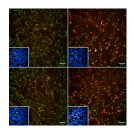


May 17, 2024 Version 2

KAT8 compound inhibition inhibits the initial steps of PINK1-dependent mitophagy V.2

DOI

dx.doi.org/10.17504/protocols.io.eq2ly7m8qlx9/v2



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OPEN ACCESS



DOI: dx.doi.org/10.17504/protocols.io.eq2ly7m8qlx9/v2

Protocol Citation: Capucine de Talhouet, Benjamin O'Callaghan, Noemi Esteras Gallego, Helene Plun-Favreau 2024. KAT8 compound inhibition inhibits the initial steps of PINK1-dependant mitophagy. protocols.io https://dx.doi.org/10.17504/protocols.io.eq2ly7m8qlx9/v2 Version created by Capucine de Talhouet

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Protocol status: Working
We use this protocol and it's

working

Created: May 30, 2023

Last Modified: May 17, 2024

Protocol Integer ID: 100051

Keywords: KAT8, MG149, Parkinson's disease, mitophagy, autophagy, TMRM, mitoKeima

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Funders Acknowledgement: The Michael J. Fox Foundation for Parkinson's Research (MJFF) and the Aligning Science Across Parkinson's (ASAP) initiative. Grant ID: ASAP-000478

Abstract

It has recently been shown that KAT8, a genome-wide association study candidate risk gene for Parkinson's Disease, is involved in PINK1/Parkin-dependant mitophagy. The KAT8 gene encodes a lysine acetyltransferase and represents the catalytically active subunit of the non-specific lethal epigenetic remodelling complex. In the current study, we show that contrary to KAT5 inhibition, dual inhibition of KAT5 and KAT8 via the MG149 compound inhibits the initial steps of the PINK1-dependant mitophagy process. More specifically, our study shows that following mitochondrial depolarisation induced by mitochondrial toxins, MG149 treatment inhibits PINK1-dependant mitophagy initiation by impairing PINK1 activation, and subsequent phosphorylation of Parkin and ubiquitin. While this inhibitory effect of MG149 on PINK1-activation is potent, MG149 treatment in the absence of mitochondrial toxins is sufficient to depolarise the mitochondrial membrane, recruit PINK1 and promote partial downstream recruitment of the autophagy receptor p62, leading to an increase in mitochondrial delivery to the lysosomes. Altogether, our study provides additional support for KAT8 as a regulator of mitophagy and autophagy processes.

Materials

Mitochondrial toxins

Oligomycin (inhibitor of mitochondrial complex V) was purchased from Cayman Chemicals (11341) and from Sigma-Aldrich (O4876) Antimycin A (inhibitor of mitochondrial complex III) was purchased from Sigma-Aldrich (A8674)

Antibodies used for immunoblotting

rabbit anti-phospho-ubiquitin (Ser65) (Cell Signaling, 37642, 1:1000)

mouse anti-TIM23 (BD Biosciences, 611223, RRID:AB_398755, 1:1000)

mouse anti-TOM20 (Santa Cruz, sc-11415, RRID:AB_2207533,1:1000)

mouse anti-GAPDH (Abcam, ab110305, RRID:AB_10861081, 1:1000)

rabbit anti-pParkin(Ser65) (Abcam/Michael J.Fox Foundation, MJF17, 1:1000)

rabbit anti-MFN2 (Cell Signaling, 9482, 1:1000)

mouse anti-OPA1 (BD Biosciences, 612607, 1:1000)

rabbit anti-LC3B (Novus Biologicals, NB100-2220, 1:1000)

IRDye 680LT donkey anti mouse (LI-COR Biosciences, 925-68022, RRID:AB_2814906, 1:20000)

IRDye 800CW donkey anti rabbit (LI-COR Biosciences, 925-32213, RRID:AB_2715510, 1:20000)

rabbit monoclonal anti-PINK1 antibody generated as described in the following manuscript: Sci Rep. 2015. DOI: https://doi.org/10.1038/s41598-018-26949-6

Antibodies used for immunofluorescence

mouse anti TOM20 (Santa Cruz, sc-17764, RRID:AB_628381 1:1000)

rabbit anti phospho-ubiquitin (Ser65) (Cell Signaling, 37642, 1:1000)

mouse anti-p62 (BD Biosciences, 610832, 1:1000)

AlexaFluor 488 goat anti rabbit (Invitrogen, A11008, 6 RRID:AB_143165,1:2000)

AlexaFluor 568 goat anti mouse (Invitrogen, A11004, 7 RRID:AB_2534072, 1:2000)



Cell Culture

Maintain cells in culture in a humified 5% CO₂ incubator at 37 °C in Dulbecco's modified Eagle's medium (DMEM, Gibco, 11995-065) supplemented with 10% heatinactivated foetal bovine serum (FBS, Gibco). Change cell medium every 3 days.

Note

Parkin over-expressing (POE) SHSY5Y cells are a gift from H. Ardley (Ardley, H., et al., 2003).

DOI: https://doi.org/10.1091/mbc.e03-02-0078

Mt-Keima POE SHSY5Y cells were a gift of C. Luft (Soutar, M., et al., 2019).

DOI: https://doi.org/10.1080/15548627.2019.1603549

PINK1-HA overexpressing SHSY5Y cells were a gift from E. Deas (Deas, E.., et al., 2011).

DOI: https://doi.org/10.1093/hmg/ddq526

2 Split cells at 75% to 90% confluency using 4 2 mL of Trypsin.

Cell Treatment (Small Molecule Modulator and Mitophagy Induction)

Dilute each compound in DMSO to reach concentrations determined according to their speculated toxicity and IC $_{50}$, [M] 10 micromolar (μ M) for NU9056 and [M] 100 micromolar (μ M) for MG149.

Note

MG149 targets KAT5 and KAT8 with IC $_{50}$ values of [M] 74 micromolar (μ M) and [M] 47 micromolar (μ M) respectively; NU9056 targets KAT5 with an IC $_{50}$ value of [M] 2 micromolar (μ M) (Li, L., *et al.*, 2020).

DOI: https://doi.org/10.1172/JCI131145

- 4 Treat cells with DMSO, [M] 10 micromolar (μM) NU9056, or [M] 100 micromolar (μM) 6h MG149 for 06:00:00 before collecting / fixing for downstream experiments.
- Treat cells with either DMSO or [M] 10 micromolar (μM) oligomycin/

 [M] 10 micromolar (μM) antimycin for 03:00:00 to induce mitophagy before collecting / fixing for downstream experiments.



Note that both small molecule modulator and O/A are often added to the same wells / cells to look at the impact of MG149 on mitophagy initiation.

Mitochondrial Enrichment

2h 10m

- 6 Incubate cells Overnight at \$\mathbb{8} -80 \circ O in mitochondrial homogenization buffer (250mM sucrose, 1mM EDTA, 10mM Tris (pH 7.4) supplemented with protease and phosphatase inhibitors).
- 7 Thaw, scrape into 1.5mL centrifuged tubes, and triturate 20x, keeping samples 2 On ice
- 8 Freeze tubes at 4°-80 °C for 60 01:00:00 , then thaw 4° On ice . 1h
- 9 Remove cell debris / non-lysed cells by centrifugation 1500 rcf, 4°C, 00:20:00 . 20m
- 10 Centrifuge the supernatant 12500 x g, 4°C, 00:20:00 to collect the mitochondrial 20m enriched pellet (aka the mitochondrial fraction) and save the supernatant (aka the cytoplasmic fraction).
- 11 Centrifuge the cytoplasmic fraction 12500 x g, 4°C, 00:20:00 to remove membrane 30m contamination and wash the mitochondrial fraction twice in \$\to\$ 500 \(\mu\text{L}\) of fractionation buffer (12500 x g, 4°C, 00:10:00 x 2) to remove cytoplasmic contaminants.
- 12 Determine the relative ratios of sample differences for the mitochondrial enriched fraction using the protein concentration of the cytoplasmic fraction (BSA assay). Re-suspend the mitochondrial enriched pellet accordingly using 1x NuPAGE® LDS sample buffer supplemented with 10 mM dithiothreitol (DTT).
- 13 Sonicate the mitochondrial fraction (10 microns power amplitude for 2 x 10 seconds, "flick" in between) and heat to 4 70 °C for 6 00:10:00 .

Cell Lysis



10m

14 Lyse cells in whole-cell lysis buffer (50mM Tris (pH 7.4), 0.1mM EGTA, 1mM EDTA, 0.27M Sucrose, 1% triton-x-100, supplemented with protease and phosphatase inhibitors) If On ice and incubated (Overnight at I -80 °C .



- 15 Centrifuge the lysates 1600 x g, 4°C, 00:20:00 and protein quantify the supernatant 20m by BSA assay.
- 16 Make LDS preps with 10 mM dithiothreitol (DTT), and heat to 🖁 70 °C for 🚫 00:10:00 . 10m

Western Blotting



- 17 Separate the LDS preps (either after cell lysis or mitochondrial enrichment) by SDS-PAGE and transfer them to nitrocellulose membrane.
- 18 Incubate the membrane Overnight at 4 °C with primary antibodies diluted in 3% milk in PBS-Tween.
- 19 Wash and incubate the membrane for 60 01:00:00 at 8 Room temperature in secondary 1h antibodies diluted in PBS-Tween.
- 20 Detect the protein bands using Odyssey CLx LI-COR.

Immunofluorescence



- 21 Fix cells with 4% Paraformaldehyde (PFA, Sigma-Aldrich) for 00:20:00 at 20m
 - Room temperature .
- 22 Remove PFA solution, block and permeabilize the cells with a mix of 10% FBS, 0.25% Triton X-1h 100 in PBS for 60 01:00:00 at 8 Room temperature .
- 23 Immunostain the cells with pUb(Ser65) and TOM20 primary antibodies diluted in 10% FBS/PBS 2h for 👏 02:00:00 at 🖁 Room temperature .

Note

rabbit anti-pUb(Ser65) (Cell Signaling, 37642) and mouse anti-TOM20 (Santa Cruz, sc-17764) primary antibodies are both added at a 1:1000 dilution in 10% FBS/PBS.

24 Wash wells three times using PBS 1X and incubate with secondary antibodies for 1h 01:00:00 at \$\mathbb{E}\$ Room temperature , before washing again three times using PBS 1X.



AlexaFluor 568 anti-mouse, 488 anti-rabbit secondary antibodies, and Hoechst 33342 (Thermo Scientific) are all added at a 1:2000 dilution in 10% FBS/PBS.

Image cells on the Opera Phenix (Perkin Elmer), acquiring confocal z-stacks for multiple fields of view across 3 individual wekks per experimental condition, using the 40X water objective.

Note

Analyse images using Columbus 2.8 analysis system (Perkin Elmer) to measure the integrated intensity of pUb(Ser65) within the whole cell.

For visualisation purposes, select brightness and contrast settings on control and apply the same settings to all other conditions.

Measuring Mitochondrial Membrane Potential using TMRM

40m

- 26 Seed POE-SHSY4Y cells in a 96-well CellCarrier Ultra plate.
- Incubate live cells in [M] 25 nanomolar (nM) Tetramethylrhodamine, Methyl Ester, Perchlorate (TMRM) (Sigma Aldrich) diluted in cell media for 00:40:00 at 37 °C and 4% CO₂.
- Image cells on the Opera Phenix (PerkinElmer) at 37 °C and 4% CO₂, acquiring confocal z-stacks for multiple fields of view across 3 individual wells per experimental condition using the 40X water objective.

Note

Analyse images using the Columbus 2.8 analysis system (Perkin Elmer) to measure the TMRM signal intensity of maximum intensity projections.

Measuring Mitochondrial Delivery to Lysosomes using mitoKeima Reporter



- Seed mt-Keima expressing POE SHSY5Y cells in a 96-well CellCarrier Ultra plate and treat them according to desired conditions (MG149, NU9056, O/A as described in <u>5 go to step #3</u>).
- Before imaging, replace cell media with phenol-free DMEM + 10% FBS containing the treatments previously administered , and adding Hoechst 33342 (1:10000) to stain the nuclei.



Image cells immediately on the Opera Phenix (PerkinElmer) at 37 °C with 5% CO₂, acquiring multiple single-plane fields of view, using the 63X water objective.

Note

Analyse images in using the Columbus 2.8 analysis system (Perkin Elmer) to measure the mitophagy index.

To analyse:

- Identify cells using the Hoechst 33342 channel, before measuring the area of the cytoplasmic and lysosomal mitoKeima
- Calculate the mitophagy index = the ratio between the total area of lysosomal mitochondria and the total area of mitoKeima (cytoplasmic + lysosomal areas) per well.

RT-qPCR

- Extract total RNA from cells using the Monarch Total RNA Miniprep Kit (New England Bioscience) with inclusion of the optional on-column DNAse treatment.
- Reverse transcribe RNA with SuperScript™ IV reverse transcriptase and random hexamers (Invitrogen).
- 34 Dilute the cDNA product then subject to quantitative real-time PCR (qPCR) using Power SYBR™ Green Master Mix (Applied Biosystems) and gene specific primer pairs on a QuantStudio™ 7 Flex Real-Time PCR System (Applied Biosystems).

	A	В	С	D
Γ		Sequence 5'-3'		
	Gene Target	Forward Reverse		Product Size / bp
	RPL18A	CCCACAACATGTACCGGGAA	TCTTGGAGTCGTGGAACTGC	180
	PINK1	GTGGAACATCTCGGCAGGTT	CCTCTCTTGGATTTTCTGTAAGTGAC	129

List of primer pairs used for RT-qPCR of target genes

Calculate relative mRNA expression levels using the $2^{-\Delta\Delta Ct}$ method and *RPL18A* as the housekeeping gene.

Statistical Analysis



- 36 Use a minimum of 3 biological replicates for each experiments (N numbers shown in legends refer to the number of replicate experiments).
 - For each imaging experiments, calculate the mean values of every condition from a minimum of 3 technical replicates.
 - Normalise intensity and integrated density measurements to control conditions.
- 37 Subject data to a two-way ANOVA with Dunnett's post-hoc analysis for multiple comparisons unless stated otherwise.
 - Express data as the mean ± standard deviation (SD) from replicate experiments.
 - Set statistical significance at a p-value of <0.05.

Use GraphPad Prism 6 (La Jolla, California, USA) for statistical analyses and graph production.

(Present data with error bars showing mean ± standard deviation (SD) from replicate experiments).

Measuring p62 recruitment to the OMM and mitochondrial delivery to lysosomes using mitoSRAI

3h

- Seed mitoSRAI POE SHSY5Y cells in a 96-well CellCarrier Ultra plate and treat them according to desired conditions (MG149, NU9056, O/A as described in <u>5 go to step #3</u>).
- Fix cells as described in go to step #21 to 22
- Immunostain the cells with p62 primary antibodies diluted in 10% FBS/PBS for 02:00:00

at Boom temperature.

Note

mouse anti-SQSTM1/p62 (Abcam, ab56416) primary antibody added at a 1:1000 dilution in 10% FBS/PBS.

Wash wells three times using PBS 1X and incubate with secondary antibodies for 01:00:00 at Room temperature, before washing again three times using PBS 1X.

1h

Note

AlexaFluor 568 anti-mouse secondary antibodies, and draq5 (Thermo Scientific) are all added at a 1:2000 dilution in 10% FBS/PBS.



Image cells on the Opera Phenix (Perkin Elmer), acquiring confocal z-stacks for multiple fields of view across 3 individual wells per experimental condition, using the 63X water objective.

Note

Analyse images using Columbus 2.8 analysis system (Perkin Elmer) to measure the SRAI mitophagy index

- corresponds to the ratio between the area of lysosomal mitochondria (corresponding to the total YPet negative TOLLES area) and the total mitochondrial area (corresponding to the total TOLLES area) per well.

Analyse images using Columbus 2.8 analysis system (Perkin Elmer) to measure p62 recruitment to teh mitochondria using TOLLEs as a mitochondrial marker

- calculated as the ratio of p62 signal intensity inside vs. outside TOLLES

For visualisation purposes, select brightness and contrast settings on control and apply the same settings to all other conditions.

Generating PINK1 KO POE SH-SY5Y cell line

PINK1 was functionally knocked out from the POE SH-SY5Y cell line through CRISPR-Cas9 editing by adapting previously published protocols (DOI: 10.15252/emmm.202013579)

Note

Single guide RNA (sgRNA) sequences targeting Exon 1 of PINK1 (NM_032409) were designed using the Horizon Discovery CRISPR Design Tool (https://horizondiscovery.com/en/ordering-and-calculation-tools/crispr-design-tool)

A	В	С	D
SgRNA ID	5'-3' sgRNA Sequence (PAM)	5'-3' Forward ssDNA Oligo with BbSI Overhang	5'-3' Reverse ssDNA Oligo with BbSI Overhang
PINK1_Seq1	CGCCACCATGGCGGTGCGAC(AGG)	caccgCGCCACCATGGCGGTGCGAC	aaacGTCGCACCGCCATGGTGGCGc
PINK1_Seq2	ACCGGGCGCGGAGCCTCGCA(GGG)	caccgACCGGGCGCGGAGCCTCGCA	aaacTGCGAGGCTCCGCGCCCGGTc

5'-3' Forward and Reverse ssDNA sequences generated (Sigma Aldrich) inclusive of 5' overhangs suitable for cloning into BbSI digested pSpCas9(BB)-2A-GFP plasmid



Materials:

pSpCas9(BB)-2A-GFP Addgene plasmid no. 48138 was a gift from Feng Zhang; (RRID:Addgene_48138)
IRDye 800CW donkey anti-rabbit secondary (LI-COR Biosciences, 925-32213, RRID:AB_2715510, 1:2000)
pUb(Ser65) (CST, 62802, RRID:AB_2799632, 1:1000)

- A subconfluent 6-well dish of POE SH-SY5Ys were transfected with 1ug of each PINK1_Seq1 and PINK1_Seq2 sgRNA encoding pSpCas9(BB)-2A-GFP plasmid using lipofectamine 3000 (Invitrogen)
- 45 24 h post-transfection cells were trypsinised and subjected to fluorescence-activated cell sorting and clonal selection of GFP-positive cells: isolated clones were expanded and phenotypically screened for clones which did not show pUb(Ser65) signal after 1.5 h O/A treatment
- Confluent wells of a 96-well were prepared for immunofluorescence staining of pUb(Ser65) with IRDye 800CW donkey anti-rabbit secondary
- 47 gDNA was extracted from a positive clone using the Wizard Genomic DNA Purification Kit (Promega) and subjected to whole genome sequencing (Novogene) which revealed homozygous 22 bp deletion and frameshift in exon 1 of PINK1 (c.156_177delGGGCGCGGAGCCTCGCAGGGTC, p.A54Sfs*46).

Generating mitoSRAI POE SH-SY5Y cell line

- 48 MitoSRAI cDNA was first PCR amplified from mitoSRAI_pcDNA3 with inclusion of 5' EcoRI and 3' NotI restriction sites
- The PCR product was cloned into pLVX-EF1α-IRES-Puro using EcoRI and NotI restriction sites to generate pLVX-EF1α-mitoSRAI-IRES-Puro.

Note

Material:

mitoSRAI_pcDNA3: provided by the RIKEN BRC through the National BioResource Project of the MEXT, Japan; cat. RDB18223 pLVX-EF1α-IRES-Puro: Clontech, Takara Bio, 631988 pMD2.G (Addgene plasmid no. 12259, RRID:Addgene_12259) and pCMVR8.74 (Addgene plasmid #22036, RRID:Addgene_22036) were gifts from Didier Trono

50 Lenti-X 293 T HEK cells cultured in DMEM 10% FBS media were transfected with pMD2.G, pCMVR8.74 and pLVX-EF1α-mitoSRAI-IRES-Puro at a 1:1:2 molar mass ratio using Lipofectamine 3000 (Invitrogen)



- The next day, a full media change was performed using DMEM 10% FBS and cells cultured for further 24
- The lentivirus containing media was collected and diluted 1:2 with DMEM 10% FBS before filtering through 0.44 µm PES filters
- 1×106 POE SHSY5Ys were reverse transduced with the 1:2 lentivirus supernatant dilution in the presence of $10 \,\mu\text{g/mL}$ polybrene (Sigma Aldrich)
- Bulk populations of cells stably expressing mitoSRAI were established through selection with 1 ug/mL puromycin (M P BIOMEDICALS UK)

 Puromycin was maintained during routine culture but withdrawn when seeding for experimental assays.

Protocol references

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