fed standard yeast-molasses-sugar-agar formula. In the case of drug treatments, low melt agar fly food formula was composed as previously described [54]. Probucol was added to low melt agar fly food at 250 μ M concentration, and the equivalent volume of DMSO was used in control vials. DMSO concentration never exceeded 1% in experiments. Fly stocks used in the study and sources are listed in Materials table (Table 1).

In silico screen for mitophagy enhancers in Drugbank

We have previously described the natural language processing methodology employed by IBM Watson for Drug discovery predictive analytics [11,12]. In brief, a set of candidate drugs were ranked according to semantic similarity to a training set of drugs known to have the desirable biological effect using natural language processing applied to published abstracts obtained from Medline.

- **Training set:** A training set consisting of compounds listed in S1A Fig was composed with reference to a review about pharmacological modulators of mitophagy [2]. This list was curated to filter out any that were associated with mitochondrial damage or apoptosis (S2 Dataset). Compounds associated with key words such as apoptosis, depolarization, and mitochondrial damage were excluded. The result was a set of 7 compounds with proven ability to induce mitophagy without an association with mitochondrial damage or apoptosis; Olaparib (PARP inhibitor), nicotinamide (NAD⁺ accumulation), pifitherin- α (p53 inhibitor), kinetin (PINK1 neo-substrate), and the SIRT1 activators resveratrol, fisetin, and SRT1720.
- **Candidate set:** A total of 3,231 final candidates were filtered from the entire DrugBank database (www.drugbank.ca). Candidates with less than 5 published abstracts were removed.

Model validation

1. Leave-one-out cross validation: Leave-one-out cross validation was performed. The ranking was run 7 times, with each training drug in turn removed from the training set and ranked among the other 3,231 candidate drugs. Receiver operating characteristic (ROC) curves were generated across the range of possible ranks to assess the model's performance in a binary classification task. Data were analyzed in Python using the scikit-learn library.

- 2. **Retrospective analysis:** Olaparib was first published as having a mitophagy inducing effect in 2015 [55]. To further validate our methodology, a ranking was performed restricted to abstracts published up to and including 2014. Olaparib was omitted from the known set and placed in the candidate list.
- **ROC Curve:** An ROC curve was constructed using these data, and the area under the curve (AUC) was calculated to assess the predictive power of the model (S1B Fig). An AUC value of 1 indicates perfect predictive ability, while <0.5 indicates predictive ability worse than random chance. The AUC for the model created by IBM Watson for Drug Discovery was 0.9513.
- **Ranking of candidate set and selection of candidates for validation:** The final ranking was applied resulting in a list of 3,231 candidate drugs rank ordered according to semantic similarity to the training set.

Following post hoc analysis, the top 79 molecules ranked as bearing highest semantic similarity to the training set molecules were purchased from Sigma. The identities of the compounds in our custom library are described in <u>S3 Dataset</u>. These molecules were arrayed in a 96-well format at 10 mM concentration in DMSO.

Screening

A total of 20,000 HeLa cells stably expressing mito-DsRed and GFP Parkin were seeded into clear, flat bottom, black polystyrene 96-well plates (Corning, CLS3603) and incubated at 37°C with 5% CO_2 overnight. A final concentration of 1 μ M of the compound in media was added to the cells for 2 hours prior to the addition of CCCP to a final concentration of 10 μ M. CCCP treatment was stopped after 24-hour incubation by washing cells with PBS followed by fixation in 4% PFA. Imaging was performed on the Cytell Cell Imaging System at 10X magnification. Eight fields were captured with the same ROI positions in each well.

Image analysis for mitochondrial clearance

The mitochondrial clearance screen was quantified first by segmenting every cell in each image. Cells were stained with DAPI to visualize nuclei, and whole-cell segmentation was also performed in the GFP channel, based on Parkin distribution. Cells were categorized into two groups, based on mito-DsRed intensity, a matrix-targeted fluorophore that serves as a mito-chondrial marker. Cells were classified as either positive or negative for mito-DsRed signal based on a fluorescence intensity cutoff determined by assessing the fluorescence intensity signal for both the negative and positive controls for the screen, DMSO pretreated cells treated with DMSO (negative) or CCCP (positive) for 24 hours.

The average mitochondrial clearance (% of cells with low/no mito-DsRed signal) for the positive control (DMSO+24 hour CCCP) and negative control (DMSO+24 hour DMSO) wells was determined and used to calculate the Z-factor, which is a metric indicative of screening robustness [56]. Normalized MAD z-scores were calculated for each of the compounds, as implemented in other mitophagy-related screens [57], across two independent screening replicates. Molecules with the highest z-scores had the strongest effect on mitochondrial clearance.

mitoQC assay to assess mitochondria-to-lysosome targeting in cells

HeLa cells were seeded onto coverslips in 6-well plates. The next day, they were transfected with Flag-Parkin using lipofectamine 2000 (Invitrogen, 11668019). Twenty-four hours later, cells were treated with 10 μ M CCCP (or DMSO as a control) in combination with 0.5 mM leupeptin and 2 μ M E-64 in combination with 1 μ M of the indicated small molecule treatments or DMSO for 6 hours prior to fixation.

Images were acquired on the Leica SP8 microscope. All image settings were kept consistent throughout imaging for each experiment. The ImageJ Plug-In developed by Garriga and colleagues was used to analyze the percentage of mitochondria, which appear red-only [58]: https://github.com/graemeball/mQC_counter. At least 40 cells were quantified per treatment group in each replicate. Cells with at least 5 red-only puncta representing mito-lysosomes were classified mitophagic, similar to the analysis described by Allen and colleagues [20].

In vivo mitoQC assay to assess mitochondria-to-lysosome targeting

Seven-day old flies were placed into vials containing low melt agar in combination with the indicated treatments. Following 24-hour incubation, flies were incubated in whole-fly fixation reagent containing 1% PFA and 0.1% PBST (Tween-20 diluted in PBS) overnight at 4°C. Fly brains were dissected following PBS wash and subsequently fixed in 4% PFA for 20 minutes. After a 5-minute PBST wash and a 5-minute PBS wash, fly brains were incubated in Fluoro-mount. Following tissue dissection, all steps were performed protected from light. Brains were mounted onto slides and covered with #1.5 coverslips and sealed with nail polish. At least 3 independent biological trials were performed for each experiment, and at least 2 fly brains were imaged for each treatment.

Zebrafish locomotor and dopaminergic neuron assays

Zebrafish locomotion was evaluated in 96-well plate format using Zebrabox video tracking system. The Tg(*dat:EGFP*) zebrafish model and pertinent details to the quantification of dopaminergic neurons in the ventral diencephalon region of the brain has previously been described in detail [27]. For confocal microscopy of dopaminergic neurons, embryos were cleaned and housed in a 28°C incubator. Twenty-four hours later, 1 mM MPP⁺ was administered with probucol at 50 μ M concentration and DMSO control. Microscopy was performed the next day. For locomotor assays, instead of performing microscopy, media was changed to fresh MPP⁺ and drug for 2 additional days, then on the on the final day, the movement trajectory of the zebrafish was tracked using ZebraBox.

Measuring LD and lysosome area

ImageJ was used to create maximum intensity projections of image z-stacks. The object counter Plugin was then employed to identify circular objects in the images and measure the area of puncta. Since variation in cell size was evident across experiments, the area of the cell structure of interest was normalized to the total cell area. TH staining was used to determine total area of dopaminergic neurons, while the brightfield was used to define cell area in culture.

Statistical analysis and figures

Unpaired Student *t* tests were performed to make pairwise comparisons. Where multiple comparisons were made, ANOVA statistical analysis was performed with Dunnett's multiple comparison correction. Independent biological replicates were used to compute statistics using GraphPad Prism v 9.0 software. Where possible, each independent biological replicate is displayed in the graphs. Error bars on graphs represent the SEM, as several technical replicates were averaged to obtain each independent replicate value. BioRender (www.biorender.com) was used to make Figs 4F, S1C and S3A.

Supporting information

S1 Fig. In silico screen to identify candidate mitophagy enhancers in DrugBank database of drugs amenable to repurposing. (A) Leave-one-out cross validation and retrospective analyses were performed to evaluate the ability of the model to identify bona fide mitophagy enhancers. Similarity scores between each chemical assessed and the mitophagy enhancer training set were calculated and used to assign a rank out of 3,231 DrugBank molecules from most to least similar in ascending order. (B) Leave-one-out cross validation results were used to construct a receiver operating characteristic curve, which demonstrates the predictive performance of the model. The area under the curve for the ROC curve is 0.9513. (C) Following validation, the model was deployed to identify new mitophagy enhancers from the DrugBank candidates based on information from a wide array of sources including PubMed literature, patent filings, and biological databases. (PDF)

S2 Fig. Cell-based mitochondrial clearance screen to evaluate candidates identified in silico. (A) Normalized MAD z-scores of 79 candidate DrugBank molecules screened in the mitochondrial clearance screen. Probucol, 3-methoxybenzamide, and lapatinib are highlighted among other compounds in descending order of rank. (B) Chemical structures of hit molecules highlighted in A. (C) HeLa cells expressing GFP-Parkin and mito-DsRed. Cells were pretreated with small molecules $(1 \,\mu M)$ for 2 hours followed by 24-hour treatment with CCCP $(10 \,\mu\text{M})$ to induce prolonged mitophagy, resulting in loss of mito-DsRed signal from many cells (mitochondrial clearance). Arrows denote cells that retained mitochondrial signal. The percentage of remaining cells with no/low mito-DsRed signal was calculated as the screening readout. (D) Immunoblotting using antibodies for outer mitochondrial membrane protein VDAC1 was performed on lysates from cells pretreated with small molecule (1 μ M) prior to CCCP (10 µM) time course. (E) Quantification of VDAC1 levels normalized to Ponceau staining to assess protein loading. Data information: Normalized MAD z-score values were calculated based on two independent screening replicates in A. Four independent biological replicates were performed for E. Bars represent mean values, and error bars represent SEM. * indicates p-value < 0.05. Statistical analysis was performed using an unpaired two-sided Student *t* test to compare DMSO and probucol at each time point. The data underlying the graphs shown in the figure can be found in S1 Data and S3 Dataset. (PDF)

S3 Fig. Mitochondrial clearance screening workflow. (A) On Day 1, GFP-Parkin mito-DsRed HeLa cells are seeded in 96-well plates. On Day 2, cells are pretreated with 1 μ M concentration of the small molecule library for 2 hours prior to the addition of 10 μ M CCCP for 24 hours. Cells are then fixed and DAPI staining is performed to visualize nuclei. Cell Profiler and Cell Profiler Analyst tools are used to differentiate cells that retain mito-DsRed signal and ones with no/low mito-DsRed signal. (B) Mitochondrial clearance values for positive control wells pretreated with DMSO for 2 hours in place of small molecules and followed by 24-hour treatment with CCCP to induce mitophagy and negative control wells containing cells treated with DMSO alone. (C) Z-factor values calculated from the positive and negative control replicate wells in each of the independent biological screening replicates. The data underlying the graphs shown in the figure can be found in <u>S1 Data</u>. (PDF)

S4 Fig. Immunoblotting to assess ATP5A levels following prolonged mitochondrial damage. (A) Immunoblotting whole-cell lysates using antibodies for inner mitochondrial membrane protein ATP5A. HeLa cells stably expressing GFP-Parkin were treated with the indicated drugs at 1 μ M concentration in combination with 10 μ M CCCP. Irrelevant lane in the center of the blot was removed for clarity, but both right and left side of blot and Ponceau correspond to the same image from the same membrane. Ponceau staining was used to visualize protein loading. (B) Densitometry analysis was performed to assess ATP5A levels normalized to Ponceau loading. The data underlying the graphs shown in the figure can be found in S1 Data.



S5 Fig. PINK1-mediated phosphorylation of mitochondrial ubiquitin and Parkin recruitment are not affected by probucol treatment. (A) HeLa cells expressing GFP-Parkin were pretreated with probucol or DMSO for 2 hours prior to 1-hour treatment with CCCP. Immunostaining with antibody specific for phospho-Ubiquitin S65 (p-Ub S65) was performed on cells. (B) The percentage of cells that are positive for mitochondrial p-Ub S65 signal and in which (C) Parkin distribution is mitochondrial was calculated for each treatment. (D) Wholecell lysates from treatments as described in A were probed with antibody against p-Ub S65. Ponceau staining was used to assess protein loading. Three independent biological replicates were performed for all experiments and are represented by data points, and bars depict means. Error bars display SEM. ANOVA statistical analysis with Dunnett's multiple comparison correction was performed; * and ** indicate *p*-value <0.05 and 0.01, respectively. The data underlying the graphs shown in the figure can be found in <u>S1 Data</u>. (PDF)

S6 Fig. Probucol improves climbing defects and survival decline, which arise from mtDNA mutation $mt:ColI^{T300I}$. Heteroplasmic flies with $mt:ColI^{T300I}$ in approximately 90% of their mtDNA were fed food supplemented with either DMSO or probucol. (A) The percentage of flies climbing beyond a height of 12.5 cm is displayed, in addition to (B) the survival of the flies in both groups. As a control, heteroplasmic $mt:ColI^{T300I}$ were maintained at permissive temperature of 18°C. Three independent biological replicates were performed for both A and B, and at least 20 flies were included in each replicate. Bars represent mean values in A, and error bars represent SEM. For A, unpaired Student *t* tests were performed to evaluate differences between DMSO and probucol. For B, log-rank test analysis was performed to compare the survival of the two treatment groups housed at 29°C. * indicates *p*-value <0.05. The data underlying the graphs shown in the figure can be found in S1 Data. (TIFF)

S7 Fig. Effects of ABCA1 manipulations on mitophagy. (**A**) Immunoblotting was performed to assess ABCA1 levels in lysates derived from the brains of flies in which human ABCA1 transgene expression is driven by pan-neuronal elav-Gal4. (**B**) Immunoblotting was performed using antibodies against ABCA1 and tubulin as a loading control on HeLa cells lysates from cells transfected with either pLKO1 control or shABCA1 vectors. The blots pictured in A and B are representative of 3 independent biological replicates. (**C**) HeLa cells expressing the mitoQC reporter and Cerulean-Parkin were transfected with expression of the mitoQC reporter in the presence and absence of the human ABCA1 transgene induced by the TH-Gal4 driver. Food containing the indicated combinations of probucol and paraquat was

administered. (PDF)

S8 Fig. Lipid droplet area expansion following CCCP treatment is reduced by probucol treatment. HeLa cells were treated with the indicated combination of DMSO, probucol, CCCP, and DGAT inhibitors. BODIPY staining was performed to visualize LDs and immunostaining with ATP5A antibody. (PDF)

S9 Fig. Effect of probucol on LC3 lipidation under basal conditions, following mitochondrial depolarization and starvation. (A) HEK293 cells were either incubated in DMEM, DMEM with CCCP, or HBSS media for 6 hours. Lysates were separated by SDS-PAGE, and immunoblotting was performed using antibodies that recognize LC3 and GAPDH, as a loading control. (B) Densitometry was performed to measure the levels of lipidated LC3-II, which were normalized to the GAPDH loading control. Unpaired *t* tests were used to evaluate differences between DMSO and probucol. * indicates *p*-value <0.05. The data underlying the graphs shown in the figure can be found in S1 Data. (PDF)

S1 Dataset. In silico candidates from DrugBank ranked in order of similarity to mitophagy enhancer training set.

(XLSX)

S2 Dataset. In silico candidate molecules associated with terms "mitochondrial damage" and "apoptosis."

(XLS)

S3 Dataset. Results of cell-based mitochondrial clearance screen. (XLSX)

S1 Data. Source data. (XLSX)

S1 Raw Images. Uncropped images of data in figures. (SVG)

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Author Contributions

Conceptualization: Natalia Moskal, G. Angus McQuibban.

Data curation: Natalia Moskal, Vijay Narasimhan.

Formal analysis: Natalia Moskal, Naomi P. Visanji, Vijay Narasimhan.

Funding acquisition: Peter N. Lewis, G. Angus McQuibban.

Investigation: Natalia Moskal, Naomi P. Visanji, Olena Gorbenko, Vijay Narasimhan, Hannah Tyrrell, Jess Nash.

Methodology: Natalia Moskal, Vijay Narasimhan.

Supervision: Peter N. Lewis, G. Angus McQuibban.

Writing - original draft: Natalia Moskal.

Writing – review & editing: Naomi P. Visanji, Olena Gorbenko, Peter N. Lewis, G. Angus McQuibban.

References

- Poewe W, Seppi K, Tanner CM, Halliday GM, Brundin P, Volkmann J, et al. Parkinson disease. Nat Rev Dis Primers. 2017; 3:1–21.
- Kitada T, Asakawa S, Hattori N, Matsumine H, Yamamura Y, Minoshima S, et al. Mutations in the parkin gene cause autosomal recessive juvenile parkinsonism. Nature [Internet]. 1998 Apr 9; 392(6676):605– 608. https://doi.org/10.1038/33416 PMID: 9560156
- Valente EM, Abou-sleiman PM, Caputo V, Muqit MMK, Harvey K, Gispert S, et al. Hereditary Early-Onset Parkinson's Disease Caused by Mutations in PINK1. Science. 2004; 304(May):1158–1161. https://doi.org/10.1126/science.1096284 PMID: 15087508
- Ordonez DG, Lee MK, Feany MB. α-synuclein Induces Mitochondrial Dysfunction through Spectrin and the Actin Cytoskeleton. Neuron. 2018; 97(1):108–124.e6.
- Li H, Ham A, Ma TC, Kuo SH, Kanter E, Kim D, et al. Mitochondrial dysfunction and mitophagy defect triggered by heterozygous GBA mutations. Autophagy. 2019; 15(1):113–130. https://doi.org/10.1080/ 15548627.2018.1509818 PMID: 30160596
- Wauters F, Cornelissen T, Imberechts D, Martin S, Koentjoro B, Sue C, et al. LRRK2 mutations impair depolarization-induced mitophagy through inhibition of mitochondrial accumulation of RAB10. Autophagy. 2020; 16(2):203–222. <u>https://doi.org/10.1080/15548627.2019.1603548</u> PMID: 30945962
- LaVoie MJ, Ostaszewski BL, Weihofen A, Schlossmacher MG, Selkoe DJ. Dopamine covalently modifies and functionally inactivates parkin. Nat Med. 2005; 11(11):1214–1221. <u>https://doi.org/10.1038/</u> nm1314 PMID: 16227987
- Chung KKK, Thomas B, Li X, Pletnikova O, Troncoso JC, Marsh L, et al. S-nitrosylation of parkin regulates ubiquitination and compromises parkin's protective function. Science (1979). 2004; 304 (5675):1328–1331.
- Tokarew JM, El-Kodsi DN, Lengacher NA, Fehr TK, Nguyen AP, Shutinoski B, et al. Age-associated insolubility of parkin in human midbrain is linked to redox balance and sequestration of reactive dopamine metabolites. Acta Neuropathol. 2021; 141(5):725–754. <u>https://doi.org/10.1007/s00401-021-02285-4</u> PMID: 33694021
- 10. Georgakopoulos ND, Wells G, Campanella M. The pharmacological regulation of cellular mitophagy. Nat Chem Biol. 2017; 13(2):136–146. https://doi.org/10.1038/nchembio.2287 PMID: 28103219
- Maclagan LC, Visanji NP, Cheng Y, Tadrous M, Lacoste AMB, Kalia L, et al. Identifying drugs with disease-modifying potential in Parkinson's disease using artificial intelligence and pharmacoepidemiology. Pharmacoepidemiol Drug Saf. 2020; 29(8):864–872. https://doi.org/10.1002/pds.5015 PMID: 32410265
- Visanji NP, Madan P, Lacoste AMB, Buleje I, Han Y, Spangler S, et al. Using artificial intelligence to identify anti-hypertensives as possible disease modifying agents in Parkinson's disease. Pharmacoepidemiol Drug Saf. 2021; 30(2):201–209. https://doi.org/10.1002/pds.5176 PMID: 33219601
- Chen KS, Menezes K, Rodgers JB, O'Hara DM, Tran N, Fujisawa K, et al. Small molecule inhibitors of α-synuclein oligomers identified by targeting early dopamine-mediated motor impairment in C. elegans. Mol Neurodegener. 2021; 16(1):1–25.
- Ashburn TT, Thor KB. Drug repositioning: Identifying and developing new uses for existing drugs. Nat Rev Drug Discov. 2004; 3(8):673–683. https://doi.org/10.1038/nrd1468 PMID: 15286734
- Narendra D, Tanaka A, Suen DF, Youle RJ. Parkin is recruited selectively to impaired mitochondria and promotes their autophagy. J Cell Biol. 2008; 183(5):795–803. <u>https://doi.org/10.1083/jcb.200809125</u> PMID: 19029340
- Villa E, Marchetti S, Ricci JE. No Parkin Zone: Mitophagy without Parkin. Trends Cell Biol [Internet]. 2018; 28(11):882–895. https://doi.org/10.1016/j.tcb.2018.07.004 PMID: 30115557
- Pickrell AM, Youle RJ. The roles of PINK1, Parkin, and mitochondrial fidelity in parkinson's disease. Neuron. 2015; 85(2):257–273. https://doi.org/10.1016/j.neuron.2014.12.007 PMID: 25611507

- Wang Y, Serricchio M, Jauregui M, Shanbhag R, di Paolo CT, Kim PK, et al. Deubiquitinating enzymes regulate PARK2- mediated mitophagy. Autophagy. 2015; 8627. <u>https://doi.org/10.1080/15548627</u>. 2015.1034408 PMID: 25915564
- Hasson SA, Kane LA, Yamano K, Huang CH, Sliter DA, Buehler E, et al. High-content genome-wide RNAi screens identify regulators of parkin upstream of mitophagy. Nature. 2013; 504(7479):291–295. https://doi.org/10.1038/nature12748 PMID: 24270810
- Allen GFG, Toth R, James J, Ganley IG. Loss of iron triggers PINK1/Parkin-independent mitophagy. EMBO Rep. 2013; 14(12):1127–1135. https://doi.org/10.1038/embor.2013.168 PMID: 24176932
- Chaudhuri A, Bowling K, Funderburk C, Lawal H, Inamdar A, Wang Z, et al. Interaction of genetic and environmental factors in a Drosophila parkinsonism model. J Neurosci. 2007; 27(10):2457–2467. https://doi.org/10.1523/JNEUROSCI.4239-06.2007 PMID: 17344383
- 22. Moskal N, Riccio V, Bashkurov M, Taddese R, Datti A, Lewis PN, et al. ROCK inhibitors upregulate the neuroprotective Parkin-mediated mitophagy pathway. Nat Commun. 2020; 11(1):1–14.
- Wen L, Wei W, Gu W, Huang P, Ren X, Zhang Z, et al. Visualization of monoaminergic neurons and neurotoxicity of MPTP in live transgenic zebrafish. Dev Biol. 2008; 314(1):84–92. <u>https://doi.org/10.1016/j.ydbio.2007.11.012</u> PMID: 18164283
- Chen Z, Qi Y, French S, Zhang G, Garcia RC, Balaban R, et al. Genetic mosaic analysis of a deleterious mitochondrial DNA mutation in Drosophila reveals novel aspects of mitochondrial regulation and function. Mol Biol Cell. 2015; 26(4):674–684. https://doi.org/10.1091/mbc.E14-11-1513 PMID: 25501370
- Ma H, Xu H, O'Farrell PH. Transmission of mitochondrial mutations and action of purifying selection in Drosophila melanogaster. Nat Genet. 2014; 46(4):393–397. <u>https://doi.org/10.1038/ng.2919</u> PMID: 24614071
- Kandul NP, Zhang T, Hay BA, Guo M. Selective removal of deletion-bearing mitochondrial DNA in heteroplasmic Drosophila. Nat Commun. 2016; 7:1–11. <u>https://doi.org/10.1038/ncomms13100</u> PMID: 27841259
- Xi Y, Yu M, Godoy R, Hatch G, Poitras L, Ekker M. Transgenic zebrafish expressing green fluorescent protein in dopaminergic neurons of the ventral diencephalon. Dev Dyn. 2011; 240(11):2539–2547. https://doi.org/10.1002/dvdy.22742 PMID: 21932324
- Nguyen TB, Louie SM, Daniele JR, Tran Q, Dillin A, Zoncu R, et al. HHS Public Access. 2018; 42(1):9– 21.
- Long M, Sanchez-Martinez A, Longo M, Suomi F, Stenlund H, Johansson AI, et al. DGAT 1 activity synchronises with mitophagy to protect cells from metabolic rewiring by iron depletion. EMBO J. 2022:1–25.
- Benador IY, Veliova M, Liesa M, Shirihai OS. Mitochondria Bound to Lipid Droplets: Where Mitochondrial Dynamics Regulate Lipid Storage and Utilization. Cell Metab. 2019; 29(4):827–835. <u>https://doi.org/ 10.1016/j.cmet.2019.02.011</u> PMID: 30905670
- 31. Mizushima N, Yoshimori T. How to interpret LC3 immunoblotting. Autophagy. 2007; 3(6):542–545. https://doi.org/10.4161/auto.4600 PMID: 17611390
- Liebl MP, Meister SC, Frey L, Hendrich K, Klemmer A, Wohlfart B, et al. Robust LC3B lipidation analysis by precisely adjusting autophagic flux. Sci Rep. 2022; 12(1):1–14.
- Mak SK, McCormack AL, Manning-Bog AB, Cuervo AM, di Monte DA. Lysosomal degradation of αsynuclein in vivo. J Biol Chem. 2010; 285(18):13621–13629.
- Zhu JH, Gusdon AM, Cimen H, van Houten B, Koc E, Chu CT. Impaired mitochondrial biogenesis contributes to depletion of functional mitochondria in chronic MPP b toxicity: dual roles for ERK1 / 2. Cell Death Dis 2012; 3(5):e312–10. https://doi.org/10.1038/cddis.2012.46 PMID: 22622131
- Ruiz LM, Jensen EL, Rossel Y, Puas GI, Gonzalez-Ibanez AM, Bustos RI, et al. Non-cytotoxic copper overload boosts mitochondrial energy metabolism to modulate cell proliferation and differentiation in the human erythroleukemic cell line K562. Mitochondrion. 2016; 29:18–30. <u>https://doi.org/10.1016/j.mito.</u> 2016.04.005 PMID: 27094959
- Tsang T, Posimo JM, Gudiel AA, Cicchini M, Feldser DM, Brady DC. Copper is an essential regulator of the autophagic kinases ULK1/2 to drive lung adenocarcinoma. Nat Cell Biol. 2020; 22(4):412–424. https://doi.org/10.1038/s41556-020-0481-4 PMID: 32203415
- Laker RC, Drake JC, Wilson RJ, Lira VA, Lewellen BM, Ryall KA, et al. Ampk phosphorylation of Ulk1 is required for targeting of mitochondria to lysosomes in exercise-induced mitophagy. Nat Commun. 2017; 8(1):548. https://doi.org/10.1038/s41467-017-00520-9 PMID: 28916822
- Wang C, Wang H, Zhang D, Luo W, Liu R, Xu D, et al. Phosphorylation of ULK1 affects autophagosome fusion and links chaperone-mediated autophagy to macroautophagy. Nat Commun. 2018; 9(1):3492. https://doi.org/10.1038/s41467-018-05449-1 PMID: 30154410

- Kume S, Uzu T, Horiike K, Chin-Kanasaki M, Isshiki K, Araki S, et al. Calorie restriction enhances cell adaptation to hypoxia through Sirt1-dependent mitochondrial autophagy in mouse aged kidney. J Clin Invest. 2010; 120(4):1043–1055. https://doi.org/10.1172/JCI41376 PMID: 20335657
- Brakedal B, Dölle C, Riemer F, Ma Y, Nido GS, Skeie GO, et al. The NADPARK study: A randomized phase I trial of nicotinamide riboside supplementation in Parkinson's disease. Cell Metab. 2022; 34 (3):396–407.e6. https://doi.org/10.1016/j.cmet.2022.02.001 PMID: 35235774
- Wu CA, Tsujita M, Hayashi M, Yokoyama S. Probucol inactivates ABCA1 in the plasma membrane with respect to its mediation of apolipoprotein binding and high density lipoprotein assembly and to its proteolytic degradation. J Biol Chem. 2004; 279(29):30168–30174. <u>https://doi.org/10.1074/jbc.</u> M403765200 PMID: 15140889
- 42. Stone SJ, Levin MC, Zhou P, Han J, Walther TC, Farese R, et al. The endoplasmic reticulum enzyme DGAT2 is found in mitochondria-associated membranes and has a mitochondrial targeting signal that promotes its association with mitochondria. J Biol Chem. 2009; 284(8):5352–5361. https://doi.org/10. 1074/jbc.M805768200 PMID: 19049983
- Singh R, Kaushik S, Wang Y, Xiang Y, Novak I, Komatsu M, et al. Autophagy regulates lipid metabolism. Nature. 2009; 458(7242):1131–1135. https://doi.org/10.1038/nature07976 PMID: 19339967
- 44. Schulze RJ, Krueger EW, Weller SG, Johnson KM, Casey CA, Schott MB, et al. Direct lysosome-based autophagy of lipid droplets in hepatocytes. Proc Natl Acad Sci U S A. 2020; 117(51):32443–32452. https://doi.org/10.1073/pnas.2011442117 PMID: 33288726
- Thazar-Poulot N, Miquel M, Fobis-Loisy I, Gaude T, Dupont N, Chauhan S, et al. Neutral lipid stores and lipase PNPLA5 contribute to autophagosome biogenesis. Curr Biol. 2015; 24(6):4158–4163.
- 46. Korolchuk VI, Saiki S, Lichtenberg M, Siddiqi FH, Roberts EA, Imarisio S, et al. Lysosomal positioning coordinates cellular nutrient responses. Nat Cell Biol. 2011; 13(4):453–462. <u>https://doi.org/10.1038/ncb2204</u> PMID: 21394080
- Johnson DE, Ostrowski P, Jaumouillé V, Grinstein S. The position of lysosomes within the cell determines their luminal pH. J Cell Biol. 2016; 212(6):677–692. https://doi.org/10.1083/jcb.201507112 PMID: 26975849
- Brekk OR, Honey JR, Lee S, Hallett PJ, Isacson O. Cell type-specific lipid storage changes in Parkinson's disease patient brains are recapitulated by experimental glycolipid disturbance. Proc Natl Acad Sci U S A. 2020; 117(44):27646–27654. https://doi.org/10.1073/pnas.2003021117 PMID: 33060302
- 49. Ya L, Lu Z. Differences in ABCA1 R219K polymorphisms and serum indexes in alzheimer and parkinson diseases in Northern China. Med Sci Monit. 2017; 23:4591–4600. <u>https://doi.org/10.12659/msm.</u> 903636 PMID: 28943632
- Shi Z, Tian Y, Zhao Z, Wu Y, Hu X, Li J, et al. Association between the ABCA1 (R219K) polymorphism and lipid profiles: a meta-analysis. Sci Rep. 2021; 11(1):1–10.
- Ray A, Martinez BA, Berkowitz LA, Caldwell GA, Caldwell KA. Mitochondrial dysfunction, oxidative stress, and neurodegeneration elicited by a bacterial metabolite in a *C. elegans* Parkinson's model. Cell Death Dis. 2014; 5(1):e984–e912.
- Andres AM, Hernandez G, Lee P, Huang C, Ratliff EP, Sin J, et al. Mitophagy is required for acute cardioprotection by simvastatin. Antioxid Redox Signal. 2014; 21(14):1960–1973. <u>https://doi.org/10.1089/ars.2013.5416 PMID: 23901824</u>
- Neufeld EB, Remaley AT, Demosky SJ, Stonik JA, Cooney AM, Comly M, et al. Cellular Localization and Trafficking of the Human ABCA1 Transporter. J Biol Chem. 2001; 276(29):27584–27590. https://doi.org/10.1074/jbc.M103264200 PMID: 11349133
- Neumuller RA, Craig-Muller S, Markstein M, Perrimon N, Dettorre S, Cho J. Systematic screen of chemotherapeutics in Drosophila stem cell tumors. Proc Natl Acad Sci U S A. 2014; 111(12):4530–4535. https://doi.org/10.1073/pnas.1401160111 PMID: 24616500
- Arun B, Akar U, Gutierrez-Barrera AM, Hortobagyi GN, Ozpolat B. The PARP inhibitor AZD2281 (Olaparib) induces autophagy/mitophagy in BRCA1 and BRCA2 mutant breast cancer cells. Int J Oncol. 2015; 47(1):262–268. https://doi.org/10.3892/ijo.2015.3003 PMID: 25975349
- Zhang J, Chung TDY, Oldenburg KR. A Simple Statistical Parameter for Use in Evaluation and Validation of High Throughput Screening Assays. J Biomol Screen. 1999. <u>https://doi.org/10.1177/</u> 108705719900400206 PMID: 10838414
- Hasson SA, Fogel AI, Wang C, MacArthur R, Guha R, Heman-Ackah S, et al. Chemogenomic Profiling of Endogenous PARK2 Expression Using a Genome-Edited Coincidence Reporter. ACS Chem Biol. 2015; 10(5):1188–1197. https://doi.org/10.1021/cb5010417 PMID: 25689131
- Montava-Garriga L, Singh F, Ball G, Ganley IG. Semi-automated quantitation of mitophagy in cells and tissues. Mech Ageing Dev [Internet]. 2020; 185(December 2019):111196. <u>https://doi.org/10.1016/j.</u> mad.2019.111196 PMID: 31843465