#### PARK15/FBXO7 is dispensable for PINK1/Parkin mitophagy in iNeurons and HeLa cell systems

Felix Kraus<sup>1,2</sup>, Ellen A. Goodall<sup>1,2</sup>, Ian R. Smith<sup>1</sup>, Yizhi Jiang<sup>1</sup>, Julia C. Paoli<sup>1</sup>, Jiuchun Zhang<sup>1</sup>, Joao A. Paulo<sup>1</sup>, and J. Wade Harper<sup>1,2\*</sup> 

<sup>1</sup>Department of Cell Biology, Blavatnik Institute, Harvard Medical School, 240 Longwood Ave, Boston MA 02115, USA

<sup>2</sup>Aligning Science Across Parkinson's (ASAP) Collaborative Research Network, Chevy

Chase, MD 20815, USA

\*Correspondence: wade harper@hms.harvard.edu (J.W.H.) 

#### 18 ABSTRACT

19 The protein kinase PINK1 and ubiquitin ligase Parkin promote removal of damaged 20 mitochondria via a feed-forward mechanism involving ubiquitin (Ub) phosphorylation, Parkin 21 activation, and ubiquitylation of mitochondrial outer membrane proteins to support 22 recruitment of mitophagy receptors. The ubiquitin ligase substrate receptor FBXO7/PARK15 23 is mutated in an early-onset parkinsonian-pyramidal syndrome. Previous studies have 24 proposed a role for FBXO7 in promoting Parkin-dependent mitophagy. Here, we 25 systematically examine the involvement of FBXO7 in depolarization-dependent mitophagy in 26 the well-established HeLa and induced-neurons cell systems. We find that FBXO7<sup>-/-</sup> cells 27 have no demonstrable defect in: 1) kinetics of pUb accumulation, 2) pUb puncta on 28 mitochondria by super-resolution imaging, 3) recruitment of Parkin and autophagy 29 machinery to damaged mitochondria, 4) mitophagic flux, and 5) mitochondrial clearance as 30 quantified by global proteomics. Moreover, global proteomics of neurogenesis in the 31 absence of FBXO7 reveals no obvious alterations in mitochondria or other organelles. These 32 results argue against a general role for FBXO7 in Parkin-dependent mitophagy and point to 33 the need for additional studies to define how FBXO7 mutations promote parkinsonian-34 pyramidal syndrome.

- 35
- 36

#### 37 INTRODUCTION

38 Organelle quality control underlies cellular health and is often defective in disease 39 and pathological states. Arguably the best understood such quality control pathway is 40 mitophagy, wherein damaged or supernumerary mitochondria are targeted for removal from 41 the cell via selective autophagy (Goodall et al, 2022; Harper et al, 2018; McWilliams & Mugit, 42 2017; Pickrell & Youle, 2015). In this process, individual organelles are marked for 43 sequestration within a double-membrane vesicle called an autophagosome, which then 44 fuses with a lysosome to facilitate degradation of the organelle by resident lysosomal 45 hydrolases (Dikic & Elazar, 2018; Stolz *et al*, 2014). Multiple forms of mitophagy have been 46 reported, which differ in the types of regulatory mechanisms involved in marking the 47 organelle for degradation, but these fall into two primary types - ubiquitin-dependent and 48 ubiquitin-independent (Harper et al., 2018; Pickrell & Youle, 2015). Our understanding of 49 ubiquitin-dependent mitophagy has been advanced by the discovery of a signalling pathway 50 composed of the PINK1 protein kinase and the Parkin (also called PRKN) ubiquitin (Ub) E3 51 ligase, which marks damaged mitochondria for elimination (Narendra et al, 2008). Parkin 52 and PINK1 are both mutated in early-onset recessive forms of Parkinson's disease and 53 understanding how these enzymes work has been a major focus of the field (Ng et al, 2021; 54 Pickrell & Youle, 2015).

55 In healthy mitochondria, PINK1 is imported into the mitochondrial translocon and 56 rapidly processed for degradation (Jin et al, 2010; Yamano & Youle, 2013). In response to 57 mitochondrial damage - such as depolarization, accumulation of mitochondrial misfolded 58 proteins, or defects in mitochondrial fusion – PINK1 is stabilized on the mitochondrial outer 59 membrane (MOM) in association with the translocon (Lazarou et al, 2012), where it can 60 phosphorylate Ser65 on Ub already conjugated to proteins on the MOM (Kane et al, 2014; 61 Kazlauskaite et al, 2015; Koyano et al, 2014; Ordureau et al, 2014; Wauer et al, 2015). 62 Accumulation of phospho S65-Ub (referred to henceforth as pUb) on the MOM promotes 63 recruitment of Parkin – a pUb-binding protein – thereby facilitating phosphorylation of Parkin 64 on S65 of its N-terminal Ub-like (UBL) domain and activation of its Ub ligase activity 65 (Gladkova et al, 2018; Kane et al., 2014; Kazlauskaite et al., 2015; Ordureau et al., 2014; 66 Sauve et al, 2018; Wauer et al., 2015). Parkin then ubiquitylates numerous MOM proteins, 67 resulting in both the accumulation of additional pUb and further MOM protein ubiquitylation 68 (Antico et al, 2021; Bingol et al, 2014; Ordureau et al, 2020; Sarraf et al, 2013). MOM 69 ubiquitylation then promotes recruitment of selective autophagy cargo receptors including 70 OPTN, CALCOCO2 (also called NDP52), and SQSTM1 (also called p62) to facilitate the 71 assembly of an autophagosome around the ubiquitylated organelle (Evans & Holzbaur, 72 2020; Heo et al, 2015; Lazarou et al, 2015; Wong & Holzbaur, 2014). Thus, PINK1, Parkin, 73 and pUb function in a positive feedback loop to promote selective ubiquitylation and 74 elimination of the damaged organelle (Goodall et al., 2022).

75 Previous work (Burchell et al, 2013) concluded that FBXO7, the product of an early-76 onset parkinsonian-pyramidal syndrome risk gene also designated as PARK15 (Di Fonzo et 77 al, 2009; Houlden & Singleton, 2012; Paisan-Ruiz et al, 2010), is a positive modulator of 78 Parkin-dependent mitophagy (**Supplemental Figure 1A**). FBXO7 is a member of the F-box 79 family of proteins, which assemble with SKP1, CUL1, and RBX1 to form a modular SCF Ub 80 ligase complex wherein the F-box protein to binds substrates (Jin et al, 2004). FBXO7 is 81 characterized by an N-terminal UBL domain, a central FP domain with structural similarity to 82 a domain in the proteasome inhibitory factor PI31/PSMF1, followed by the F-box motif, and 83 a C-terminal proline rich region (Kirk et al, 2008). Patient mutations in FBXO7 are found in 84 multiple regions of the protein (Di Fonzo et al., 2009; Houlden & Singleton, 2012; Paisan-85 Ruiz et al., 2010). Initial studies on FBXO7 proposed a functional interaction with the 86 Parkin/PINK1 pathway and concluded that FBXO7 functions as a positive regulator of 87 mitophagy through direct interaction with Parkin and with PINK1 (Burchell et al., 2013). 88 Depletion of FBXO7 by siRNA in cell lines was reported to reduce depolarization-dependent 89 loss in mitochondria based on immunoblotting of matrix proteins and likewise to reduce 90 depolarization-dependent ubiquitylation of MFN1, a substrate of Parkin (Burchell et al., 91 2013). However, studies that further substantiate a functional link between FBXO7 and

mitophagy are lacking, although FBXO7 depletion has recently been reported to alter
 mitochondrial dynamics (Al Rawi *et al*, 2022).

94 Given the strong genetic association of FBOX7/PARK15 with Parkinson's Disease 95 (Houlden & Singleton, 2012) and major advances in our understanding of Parkin-dependent 96 mitophagy (Goodall et al., 2022) since initial links between FBXO7 and mitophagy were 97 reported, we set out to elucidate how FBXO7 might function within the positive feedback 98 loop to promote mitophagy (Supplemental Figure 1A). Relevant advances include the 99 identification of pUb as a sentinel response to PINK1/Parkin activation (Kane et al., 2014; 100 Kazlauskaite et al., 2015; Koyano et al., 2014; Ordureau et al., 2014; Wauer et al., 2015), 101 the development of highly quantitative mitophagic flux assays using a mitochondrially 102 localized Keima fluorescent reporter protein (Katayama et al, 2011; Lazarou et al., 2015), 103 and the development of genetically tractable iNeuron systems for functional analysis of the 104 Parkin pathway in neuronal cell types (Ordureau et al., 2020; Ordureau et al, 2018). 105 Unexpectedly, we find no evidence of a role for FBXO7 in the kinetics of depolarization-106 dependent pUb accumulation, or mitophagic flux in either iNeurons or the conventional HeLa 107 cell used extensively to examine Parkin and PINK1-dependent mitophagy (Narendra et al., 2008). Moreover, global proteomics of FBXO7<sup>-/-</sup> HeLa cells undergoing mitophagy revealed 108 109 no defect in mitochondrial elimination by autophagy. Taken together with previous studies 110 that failed to verify Parkin-FBXO7 interactions during mitophagy (Sarraf et al., 2013), these 111 data indicate that FBXO7 does not play a general role as a positive modulator of Parkin and 112 PINK1-dependent mitophagy in HeLa and iNeuron systems. This work suggests that 113 elucidation of FBXO7 biochemical functions is needed to understand how it might contribute 114 to suppression of early-onset parkinsonian-pyramidal syndrome.

115

#### 116 **RESULTS**

# 117 A tool kit for functional analysis of cells lacking FBX07

To test if FBXO7 is a pivotal amplifier of the PINK1/Parkin pathway, we employed 118 119 CRISPR-Cas9-based gene editing to generated knockouts for FBXO7 in HeLa HFT/TO-120 PRKN cells (Ordureau et al, 2015) and confirmed insertion of a frameshift mutation by 121 sequencing and immunoblotting (Supplemental Figure 1 B-D). HeLa HFT/TO-PRKN cells 122 contain a DOX-inducible PRKN cassette allowing regulated expression of Parkin to facilitate 123 an analysis of mitophagic flux in response to mitochondrial damage (Ordureau et al., 2015). 124 Three independent FBXO7<sup>-/-</sup> clones were identified (clones 27, 41 and 45) (Supplemental 125 Figure 1C,D). In parallel, we generated two independent human ES cell clones lacking 126 FBXO7 via CRISPR-Cas9 (clones 52 and 89) (Supplemental Figure 1B,E,F). Here, an ES 127 cell line harbouring a DOX inducible NGN2 gene was used, allowing for conversion of these 128 cells to cortical-like iNeurons over a two week or greater time course (Ordureau et al., 2020;

129 Zhang *et al*, 2013). We have previously demonstrated that iNeurons derived from these cells 130 exhibit Parkin and PINK1-dependent mitophagic flux in response to mitochondrial damage 131 (Ordureau *et al.*, 2020). Both clones displayed normal karyotypes and the absence of 132 FBXO7 was verified using MiSeq sequencing (**Supplemental Figure 1F,G**).

133

# Robust depolarization-dependent Ub phosphorylation by PINK1 in cells lacking FBX07

136 The earliest known event in Parkin-dependent mitophagy is accumulation of PINK1 137 on the mitochondrial outer membrane followed by rapid S65 phosphorylation of pre-existing 138 Ub linked with mitochondrial proteins (Supplemental Figure 1A), as has been worked out in 139 detail using the well-validated HeLa cell system with ectopically expressed Parkin (Kane et 140 al., 2014; Lazarou et al., 2012; Narendra et al, 2010; Yamano & Youle, 2013). In the 141 presence of Parkin, Ub phosphorylation is enhanced as a result of increased Ub conjugation 142 to mitochondrial proteins, thereby providing additional substrate molecules for 143 phosphorylation by PINK1 (Kane et al., 2014; Koyano et al., 2014; Ordureau et al., 2014). To 144 examine pUb accumulation in the context of HeLa cells lacking FBXO7, we induced Parkin 145 expression (16 h) and then depolarized mitochondria for 1 or 6 h with antimycin 146 A/Oligomycin A (AO), which inhibit Complex III and Complex IV activity in the electron 147 transport chain, respectively. As expected, control HeLa cells display time-dependent 148 accumulation of pUb as measured by immunoblotting, but the absence of FBXO7 had no impact on the extent of pUb accumulation, as assessed in all three FBXO7<sup>-/-</sup> cell lines in 149 150 biological triplicate analyses (Figure 1A,B).

Similarly, we assessed depolarization-dependent pUb accumulation in the iNeuron system. control or FBXO7<sup>-/-</sup> ES cells were converted to day 12 iNeurons and depolarized with AO in duplicate (**Figure 1C**). pUb was detected by immunoblotting at 6 and 24 h and was indistinguishable between WT and two independent FBXO7<sup>-/-</sup> clones (**Figure 1D**). Taken together, these data do not support a universal role for FBXO7 in promoting the earliest step in mitophagy signalling – PINK1-dependent Ub phosphorylation – in the HeLa cell system or iNeurons.

- 158
- 159

#### 9 Super-resolution imaging of pUb formation independent of FBX07

While depolarization-dependent Ub phosphorylation was not impaired, we considered the possibility that the organization of pUb on damaged mitochondria might be altered in cells lacking FBXO7. To our knowledge, super-resolution microscopy has not been used to examine the kinetics and spatial properties of pUb accumulation in response to depolarization. We therefore optimized conditions to monitor pUb via volumetric 3D-SIM super-resolution microscopy in both HeLa and iNeuron systems (see **EXPERIMENTAL** 

166 **MATERIALS AND METHODS**). After AO treatment of HeLa cells for 1, 3 or 6 h, the 167 mitochondria matrix protein HSP60 and pUb were examined by super-resolution imaging. In 168 control cells at 1 h, fragmented mitochondria were often proximal to pUb foci with an 169 average volume of 5.71 µm<sup>3</sup>, which were not observed in cells lacking PINK1 (Figure 2A-C, 170 Supplemental Figure 2A,B). The volume of pUb increased ~8-fold over 6 h (47.88 µm<sup>3</sup>). 171 consistent with the known feed-forward response (Figure 2B, Supplemental Figure 2A). At 172 6 h of AO treatment, we observed a near complete coating of mitochondrial fragments with 173 pUb, as quantified by the significant increase in pUb object volume (Figure 2B,C, 174 Supplemental Figure 2A). Consistent with analysis of pUb by immunoblotting, two FBXO7<sup>-/-</sup> 175 clones did not show any significant differences in pUb recruitment, volume or overall 176 morphological differences compared to control cells, indicating that FBXO7 is not required 177 for the feed-forward amplification of PINK1/Parkin mitophagy after AO-treatment (Figure 2A-178 **C**). Analogous super-resolution experiments in day 12 iNeurons with 0.5, 1, and 3 h of AO 179 revealed similar results, with no obvious effect of FBXO7 deletion on pUb recruitment, 180 volume or overall morphology when compared to WT iNeurons (Figure 2D-F, 181 Supplemental Figure 3A). As expected (Ordureau et al., 2020; Ordureau et al., 2018), 182 iNeurons lacking PINK1 did not accumulate pUb (Figure 2D,E). These results were also 183 confirmed in iNeurons treated with AO for 1 or 6 h using conventional immunofluorescence 184 and image analysis of HSP60 and pUb co-localization (Figure 2G,H). To ensure that kinetic 185 effects were not obscured by overt depolarization, we reduced the concentrations of AO 186 used by 10-fold, but again no defect in pUb recruitment to mitochondria was observed 187 (Supplemental Figure 3B,C). These data indicate that the absence of FBXO7 does not 188 alter the ability of PINK1-dependent pUb foci to accumulate on damaged mitochondria in 189 either HeLa or iNeuron systems.

- 190
- 191

# Parkin and autophagy regulator recruitment in FBXO7<sup>-/-</sup> cells

192 Parkin recruitment depends upon the accumulation of pUb on damaged mitochondria 193 (Supplemental Figure 1A) (Okatsu et al, 2015). Given that the existing model for FBXO7 in 194 Parkin-dependent mitophagy posited that Parkin and FBXO7 physically interact and that 195 FBXO7 promotes Parkin function (Burchell et al., 2013), we directly tested whether FBXO7 196 is required for Parkin recruitment to mitochondria using the HeLa cell system. Control or 197 FBXO7<sup>-/-</sup> cells were stably transduced with a lentivirus expressing GFP-Parkin and subjected 198 to imaging with or without depolarization with AO (Supplemental Figure 4A). We used 199 CellProfiler (Stirling et al, 2021) to analyse the characteristics of GFP-Parkin and 200 mitochondria and their relationship to each other. By segmenting out cytosolic and 201 mitochondrially-localized Parkin, we could quantify alterations in Parkin abundance within 202 the mitochondrial mask as a measure of Parkin-translocation (Supplemental Figure 4B).

203 Our expectation was that a requirement for FBXO7 in Parkin recruitment would result in 204 reduced Parkin intensity within the mitochondrial mask in response to depolarization. First, 205 we tested whether mitochondrial morphology (mitochondrial integrated density [==mtIntDen] 206 as proxy, stained with HSP60) is changing after AO-treatment on a single-cell level. Under fed conditions, control and three FBXO7<sup>-/-</sup> clones displayed comparable mtIntDen. 207 208 Depolarization by AO led to the expected mitochondrial fragmentation and clustering, thus 209 causing an increase in local HSP60 signal and a right-shift in mtIntDen (Supplemental 210 Figure 4C, top panel). This shift was also observed to a similar extend in FBXO7<sup>-/-</sup> cells. 211 Thus, both control and FBXO7<sup>-/-</sup> cells robustly respond to AO-induced mitophagy. Next, we 212 tested if Parkin intensities are increasing on mitochondria on an "organelle object" level. As 213 expected, mtIntDen did shift to the right after AO-treatment, in line with previous 214 experiments. Parkin signal was comparable between the genotypes under fed conditions, 215 and increased in intensity after AO-treatment, consistent with Parkin accumulation inside the 216 mitochondrial area (Supplemental Figure 4C, lower panel). This Parkin shift was observed in both control and FBXO7<sup>-/-</sup> cell lines, arguing that FBXO7 is not universally required for 217 218 Parkin recruitment to damaged mitochondria.

219 Mitochondrial ubiquitylation by Parkin promotes recruitment of downstream 220 autophagy machinery, including the FIP200-ULK1 complex and Ub-bind autophagy 221 adaptors, ultimately leading to the formation of an autophagosome (Heo et al., 2015; 222 Lazarou et al., 2015; Ravenhill et al, 2019; Vargas et al, 2019). Lipidation of LC3 is 223 associated with autophagosome formation and is thought to contribute to recruitment of 224 autophagy receptors. As expected, depolarization of Parkin-expressing HeLa cells resulted 225 in recruitment of both p62 and FIP200 to mitochondria in a manner that dependent on 226 PINK1, as revealed by immunostaining and confocal imaging (Supplemental Figure 4D). 227 However, FBXO7<sup>-/-</sup> cells were as proficient as control cells in recruitment of p62 or FIP200 228 after 3 h or 16 h or AO-induced mitophagy (Supplemental Figure 4D-G). Previous studies 229 using siRNAs targeting FBXO7 also reported that LC3 lipidation in response to mitochondrial 230 depolarization is defective compared with non-depleted cells (Burchell et al., 2013). 231 However, all three FBXO7<sup>-/-</sup> HeLa cell clones underwent depolarization-dependent LC3 232 lipidation to similar extents, consistent with a functional Parkin pathway (Supplemental 233 Figure 4H,I). Taken together, these data indicate that FBXO7 is not universally required to 234 promote either Parkin-dependent recruitment to damaged mitochondria or subsequent steps 235 that depend upon Ub chain assembly on mitochondria such as recruitment of autophagy 236 machinery and initiation of LC3 lipidation.

Additionally, we have not able to reproduce a previously reported association between overexpressed FBXO7 and PINK1 (Huang *et al*, 2020) in either HEK293T or HCT116 cells. Briefly, as part of our BioPlex project (Huttlin *et al*, 2021), we ectopically

240 expressed PINK1-HA and performed interaction proteomics. While we identified the known 241 protein kinase chaperone HSP90 subunits in association with PINK1, we did not detect 242 FBXO7 (Supplemental Figure 4J, left panel; Supplemental Table S1). Likewise, FBXO7 243 associated reciprocally with members of the CUL1-SKP1-RBX1, the COP9/Signalosome, 244 and both core particle and regulatory subunits of the proteasome in HEK293T and/or 245 HCT116 cells, as expected (Bader et al, 2011; Liu et al, 2019; Vingill et al, 2016), but neither 246 PINK or Parkin were detected (Supplemental Figure 4J,K; Supplemental Table S2) 247 (Huttlin et al., 2021) (see METHODS). These results are consistent with our previous Parkin 248 interaction proteomics experiments that failed to demonstrate an interaction with FBXO7 249 during mitophagy induction (Sarraf et al., 2013).

250

# 251 Mitophagic flux in FBXO7<sup>-/-</sup> cells

252 A significant advance has been the development of mitophagy reporters, which 253 cumulatively measure flux of mitochondrial turnover through to the final stages of mitophagy: 254 the fusion of mitochondria-laden autophagosomes with lysosomes (Lazarou et al., 2015). 255 mKeima is a fluorescent protein that undergoes a pH-dependent Stokes shift, and delivery of 256 a Keima-tagged cargo from the cytoplasm to the acidic interior of the lysosome can be 257 monitored by measuring the ratio of emission at 620 nm with maximal excitation at 440 nm 258 or 586 nm under neutral or acidic conditions, respectively, by flow cytometry or microscopy 259 (Figure 3A). To monitor differences in mitophagic flux upon deletion of FBXO7, we 260 employed mtKeima, an mKeima protein targeted to the mitochondrial with a COX8-matrix 261 targeting sequence (Ordureau et al., 2020).

262 In Parkin-expressing HeLa cells, the ratio of acidic-Keima (measured with excitation 263 at 561 nm and emission 620 nm) to neutral Keima (ex. 405 nm, em. 603 nm) increases as 264 cells are treated with AO (Figure 3B, Supplemental Figure 4L). This mitophagic flux is 265 PINK1-dependent as deletion of PINK1 abolishes this shift (Figure 3B, Supplemental 266 Figure 4L). As expected, there is little flux without depolarization with AO, and inhibition of 267 the lysosomal V-ATPase with BafilomycinA1 (BafA) restores the mtKeima acidic:neutral ratio 268 to levels similar to that seen fed cells (**Supplemental Figure 4L**). In three FBXO7<sup>-/-</sup> clones, 269 AO-dependent mitophagic flux is similar to control cells with increasing flux observed with 270 increasing length of AO treatment (Figures 3B; Supplemental Figure 4L). We validated the 271 mtKeima flux results using the mitochondrial DNA clearance assay (Heo et al., 2015; 272 Lazarou et al., 2015). Staining for mtDNA foci per cell after 24 h AO treatment in control and 273 FBXO7<sup>-/-</sup> HeLa cells showed a significant decrease in mtDNA number in all samples (Figure **3G**). The remaining mtDNA levels were indistinguishable between control and FBXO7<sup>-/-</sup> cells 274 275 (Figure 3G).

276 We also measured mtKeima foci and flux in control and FBXO7<sup>-/-</sup> iNeurons at day 12 277 of differentiation (Figure 3C-E). As with HeLa cells, red-shifted mtKeima foci increased ~2-278 fold after 6 h AO treatment in control and FBXO7<sup>-/-</sup> cells (Figure 3C,D). Using flow 279 cytometry, mitophagic flux (normalized to BafA) after 6 and 24 h AO treatment was comparable between control and FBXO7<sup>-/-</sup> iNeurons, and was absent in PINK1<sup>-/-</sup> cells 280 281 (Figure 3E). Additionally, we investigated mtDNA turnover in d12 control and two 282 independent FBXO7<sup>-/-</sup> iNeuron clones. Within 6 or 24 h of depolarization of iNeurons, we 283 observed a reduction in the number of mitochondrial DNA/cell, and the effect was 284 independent of the presence or absence of FBXO7 (Figure 3F,G). Taken together, these 285 data indicate that FBXO7 is not universally required for depolarization-dependent mitophagic 286 flux in either the HeLa or iNeuron systems.

287

288

#### Proteomic analysis of HeLa cells lacking FBXO7 during mitophagy

289 As an alternative to mtKeima for measuring mitophagic flux, we examined the total 290 proteome of depolarized HeLa cells with the expectation that in control HeLa cells, 291 mitochondrial turnover would result in bulk loss of the mitochondrial proteome in depolarized 292 cells. Thus, control or FBXO7<sup>-/-</sup> HeLa cells expressing Parkin were depolarized for 16 h in 293 triplicate and total cell extracts subjected to 18-plex Tandem Mass Tagging (TMT)-based 294 proteomics (Figure 4A, Supplemental Table S3,S4) (Li et al, 2021). Replicates were highly 295 correlated, with correlation coefficients greater that 0.95 and Principal Component Analysis 296 (PCA) revealed tight clustering of replicates, with the major feature separating the samples 297 being AO-treatment (Supplemental Figure 5A,B). Through hierarchical clustering 298 (Supplemental Figure 5C), we identified major alterations in the abundance of mitochondria 299 in response to depolarization (Figure 4B, Supplemental Figure 5D-I). Indeed, the majority 300 of proteins annotated as mitochondrial were found to have reduced levels in control cells in 301 response to depolarization, as indicated in the overlap of mitochondrial proteins on the 302 volcano plots for the ~8000 proteins quantified (leftward skew of red dots in Figure 4B, left 303 panel). Importantly, the patterns of protein abundance were indistinguishable from control for 304 two independent FBXO7<sup>-/-</sup> cell lines, with leftward skew of mitochondrial proteins in the 305 volcano plot (Figure 4B,C). Proteins with altered abundance were enriched for mitochondrial 306 matrix, inner membrane, outer membrane and oxidative phosphorylation categories, 307 consistent with autophagy of entire organelles (Figure 4C). The behaviour of other 308 organelles and large protein complexes were similar for both control and FBXO7<sup>-/-</sup> cells 309 (Figure 4C). Moreover, both FBXO7 clones displayed similar behaviour to each other and to 310 the control, as indicated by correlation analysis (Pearson's R=0.82, 0.86 and 0.74, 311 respectively) (Figure 4D,E). These results are consistent with the absence of a discernible 312 alteration in depolarization-dependent mitophagic flux in cells lacking FBXO7. A previous

report proposed that depletion of FBXO7 resulted in accumulation of PINK1 proteins levels under basal conditions (Huang *et al.*, 2020), but in our experiments, PINK1 levels in the two FBXO7<sup>-/-</sup> clones were unaffected based on TMT intensities (**Figure 4F**).

- 316
- 317

#### Proteomic analysis of human ES cells lacking FBX07 during neurogenesis

318 As an initial unbiased approach to examining FBXO7 function in iNeurons, we 319 performed 18-plex TMT proteomics on control and two clones of FBXO7<sup>-/-</sup> cells at day 0, 4 320 and d12 of differentiation (Figure 5A, Supplemental Table S5-S8). We have previously 321 demonstrated dramatic remodelling of mitochondria around day 4 of differentiation to 322 support a switch from glycolysis to oxidative phosphorylation, which is accompanied by 323 BNIP3L-dependent mitophagy of a fraction of mitochondria (Ordureau et al, 2021). From the 324 ~6000 proteins quantified across all replicates and conditions, we found strong 325 correspondence of control and FBXO7<sup>-/-</sup> cells, with PC1 being drive by differentiation 326 (Supplemental Figure 6A,B). FBXO7 itself was expressed at comparable levels across the 327 time course but was not detected in FBXO7<sup>-/-</sup> iNeurons, as expected (Supplemental Figure 328 6C). Remarkably, the abundance of the proteome was largely immune to the loss of FBXO7 329 at both day 4 and day 12 of differentiation (Figure 5B-D, Supplemental Figure 6D). 330 Moreover, the abundance of mitochondrial proteins (including inner and out membrane and 331 OXPHOS components) were unchanged relative to control cells at day 4 or 12 of 332 differentiation (Figure 5B, Supplemental Figure 6D).

333 Recent reports have directly linked L250P mutations in the dimerization domain of 334 FBXO7 to MiD49/MiD51, a core adapter protein involved in mitochondrial fission (Al Rawi et 335 al., 2022). Indeed, DNM1L-mediated mitochondrial fission has been previously indicated in 336 the efficient scission of damaged parts of mitochondria and protection of the remaining healthy mitochondrial network (Burman et al, 2017). Cells expressing FBXO7<sup>L250P</sup> were 337 338 found to have reduced levels of proteasome subunits as well (Al Rawi et al., 2022). Our 339 proteomic data from iNeurons indicates no alterations in the abundance of proteasome 340 subunits or mitochondrial fission and fusion machinery (DNM1L, MFF, and MiD51) in 341 iNeurons lacking FBXO7 (Figure 5C,E). To test if loss of FBXO7 causes in morphological 342 changes in the mitochondrial network, we stained day 12 iNeurons with for the matrix marker 343 HSP60 prior to image analysis (Figure 5F,G). However, we observed no significant 344 alterations in the number of mitochondria/cell or the mean mitochondrial length in control or 345 FBXO7<sup>-/-</sup> iNeurons. The mean mitochondrial circularity was slightly reduced in FBXO7<sup>-/-</sup> 346 iNeurons treated with AO for 3h (Figure 5F). Analogous analyses of mitochondria and 347 fission/fusion proteins also did not reveal significant alterations in response to loss of FBXO7 348 in two independent clones of HeLa cells (Figure 5H,I). However, there was a ~10% 349 decrease in the abundance of proteasome subunits in HeLa cells lacking FBXO7 in the

350 absence of depolarization, but an increase after mitophagy induction (Supplemental Figure 351 **5E, Figure 4C**). If this response is a compensatory mechanism of the knockout cell line to 352 cope with cytotoxic insults is an intriguing idea. Last, we examined whether FBXO7<sup>-/-</sup> is required for the maintenance of the mitochondrial organelle pool during differentiation. 353 354 Previous work has shown significant metabolic rewiring during the differentiation from hESC 355 to iNeurons, including a switch from glycolysis to oxidative phosphorylation as the cells main 356 energy source (Ordureau et al., 2021). Since this switch is accompanied by increased 357 mitophagy, we set out to see if the absence of FBXO7 would lead to accumulation of 358 mitochondria and/or impair differentiation. FBXO7<sup>-/-</sup> cells did not display obvious differences 359 in the abundance of mitochondrial proteins, when normalized to day 0 (Figure 5B, 360 Supplemental Figure 6D). Likewise, other abundance of other organelles was unchanged 361 (Figure 5C). Finally, neither differentiation markers or autophagy proteins displayed obvious changes in control versus the two FBXO7<sup>-/-</sup> clones as measured by TMT-proteomics and 362 363 displayed using a  $T^2$ -statstic (Ordureau *et al.*, 2021) (**Supplemental Figure 7A-F**).

364

#### 365 **DISCUSSION**

366 Despite strong genetic evidence linking mutations in FBXO7 with parkinsonian-367 pyramidal syndrome (Di Fonzo et al., 2009; Houlden & Singleton, 2012; Paisan-Ruiz et al., 368 2010), our understanding of the cellular roles of FBXO7 are limited. An early study reported 369 biochemical links between FBXO7 and Parkin/PINK1-dependent mitophagy in fibroblasts 370 and SH-SY5Y cells, including interaction of overexpressed FBXO7 with both PINK1 and 371 Parkin (Burchell et al., 2013). Using siRNA to deplete FBXO7, it was also concluded that 372 FBXO7 promotes Parkin's ability to ubiquitylate the outer mitochondrial membrane substrate 373 MFN1 and also promotes clearance of mitochondria by autophagy. These biochemical and 374 physiological results led to the conclusion that FBXO7 functions as a biochemical amplifier 375 of Parkin/PINK1-deepndent mitophagy (Burchell et al., 2013). However, follow-up studies 376 have been limited, and the field has experienced dramatic advances in the understanding of 377 the biochemical roles of Parkin and PINK1 in promoting mitochondrial clearance by 378 autophagy, thereby providing the tools to examine potential roles of FBXO7 in the pathway.

379 With these tools, we set out to define where in the pathway FBXO7 might operate to 380 promote mitophagy. However, in both the HeLa cells with overexpressed Parkin and 381 iNeurons with a fully endogenous Parkin/PINK1 pathway, we failed to validate a role for 382 FBXO7 in any of several steps in the pathway, including pUb accumulation, Parkin 383 recruitment to the mitochondrial outer membrane, recruitment of autophagy machinery, 384 mitophagic flux as measured by mtKeima, and mitochondrial proteome degradation. The 385 finding that deletion of FBXO7 has no obvious effect on the pathway with either endogenous 386 or overexpressed Parkin in iNeurons and HeLa cells indicates that FBXO7 does not play a

387 general or universally required role in the pathway. A previous study largely based on 388 overexpression concluded that FBXO7 may bind and regulate PINK1 levels, with <2-fold 389 increases in PINK1 by immunoblotting in FBXO7-depleted cells (Huang et al., 2020). 390 However, our quantitative proteomics experiments (Figure 4) indicated that deletion of 391 FBXO7 has no effect on PINK1 abundance, consistent with the finding that pUb 392 accumulation in response to depolarization is unaffected in FBXO7<sup>-/-</sup> cells. Moreover, our 393 previous interaction proteomics experiments failed to identify either PINK1 in association 394 with overexpressed FBXO7 (Huttlin et al., 2021) or FBXO7 in association with 395 overexpressed Parkin (Sarraf et al., 2013). Thus, our functional and biochemical results do 396 not support a biochemical linkage between FBXO7 and the PINK1-Parkin pathway.

In order to broadly examine FBXO7<sup>-/-</sup> iNeurons for pathways that are affected, we 397 398 performed global proteomic analysis during neurogenesis in vitro. However, we did not 399 observe any alterations in pathways linked with mitochondria, or other quality control 400 pathways, although a reduction in the abundance of alcohol dehydrogenase enzymes 401 among a small number of other proteins, was observed. Whether these signatures are 402 related to FBXO7 function remains to be examined. While a recent study linked the L250P 403 mutation in the dimerization domain of FBXO7 with alterations in proteins associated with 404 mitochondrial fusion and fission (Al Rawi et al., 2022), we did not observe alterations in 405 either the abundance of these proteins (MiD51, MFF, FIS1) nor did we observe changes in 406 mitochondrial morphology in either HeLa cells or iNeurons lacking FBXO7.

407 An apparent ortholog of FBXO7 was identified in Drosophila - referred to as 408 nutcracker, ntc – and shown to associate with components of the core particle of the 409 proteasome, as well as with DmPI31 (Bader et al., 2011). Association of human FBXO7 with 410 proteasome components and PI31(also called PSMF1) have also been observed in both 411 focused studies and in our previous interaction proteomic studies (Al Rawi et al., 2022; 412 Huttlin et al., 2021; Vingill et al., 2016). In this context, PI31/DmPI31 and FBXO7/ntc have 413 been proposed to functionally link proteasome trafficking via dynein motors in neuronal 414 processes (Liu et al., 2019). Our interaction proteomics analysis has also confirmed 415 association of FBXO7 with PSMF1, CRL, COP9/Signalosome, and proteasome core 416 subunits (Supplemental Figure 4J,K). Further studies are required to elucidate the 417 relationship between FBXO7 and parkinsonian-pyramidal syndrome, to understand any 418 functional relationships between FBXO7 and other Parkinson's disease risk alleles, and to 419 examine whether FBXO7's physical association with the proteasome is linked with disease.

420

#### 421 ACKNOWLEDGMENTS

This work was supported by Aligning Science Across Parkinson's (ASAP) (J.W.H.), the NIH
(R01 NS083524 to J.W.H. and K01DK098285 to J.A.P.), and the Harvard Medical School

424 Cell Biology Initiative for Molecular Trafficking and Neurodegeneration. Michael J Fox 425 Foundation administers the grant ASAP-000282 on behalf of ASAP and itself. For the 426 purpose of open access, the author has applied a CC-BY public copyright license to the 427 Author Accepted Manuscript (AAM) version arising from this submission. We thank Laura 428 Pontano Vaites for help with the BioPlex 3.0 interactome analysis. We thank Jennifer 429 Waters, Talley Lambert, Federico Gasparoli and Rylie Walsh in the Nikon Imaging Center 430 and the Cell Biology Microscopy Facility (CBMF) at Harvard Medical School for microscopy 431 support and technical support.

432

### 433 KEYWORDS

- 434 FBXO7, mitophagy, proteomics, iNeurons
- 435

# 436 DATA & CODE AVAILABILITY

437 Proteomic data and analysis files part of this study are deposited at ProteomeXchange 438 Consortium by the PRIDE partner (Deutsch et al, 2020; Perez-Riverol et al, 2022). The 439 PRIDE project identification number is PXD037797 and can be accessed for reviewers 440 under username reviewer pxd037797@ebi.ac.uk; password 65wg8Sil. Macros and 441 pipelines used in this work can be found on GitHub 442 (https://github.com/harperlaboratory/FBXO7.git) Zenodo or 443 (https://doi.org/10.5281/zenodo.7258918). Raw files associated with this work are deposited 444 on Zenodo (https://doi.org/10.5281/zenodo.7268030).

445

#### 446 AUTHOR CONTRIBUTIONS

447 Study design: J.W.H., F.K., E.A.G.; Data collection: F.K., E.A.G., J.A.P., J.Z., Y.Z.; Data 448 analysis and interpretation: F.K., E.G., I.R.S., J.A.P., J.Z., Y.Z. L.P.V.; Manuscript 449 preparation: J.W.H., F.K. with input from all authors.

450

#### 451 **DECLARATION OF INTERESTS**

452 J.W.H. is a consultant and founder of Caraway Therapeutics and a founding scientific 453 advisory board member of Interline Therapeutics. No other authors declare a conflict of 454 interest.

- 455
- 456
- 457



Figure 1: Robust depolarization-dependent Ub phosphorylation by PINK1 cells
lacking FBXO7. (A) Immunoblot for pUb on HeLa control and FBXO7<sup>-/-</sup> cells after treatment
with AO for indicated times. (B) Quantification of pUb signal relative to loading. Immunoblots
run in triplicates. Error bars depict S.D.. (C) Scheme depicting production of iNeurons for
biochemical analysis. (D) Immunoblot for pUb on extracts from day 12 iNeuron control and
FBXO7<sup>-/-</sup> cells after treatment with BafA or AO for indicated times.



469 Figure 2. Super-resolution imaging of pUb formation independent of FBXO7. (A) 3D-SIM images of HeLa control, PINK1<sup>-/-</sup> and FBXO7<sup>-/-</sup> cell lines after AO-induced mitophagy. 470 471 Cells were stained for nuclear DNA (DAPI), mitochondria (HSP60) and pUb. Zoom-ins of 472 regions of interested are enlarged in the middle panel. 3D-surface renderings of insets are 473 shown on the right. Scale bar = 5  $\mu$ m or 1  $\mu$ m. (**B,C**) Evaluation of 3D-SIM images from 474 HeLa datasets. The changes in pUb volume and minimal distances between mitochondria 475 and pUb after mitophagy-induction are plotted. Error bars depict S.D. from 8-14 measured 476 cells per condition. (D) 3D-SIM images of iN day 12 control, PINK1<sup>-/-</sup> and FBXO7<sup>-/-</sup> cell lines 477 after AO-induced mitophagy. Cells were stained for nuclear DNA (DAPI), mitochondria 478 (HSP60) and pUb. Zoom-ins of regions of interested are enlarged in the middle panel. 3D-479 surface renderings of insets are shown on the right. Scale bar = 5  $\mu$ m or 1  $\mu$ m. (E,F) 480 Evaluation of 3D-SIM images from iNeuron datasets. The changes in pUb volume and 481 minimal distances between mitochondria and pUb after mitophagy-induction are plotted. 482 Error bars depict S.D. from 7-14 measured cells per condition. (G) Confocal images of iNeuron d12 Control, PINK1<sup>-/-</sup> and FBXO7<sup>-/-</sup> cell lines after AO-induced mitophagy. Cells 483 484 were stained for nuclear DNA (Hoechst33342), mitochondria (HSP60) and pUb. Scale bar = 485 10µm and 5µm. (H) Evaluation of pUb volume after mitophagy induction. Error bars depict 486 S.D. from 3 replicates.

487





490 Figure 3. Mitophagic flux in iNeurons and HeLa cells lacking FBXO7. (A) Scheme 491 depicting mitophagic flux assay using mtKeima. See text for details. (B) Mean Acidic:Neutral 492 mtKeima per-cell ratios measured by flow cytometry for HeLa cells expressing Parkin 493 ndicating the number of hours treated with AO (Antimycin A (0.5 µM) and Oligomycin (0.5 494 μM)) or three hours with 25 nM BafilomycinA (BafA). Error bars depict S.D. from triplicate 495 measurements. (C) Indicated hESCs expressing mtKeima were differentiated to iNeurons 496 (day 12) and were subjected to image analysis. Scale bar, 5 µm. (D) The number of red-497 shifted Keima foci per cell is plotted, where each dot represents the average foci-number per 498 image stack, originating from 7-12 image stacks of iNeuron differentiations. Error bars 499 represent S.D.. (E) mtKeima mitophagy flux readout of indicated iNeuron genotypes. Cells 500 were treated with AO for either 1 or 6h and mitophagic flux measured by flow cytometry 501 (>10,000 cells). Pooled data from 2 biological replicates is shown, normalized to 6h BafA 502 treated cells. Error bars depict S.D.. (F) Bar graph showing the number of mtDNA 503 puncta/cell with or without treatment with AO (6 h) in control or FBXO7<sup>-/-</sup> day 12 iNeurons. 504 Error bars depict S.D. from triplicate differentiations. (G) Bar graph showing the number of 505 mtDNA puncta/cell with or without treatment with AO (6 h) in HeLa control or FBXO7<sup>-/-</sup> cells. 506 Error bars depict S.D. from triplicate replicates.



510 Figure 4. Proteomic analysis of HeLa cells lacking FBXO7 during mitophagy. (A) 511 Workflow for analysis of total protein abundance in HeLa cells expressing Parkin with and 512 without depolarization with AO (16 h). Cell extracts were digested with trypsin prior to 18-513 plex TMT labelling and analysis by mass spectrometry. (B) Volcano plots [Log<sub>2</sub> FC (16 h AO / Fed) versus -Log<sub>10</sub>(q-value)] for control or one of two FBXO7<sup>-/-</sup> cell lines with or without 514 515 treated with AO (16 h). Red dots, mitochondrial proteins; blue dots, autophagy proteins; grey dots, remainder of the proteome. (C) Violin plots [Log<sub>2</sub> (16 h AO/Fed)] of control or FBXO7<sup>-/-</sup> 516 517 cells depicting alterations in the abundance of mitochondrial protein (left panel) or specific

organelles or protein complexes (right panel). (**D**) Correlation plot of c27 and c41 FBXO7<sup>-/-</sup> clones. Log<sub>2</sub>FC (16 h AO for each clone relative to control cells) is plotted. (**E**) Correlation plots [Log<sub>2</sub> FC (16 h AO / Fed)] of the proteome of FBXO7<sup>-/-</sup>c27 or c41 clones against control. Red dots, mitochondrial proteins; blue dots, autophagy proteins; grey dots, remainder of the proteome. (**F**) PINK1 levels in control cells and in two FBXO7<sup>-/-</sup> clones were measured by TMT-proteomics in fed cells (n=3).



526 527 Figure 5. Proteomic analysis of human ES cells lacking FBXO7 during neurogenesis. 528 (A) Workflow for analysis of total protein abundance in ES cells undergoing NGN2-driven 529 neurogenesis with or without FBXO7. Cell extracts were digested with trypsin prior to 18plex TMT labelling and analysis by mass spectrometry. (B) Log<sub>2</sub> FC for the indicated 530 mitochondrial protein-groups in control or FBXO7<sup>-/-</sup> cells at day 0, 4 or 12 during 531 532 neurogenesis is shown in violin plots. (C) Log<sub>2</sub> FC for the indicated cellular organelle 533 proteins in control or FBXO7<sup>-/-</sup> cells at day 0, 4 or 12 during neurogenesis is shown in violin plots. (D) Volcano plots [Log<sub>2</sub> FC (FBXO7<sup>-/-</sup>/Control) versus -Log<sub>10</sub>(q-value)] for FBXO7<sup>-/-</sup> 534 535 (c52 and c89) and control cells at day 4 (left panel) or day 12 (right panel) of differentiation. 536 Proteins showing decreased or increased abundance are shown as blue or red dots. (E) 537 Relative abundance of DNM1L (Drp1), MFF, and FIS1 in control or FBXO7<sup>-/-</sup> cells at day 0, 4 538 or 12 during neurogenesis. (F,G) Mitochondrial morphology in iNeurons (iN) was assessed 539 using confocal imaging after staining cells with HSP60 to detect mitochondria and DAPI to 540 identify nuclei, in either fed cells or cells treated with AO (3 h) (left panel). The number of 541 median # of mitochondria/cell, median mitochondrial length, and median mitochondrial 542 circularity is shown. Error bars depict S.D. from biological triplicate experiments (9-12 image

stacks each), as shown in panel I left, center, and right, respectively. (H) Western Blot
analysis on HeLa control and FBXO7<sup>-/-</sup> whole cell lysate, probed for the mitochondrial fission
adapter MiD51. (I) Relative abundance of DNM1L, MFF, MiD51 and FIS1 in WT (Control) or
FBXO7<sup>-/-</sup> HeLa cells either in the fed state or after 16 h AO as determined from the
proteomics data in Figure 5A,B.

548



550 551

552 Supplemental Figure 1: Tool kit for analysis of HeLa cells and iNeurons lacking 553 FBX07. (A) Working model summarizing the suggested modes of action of FBX07 in 554 PINK1/Parkin mitophagy. (B) Targeting of gRNA sequence within FBXO7 gene along with 555 the sequences of alleles A and B. MiSeq analysis of the three clones used for this study are shown. (B) Schematic for generation of FBXO7<sup>-/-</sup> in hESC cell lines. (C) Western Blot 556 analysis on HeLa control and FBXO7<sup>-/-</sup> whole cell lysate. (D) Targeting of gRNA sequence 557 558 within FBXO7 gene along with the sequences of alleles A and B. MiSeq analysis of the three 559 HeLa clones (c27, c41, c45) used for this study are shown. (E) Western Blot analysis on 560 hESC control and FBXO7<sup>-/-</sup> whole cell lysate. Asterisk indicates non-specific band. (F) 561 Targeting of gRNA sequence within FBXO7 gene along with the sequences of alleles A and 562 B. MiSeq analysis of the two clones (c52, c89) used for this study are shown. (G) Karyotype analysis of FBXO7<sup>-/-</sup> c52 and c89 hESCs. 563

564





566 567

567 568 **Supp** 

Supplemental Figure 2. Super-resolution pUb detection in HeLa cells in response to mitochondrial depolarization. 3D-SIM images of HeLa control, PINK1<sup>-/-</sup> and FBXO7<sup>-/-</sup> cell lines after 1h, 3h or 6h, AO-induced mitophagy (related to **Figure 2A,B**). Cells were stained for nuclear DNA (DAPI), mitochondria (HSP60) and pUb. (B) Zoom-ins of regions of interested after 1h AO-induced mitophagy are depicted and 3D-surface renderings of insets are shown on the right. Scale bar = 5  $\mu$ m or 1  $\mu$ m.

- 574
- 575



576 577

578 Supplemental Figure 3. Super-resolution pUb detection in iNeurons in response to 579 mitochondrial depolarization. (A) 3D-SIM images of iNeurons day 12 control, PINK1<sup>-/-</sup> and FBXO7<sup>-/-</sup> cell lines after AO-induced mitophagy. Cells were stained for nuclear DNA (DAPI), 580 mitochondria (HSP60) and pUb. Related to Figure 2D. Scale bar = 5  $\mu$ m or 1 $\mu$ m. (B) 581 Confocal images of iNeurons day 12 control, PINK1<sup>-/-</sup> and FBXO7<sup>-/-</sup> cell lines after AO-582 583 induced mitophagy with lower AO concentrations. Cells were stained for nuclear DNA 584 (DAPI), mitochondria (HSP60) and pUb. (C) Evaluation of images depicted in B. Error bars 585 depict S.D. from 12 image stacks measured per condition. Scale bar = 10  $\mu$ m and 5  $\mu$ m.



591 592 593 analysis of Parkin recruitment to mitochondria in single cells / mitochondrial organelles. (C) 594 Std mitochondrial signal vs Parkin intensity and Std Parkin vs mitochondria with or without 595 1h AO-induced mitophagy are shown. Data from three replicates, including a total of 238 image stacks, containing 16888 cells and 237848 mitochondrial objects. (D) Recruitment of 596 597 p62 to mitochondria (not stained) in control or FBXO7<sup>-/-</sup> HeLa cells with or without treatment with AO (16 h) was examined by confocal imaging. Scale bar = 10 and 5 µm. (E,F) 598 599 Quantification of cells in panel D. Assays were performed in biological triplicate with 4 image

588 589

600 stacks taken per repeat. n = 6712 cells. Error bars depict S.D.. (G) Quantification of #p62 foci normalized per cell in control, PINK1<sup>-/-</sup> and FBXO7<sup>-/-</sup> cells with or without 3h AO 601 602 treatment. Assays were performed in biological triplicate with 4 image stacks taken per 603 repeat . n = 12646 cells. Error bars depict S.D.. (H) Control or FBXO7<sup>-/-</sup> HeLa cells 604 expressing Parkin were either left untreated or incubated with AO for 1 or 6 h and extracts 605 subjected to immunoblotting with  $\alpha$ -LC3B and  $\alpha$ -Actin as a loading control. (I) The ratio of 606 LC3B lipidation (LC3B-II/LC3B-I) was quantified in 3 biological triplicate experiments. (J) Interaction network analysis of PINK1 (left) and FBXO7 (right) based on interaction 607 proteomics data from our BioPlex Interactome (Huttlin et al., 2021) (see METHODS for 608 609 details). (K) Summary of major interactions observed for FBXO7 in the context of the 610 relevant protein complexes. (L) Ridgeline plots of mtKeima-shift analysis in HeLa control, PINK1-1- and FBXO7-1- cell lines treated with AO at the indicated times. All lines are 611 612 normalized to the BafA sample.

- 613
- 614





617 Supplemental Figure 5. Proteomic analysis of HeLa cells lacking FBXO7 during 618 mitophagy. (A,B) Correlation plots (panel A) and PCA analysis (panel B) for ~8000 proteins quantified by TMT proteomics in individual replicates for the experiment outlined in Figure 619 4A. (C) Hierarchical clustering of 8000 proteins quantified in the experiment outlined in 620 Figure 4A. (D,E) Violin plots [Log<sub>2</sub> (control / FBXO7<sup>-/-</sup>)] for FBXO7<sup>-/-</sup> c27 and c41 cells 621 622 depicting alterations in the abundance of mitochondrial proteins (left panel) or specific 623 organelles or protein complexes (right panel). (F) Venn diagrams depicting the overlap in the 624 number of proteins whose levels are reduced (left panel) or increased (right panel) for the

data shown in **Figure 4B** are shown. Proteins whose abundance is significantly reduced upon AO treatment are enriched in proteins limited with mitochondrial function (right panel), as expected for cells undergoing mitophagy. (**G**) GO-term enrichment analysis of genes significantly down in both Control and FBXO7<sup>-/-</sup> after 16h AO mitophagy. (**H,I**) GO-term enrichment analysis of unique genes significantly down (panel H) or up (panel I) in FBXO7<sup>-/-</sup> after 16h AO mitophagy.



633 634

635 Supplemental Figure 6. Proteomic analysis of human ES cells lacking FBXO7 during 636 neurogenesis. (A) Hierarchical clustering and correlation plots of control and FBXO7<sup>-/-</sup> c52 637 (top) or c89 (bottom) at indicated days of differentiation in the experiment outlined in Figure 638 5A. (B) PCA analysis of whole cell proteomic samples from control and FBXO7<sup>-/-</sup> cells. 639 Triplicates for each timepoint and genotype were used. (C) FBXO7 peptide abundance in control and FBXO7<sup>-/-</sup> cells at indicated days of differentiation. (D) Violin plots [Log<sub>2</sub> (control / 640 641 FBXO7<sup>-/-</sup>)] for FBXO7<sup>-/-</sup> c89 cells depicting alterations in the abundance of mitochondrial 642 proteins.

643

644



648 Supplemental Figure 7. Comparison of human ES cell neurogenesis with or without FBX07. (A) Hotelling plots (Log<sub>10</sub> T<sup>2</sup> statistic versus Log<sub>2</sub> FC day 12/day 0) of control (left 649 650 panel) or FBXO7<sup>-/-</sup> c52 hES cells undergoing differentiation. Selected neurogenesis factors 651 are indicated. (B) The relative abundance of selected proteins is shown in the lower 652 histograms at day 0, 4 and 12 of differentiation. (C,D) Hotelling plots similar to A,B for control and FBXO7<sup>-/-</sup> c89. (E) Patterns of changes in the abundance of autophagy proteins in 653 control (left panel) and FBXO7<sup>-/-</sup> (right panel) day 12 iNeurons relative to day 0 cells 654 displayed on a Hotelling plot. (F) Cluster analysis of control and FBXO7<sup>-/-</sup> cells comparing 655 656 day 12 of differentiation to day 0. Individual clusters are displayed on Hotelling plots for each 657 genotype. 658

#### 659 EXPERIMENTAL MATERIALS AND METHODS

660 All details and catalogue numbers can be found in the **Materials Table (Supplemental** 661 **Table S9).** Protocols associated with this work can be found on protocols.io 662 (dx.doi.org/10.17504/protocols.io.kxygx99pwg8j/v1)

663

#### 664 Cloning and plasmid generation

The construction of lentiviral expression constructs used in this study have been previously described: pHAGE-mt-mKeima in (Heo *et al.*, 2015), PB-mt-mKeimaXL (Ordureau *et al.*, 2021) and pHAGE-GFP-Parkin in (Ordureau *et al.*, 2018).

668

# 669 Cell culture, generation of lentiviral stable cell lines.

HeLa Flip-In T-Rex (HFT) expressing doxycycline-inducible WT-Parkin (HeLa HFT/TOPRKN-WT) cells (Heo *et al.*, 2015) and HEK293T were maintained in Dulbecco's modified
Eagles medium (DMEM), supplemented with 10% vol/vol fetal bovine serum (FBS), 5%
vol/vol penicillin-streptomycin (P/S), 5% vol/vol GlutaMAX and 5% vol/vol Non-essential
amino acids (NEAA) at 37°C, 5% O<sub>2</sub>.

Stable cell lines were generated using lentivirus generated from HEK293T cells.
pHAGE-mt-mKeima or pHAGE-eGFP-Parkin were co-transfected in HEK293T cells together
with the lentiviral vectors pSpax2 and pMD2.1 using Lipofectaime LTX reagent (Thermo
Fisher Scientific, 15338100), according to manufacturer's instructions. Virus was harvested,
filtered (0.45 μm) and added 8μg/ml polybrene and target cells infected for 18 h. Antibiotic
selection was performed using 1 μg/ml puromycin or FACS sorting and cells verified using
immunoblotting or fluorescence microscopy.

If not stated otherwise, Parkin expression in HeLa HFT/TO-PRKN was induced using 2
 µg/ml doxycycline o/n, and subsequent mitophagy induced using Antimycin A / Oligomycin A
 (==AO) for the indicated times.

685 H9 ES cells harboring the mitochondrial mt-mKeima flux reporter (Ordureau *et al.*, 686 2020) were generated by electroporation of  $1x10^6$  cells with 2.5 µg of pAC150-PiggyBac-687 matrix-mKeima<sup>XL</sup> along with 2.5 µg of pCMV-HypBAC-PiggyBac-Helper, as described 688 (Ordureau *et al.*, 2020). The cells were selected and maintained in E8 medium 689 supplemented with 50 µg/ml Hygromycin and Hygromycin was kept in the medium during 690 differentiation to iNeurons.

691

#### 692 Gene-Editing and iNeuron differentiation

Generation of HeLa HFT cells lacking FBXO7<sup>-/-</sup> was facilitated using CRISPR/Cas9 with 693 694 target sites determined using CHOPCHOP (Labun et al, 2019). Guide RNAs were ligated 695 into the px459 plasmid (Addgene plasmid # 62988) and cells transfected using Lipofectaime 696 LTX reagent (Thermo Fisher Scientific, 15338100), according to manufacturer's instructions. 697 Two days post-transfection, single GFP positive cells were sorted into 96-well dishes 698 containing 300µl full growth medium (composition as described above). Single cells were 699 allowed to grow into colonies, then duplicated for MiSeq analysis and maintenance. 700 Knockout candidates were confirmed by Western blot on whole cell lysates. The sgRNAs 701 were generated using GeneArt Precision gRNA Synthesis Kit (Thermo Fisher Scientific) 702 according to the manufacturer's instruction and purified using RNeasy Mini Kit (Qiagen). The 703 sgRNA target sequence: ACCGATTCACTACAGAGCAT.

704 The PINK1<sup>-/-</sup> H9 cells used here, wherein sequences in exon 1 were deleted using 705 CRISPR-Cas9 to create a null allele, were reported previously (Ordureau et al., 2018). To 706 generate FBXO7<sup>-/-</sup> H9 ES or HeLa cells, 0.6 µg sgRNA was incubated with 3 µg SpCas9 707 protein for 10 minutes at room temperature and electroporated into 2x10<sup>5</sup> WT H9 cells using 708 Neon transfection system (Thermo Fisher Scientific). Mutants were identified by Illumina 709 MiSeq and further confirmed by Western blotting. For introduction of TRE3G-NGN2 into the 710 AAVS1 site, a donor plasmid pAAVS1-TRE3G-NGN2 was generated by replacing the EGFP 711 sequence with N-terminal flag-tagged human NGN2 cDNA sequence in plasmid pAAVS1-712 TRE3G-EGFP (Addgene plasmid # 52343). Five micrograms of pAAVS1-TRE3G-NGN2, 2.5 713 μg hCas9 (Addgene plasmid # 41815), and 2.5 μg gRNA AAVS1-T2 (Addgene plasmid # 41818) were electroporated into  $1 \times 10^6$  H9 cells. The cells were treated with 0.25 µg/ml Puromycin for 7 days and surviving colonies were expanded and subjected to genotyping.

716 Human ES cells (H9, WiCell Institute) were cultured in E8 medium (Chen et al, 2011) 717 on Matrigel-coated tissue culture plates with daily medium change. Cells were passaged 718 every 4-5 days with 0.5 mM EDTA in 1× DPBS (Thermo Fisher Scientific). SpCas9 and 719 AsCas12a/AsCpf1 expression plasmids pET-NLS-Cas9-6xHis (Addgene plasmid # 62934) 720 and modified pDEST-his-AsCpf1-EC, generated by deleting the MBP sequence from 721 plasmid pDEST-hisMBP-AsCpf1-EC (Addgene plasmid # 79007), were transformed into 722 Rosetta™(DE3)pLysS Competent Cells (Novagen), respectively, for expression. SpCas9 723 and AsCas12a/AsCpf1 proteins were purified as described elsewhere (Hur et al, 2016; Zuris 724 et al, 2015). Briefly, cells expressing SpCas9 [0.5 mM isopropylthio-β-galactoside, 14-hour 725 induction] were lysed in FastBreak buffer (Promega, Inc) and the NaCl concentration 726 adjusted to 500 mM. Extracts were centrifuged at 38,000g for 10 min at 4°C and the 727 supernatant incubated with Ni-NTA resin for 1 hour. The resin was washed extensively with 728 50 mM Tris pH8.0, 500 mM NaCl, 10% glycerol, 20 mM imidazole, and 2 mM TCEP prior to 729 elution with this buffer supplemented with 400 mM imidazole. Proteins were diluted two 730 volumes/volume in PBS and fractionated on a Heparin-Sepharose column using a 0.1 to 1.0 731 M NaCl gradient. Cas9-containing fractions were stored in PBS, 20% glycerol, 2 mM TCEP 732 at -80°C. AsCpf1 expression was induced similarly, and cells pelleted by centrifugation. Cells 733 were lysed by sonication in 50 mM HEPES pH7, 200 mM NaCl, 5 mM MgCl<sub>2</sub>, 1 mM DTT, 734 and 10 mM imidazole supplemented with lysozyme (1 mg/ml) and protease inhibitors (Roche 735 complete, EDTA-free). After centrifugation (16,000g for 30 min), the supernatant was 736 incubated with Ni-NTA resin, the resin washed with 2M NaCl, and bound proteins eluted with 737 250 mM imidazole, and buffer exchanged into lysis buffer lacking MgCl<sub>2</sub> and imidazole prior 738 to storage at -80°C.

739 For human ES cell conversion to iNeurons, cells were expanded and plated at 740  $2 \times 10^4$ /cm<sup>2</sup> on Matrigel-coated tissue plates in DMEM/F12 supplemented with 1x N2, 1x 741 NEAA (Thermo Fisher Scientific), human Brain-derived neurotrophic factor (BDNF, 10 ng/ml, 742 PeproTech), human Neurotrophin-3 (NT-3, 10 ng/ml, PeproTech), mouse laminin (0.2 µg/ml, 743 Cultrex), Y-27632 (10 µM, PeproTech) and Doxycycline (2 µg/ml, Alfa Aesar) on Day 0. On 744 Day 1, Y-27632 was withdrawn. On Day 2, medium was replaced with Neurobasal medium 745 supplemented with 1x B27 and 1x Glutamax (Thermo Fisher Scientific) containing BDNF, 746 NT-3 and 1 µg/ml Doxycycline. Starting on Day 4, half of the medium was replaced every 747 other day thereafter. On Day 7, the cells were treated with Accutase (Thermo Fisher 748 Scientific) and plated at 3-4x10<sup>4</sup>/cm<sup>2</sup> on Matrigel-coated tissue plates. Doxycycline was 749 withdrawn on Day 10.

750

# 751 Immunoblotting

752 At the indicated times, ES cells, iNeurons or HeLa cells were washed on ice in 1x PBS, 753 harvested and pellet wash with 1x PBS and resuspended in 8 M urea buffer (8 M urea, 150 754 mM TRIS pH 7.4, 50 mM NaCl, PhosSTOP phosphatase inhibitor cocktail). Resuspended 755 cell lysates were sonicated for 10 seconds and debris pelleted at 13,000 rpm for 10 min. 756 Protein concentration was determined by BCA assay according to manufacturer's instructions (Thermo Fisher Scientific, 23227). Indicated amounts of proteins were 757 758 resuspended in 1xLDS + 100 mM DTT and boiled for 10 minutes at 85°C. Equal amount of 759 protein and volume were loaded run on 4%-20% Bis-Tris, 8% Tris NuPAGE gels for 5 760 minutes at 100V, 5 min at 150 V and then run at 200 V for the required time. Gels were 761 transferred via wet transfer system onto PDVF membranes for immunoblotting. 762 Chemiluminescence and colorimetric images were acquired using a BioRad ChemiDoc MP 763 imaging system. Images from Western Blots were exported and analysed using Image Lab 764 and ImageJ/FiJi (Schindelin et al, 2012).

- 765
- 766
- 767
- 768

769 770

771 PROTEOMICS

#### 772 Proteomics – general sample preparation

773 Sample preparation of proteomic analysis of whole-cell extract from HeLa, hESC, NPC and 774 iNeurons was performed according to previously published studies (Ordureau et al., 2021; 775 Ordureau et al., 2020). Cells were harvested on ice and plates were washed twice with 1x 776 PBS and detached in 1x PBS using cell scraper. After pelleting at 2000 rpm for 5 min at 4°C. 777 cells were washed 2x with 1x PBS and resuspended in 8 M urea buffer (composition stated 778 above). After sonification for 10 seconds, resuspended cells were pelleted for 10 min at 779 13000 rpm. Protein concentration was determined using BCA kit (Thermo Fisher Scientific, 780 23227).

781 Unless otherwise noted, proteomics and data analysis was performed as described 782 (Ordureau et al., 2021; Ordureau et al., 2020). Briefly, protein extracts (100 µg) were 783 subjected to disulfide bond reduction with 5 mM TCEP (room temperature, 10 min) and 784 alkylation with 25 mM chloroacetamide (room temperature, 20 min). Methanol-chloroform 785 precipitation was performed prior to protease digestion. In brief, four parts of neat methanol 786 were added to each sample and vortexed, one part chloroform was then added to the 787 sample and vortexed, and finally three parts water was added to the sample and vortexed. 788 The sample was centrifuged at 6,000 rpm for 2 min at room temperature and subsequently 789 washed twice with 100% methanol. Samples were resuspended in 100 mM EPPS pH8.5 790 containing 0.1% RapiGest and digested at 37 °C for 8h with Trypsin at a 100:1 protein-to-791 protease ratio. Samples were acidified with 1% Formic Acid for 15 min and subjected to C18 792 solid-phase extraction (SPE) (Sep-Pak, Waters).

793

## 794 *Proteomics – quantitative proteomics using TMT*

795 Tandem mass tag labeling of each sample (50 mg peptide input) was performed by adding 5 796 uL of the 25 ng/mL stock of TMTpro reagent along with acetonitrile to achieve a final 797 acetonitrile concentration of approximately 30% (v/v). Following incubation at room 798 temperature for 1 h, the reaction was guenched with hydroxylamine to a final concentration 799 of 0.5% (v/v) for 15 min. The TMTpro- labeled samples were pooled together at a 1:1 ratio. 800 The sample was vacuum centrifuged to near dryness, and following reconstitution in 1% FA, 801 samples were desalted using C18 solid-phase extraction (SPE) (50 mg, Sep-Pak, Waters), 802 according to manufacturer protocol.

803 Dried TMTpro-labeled peptides (~300 ug) were resuspended in 10 mM NH<sub>4</sub>HCO<sub>3</sub> pH 804 8.0 and fractionated using basic pH reverse phase HPLC (Wang et al, 2011). Briefly, 805 samples were offline fractionated over a 90 min run, into 96 fractions by high pH reverse-806 phase HPLC (Agilent LC1260) through an Aeris peptide xb-c18 column (Phenomenex: 250 807 mm x 3.6 mm) with mobile phase A containing 5 % acetonitrile and 10 mM NH<sub>4</sub>HCO<sub>3</sub> in LC-808 MS grade  $H_2O$ , and mobile phase B containing 90 % acetonitrile and 10 mM  $NH_4HCO_3$  in 809 LC-MS grade  $H_2O$  (both pH 8.0). The 96 resulting fractions were then pooled in a non-810 continuous manner into 24 fractions (as outlined in Figure S5 of (Paulo et al, 2016)) and sets 811 of 12 fractions (even or odd numbers) were used for subsequent mass spectrometry 812 analysis. Fractions were vacuum centrifuged to near dryness. Each consolidated fraction 813 was desalted via StageTip, dried again via vacuum centrifugation, and reconstituted in 5 % 814 acetonitrile, 1 % formic acid for LC-MS/MS processing.

815 TMT proteomics samples were subjected to analysis using a Orbitrap Fusion Lumos 816 Tribrid mass spectrometer (Thermo Fisher Scientific, San Jose, CA) online with Proxeon 817 EASY-nLC1200 liquid chromatography (Thermo Scientific). Peptides were resuspended in 5 818 % ACN/5 % FA and 10 % of the samples were loaded on a 35 cm analytical column (100mm 819 inner diameter) packed in-house with Accurcore150 resin (150 Å, 2.6 mm, Thermo Fisher 820 Scientific, San Jose, CA) for LC-MS analysis. Peptide separation was performed with a 821 gradient of acetonitrile (ACN, 0.1% FA) from 3-13 % (0-83 min) and 13-28 % (80-83 min) 822 during a 90 min run.

823 For the HeLa TMTpro proteomic samples, LC-hrMS/MS was combined with 3 824 optimized CV parameters on the FAIMS Pro Interface to reduced precursor ion interference 825 (Schweppe et al, 2019). Data-dependent acquisition (DDA) was performed by selecting the 826 most abundant precursors from each CV's (-40/-60/-80) MS<sup>1</sup> scans for hrMS/MS over a 1-827 1.5s duty cycle (1s/1.5s/1s respectively). The MS<sup>1</sup> scan parameters include a 400-1,600 m/z mass range at 60,000 resolution (at 200 Th) with 4 x  $10^5$  automated gain control (AGC) (100 828 829 %), and a maximum injection time (max IT) of 50 ms. Precursors (z=2-5) were isolated with 830 0.7 Th (quadrupole), fragmented with high energy C-trap dissociation (HCD) at 36 831 normalized collision energy (NCE), and subjected to hrMS/MS on the Orbitrap at 50,000 832 resolution (at 200 Th), 120 ms max IT, fixed first mass 110 Th, and 1.0 x 10<sup>5</sup> AGC (200%). 833 Precursors were placed on 90 s dynamic exclusion (+/- 10 ppm) to prevent redundant 834 sampling.

835 In the iNeuron TMTpro experiments, the same FAIMS and MS<sup>1</sup> parameters were 836 implemented (with 1.25 s duty cycle/CV for DDA) with the Multi-Notch SPS-MS3 acquisition 837 method (McAlister et al, 2014), to further reduce ion interference in TMT reporter quantification (Paulo et al., 2016). Most abundant precursors (with 120 s dynamic exclusion 838 839 +/- 10 ppm) were selected from  $MS^1$  scans, isolated using the quadrupole (0.6 Th isolation), 840 fragmented with collision-induced dissociation (CID) at 35 NCE, and subjected to MS/MS in 841 the ion trap (turbo scan speed, 35 ms max IT, 1.0 x  $10^4$  AGC). Using Real Time Search 842 analysis software (Erickson et al, 2019; Schweppe et al, 2020), a synchronous-precursor-843 selection (SPS) API-MS<sup>3</sup> scan was collected on the top 10 most intense b- or y-ions from the 844 matched peptide identification (determined by an online search of its respective MS/MS 845 scan). MS<sup>3</sup> scans were performed on the Orbitrap (AGC 2.0 x 10<sup>5</sup>; NCE 55; max IT 250 ms, 846 50,000 resolution at 200 Th). To increase quantitative sampling (MS<sup>3</sup> scans) of proteins 847 during each mass spectrometry injection, a 2 peptide per protein per sample closeout was 848 set. This ensures no more than two peptide-spectrum matches per protein (that pass quality 849 filters) are subjected to MS<sup>3</sup> scans, reducing redundant protein MS sampling and potentially 850 increasing proteome depth (Schweppe et al., 2020).

851

# 852 Proteomics Data Analysis

853 Mass spectrometry raw data were converted to mzXML and monoisotopic peaks were 854 reassigned using Monocle (Rad et al, 2021). Mass spectra were database searched using 855 Sequest algorithm (2019.01 rev. 5; (Eng et al, 1994)) against the Human Reference 856 Proteome Uniprot database (2019-01 SwissProt entries only; UniProt Constortium, 2015) 857 appended with sequences of common contaminates and reverse sequences of proteins as 858 decoys, for target-decoy competition (Elias & Gygi, 2007). Sequest search parameters 859 include: 50 ppm precursor tolerance, 0.9 Da product ion tolerance, trypsin endopeptidase 860 specificity (C-terminal to [KR], 2 max missed cleavages), static modifications on peptide N-861 terminus and lysines with TMTpro tags (+304.207 Da) and carbamidomethylation on 862 cysteines (57.021 Da), and variable modification of oxidation on methionines (+15.995 Da). 863 Peptide-spectrum matches were filtered at 2% false discovery rate (FDR) using linear 864 discriminant analysis (Huttlin et al, 2010), based on XCorr, DeltaCn, missed cleavages, 865 peptide length, precursor mass accurracy, fraction of matched product ions, charge state, 866 and number of modifications per peptide (additionally restricting PSM Xcorr >1 and peptide 867 length>6). Following a 2% protein FDR filter (Savitski et al, 2015), PSMs reporter ion 868 intensities were quantified (most intense centroid within 0.003 Da of theoretical TMT reporter 869 mass), filtered based on a precursor isolation specificity > 0.5, and filtered by a summed 870 signal-to-noise across TMT channels > 100.

Protein quantification was performed via the summation (weighted average) of its constituent PSMs' reporter intensities and TMT channels were normalized for protein input to total TMT channel intensity across all quantified PSMs (adjusted to median total TMT intensity for the TMT channels) (Plubell *et al*, 2017). Log<sub>2</sub> normalized summed protein reporter intensities were compared using a Student's t-test and p-values were corrected for multiple hypotheses using the Benjamini-Hochberg adjustment (Benjamini & Hochberg, 1995). Resultant q-values and mean log<sub>2</sub> fold changes between conditions were used to generate volcano plots. Hotelling  $T^2$  analysis was performed using the normalized summed protein reporter ion intensities and the timecourse (v. 1.66.0; (Tai, 2022)) package in R.

880 The annotation list for the subcellular localization of organellar protein markers was 881 derived from previously published high confidence HeLa dataset ((Itzhak et al, 2016); "high" 882 & "very high" confidence) and additional manual entries (Ordureau et al., 2021). MitoCharta 883 3.0 (Rath et al, 2021) was used for mitochondrial annotation, whereas annotations for 884 ribosome and autophagy components were used from previous studies (An et al, 2020). 885 Transcription factor and neuronal development markers were based on previously published 886 databases and publications (Lambert et al, 2018; Ordureau et al., 2021). Figures were 887 generated using a combination of Excel, R (v.4.2.0) in RStudio (2022.07.01 Build 554), 888 Perseus (v1.6.5 (Tyanova & Cox, 2018)), GraphPad Prism (v9.1.0), and Adobe Illustrator 889 (26.2.1).

890

The Supplemental Data Table S3-8 contains the quantified proteins as well as associated
 TMT reporter ratio to control channels used for quantitative analysis.

893

894 Interaction Proteomic Analysis

For interaction analysis of FBXO7 and PINK1, interaction proteomics data from the Bioplex Interactome (Huttlin *et al.*, 2021) was extracted for HEK293T and HCT116 samples (**Supplemental Table S1,2**). The number of replicates in which a peptide-spectral match (PSM) mapped to PINK1's or FBXO7's the Bioplex 3.0 interactomes was plotted via a network diagram in R. (packages to add from R include (igraph 1.3.1, tidygraph 1.2.1, and ggraph 2.0.5).

901

902 MICROSCOPY

903 and pipelines Macros used in this work can be found on GitHub 904 (https://github.com/harperlaboratory/FBXO7.git) or 905 Zenodo (https://doi.org/10.5281/zenodo.7258918).

906

907 Live-cell confocal microscopy for mitophagic flux analysis over differentiation (mt-mKeima<sup>XL</sup>) 908 For guantitative Keima-flux analysis (Ordureau et al., 2020), hESC were seeded on day 4 of 909 differentiation into 6-well 1.5 high performance glass bottom plates (Cellvis, P06-1.5H-N) 910 and further differentiated in the vessel for the indicated times until reaching appropriate 911 confluency and imaged. iNeurons were imaged using a Yokogawa CSU-W1 spinning disk 912 confocal on a Nikon Eclipse Ti-E motorized microscope. The system is equipped with a 913 Tokai Hit stage top incubator and imaging was performed at 37°C, 5% CO<sub>2</sub> and 95% 914 humidity under a Nikon Plan Apo 60×/1.40 N.A immersion oil objective lens. For ratiometic 915 imaging, mtKeimaXL were excited in sequential manner with a Nikon LUN-F XL solid state 916 laser combiner ([laser line - laser power]: 445 - 80mW, 561 - 65mW]) using a 917 Semrock Di01-T445/515/561 dichroic mirror. Fluorescence emissions were collected 918 through a Chroma ET605/52m [for 445 nm] and a 568 Chroma ET605/52m [for 561 nm], 919 filters, respectively (Chroma Technologies). Confocal images were acquired with a 920 Hamamatsu ORCA-Fusion BT CMOS camera (6.5 µm<sup>2</sup> photodiode, 16-bit) camera and NIS-921 Elements image acquisition software. Consistent laser intensity and exposure time were 922 applied to all the samples, and brightness and contrast were adjusted equally by applying 923 the same minimum and maximum display values in ImageJ/FiJi (Schindelin et al., 2012). 924 Image quantification was performed in ImageJ/FiJi using custom-written batch-macros.

925 In brief, raw confocal images of mitochondrial targeted mt-mKeima<sup>XL</sup> were divided 926 [ex:561/ex:445] resulting in a ratiometic image of only acidic Keima-puncta. These signals 927 were subjected to background subtraction (rolling kernel size 25, sliding paraboloid) and 928 converted to binary objects. The "Analyze Particles..." command (pixel size exclusion: 0.5-∞, 929 exclude edge objects) was used to measure foci-abundance and other morphological 930 parameters. Results for each image-stack saved as .csv files, together with the original 931 ratiometic .tiff file for QC purposes. Unless stated otherwise all images represent z932 projections. Statistical analysis and plotting of microscopy data was performed in Prism 933 (v9.1.0, GraphPad).

- 934
- 935
- 936
- 937

# 938 Immunocytochemical analysis

iNeurons or HeLa cells were fixed with warm 4% paraformaldehyde (Electron Microscopy 939 940 Science, #15710, purified, EM grade) in PBS at 37°C for 30 min and permeabilized with 941 0.5% Triton X-100 in PBS for 15 minutes at room temperature. After three washes with 942 0.02% Tween20 in PBS (PBST), cells were blocked for 10 min in 3% BSA-1xPBS at room 943 temperature and washed again three times in PBST. Cells were incubated for 3h in primary 944 antibodies in 3% BSA-1xPBS and washed three times with PBST. Secondary antibodies 945 (Thermo Scientific, 1:400 in 3% BSA-1xPBS) where applied for 1h at room temperature. 946 Alexa Fluor 633 Phalloidin (Thermo Fisher, 1:200, A22284) was added with secondary 947 antibodies to label F-actin. To stain nuclei, Hoechst33342 (1:10000) was added for 5 min to 948 cells in PBST and final three washes performed before mounting in Vectashield (Vector 949 Laboratories, H-1000-10). Primary and secondary antibodies used in this study can be found 950 in the Materials Table (Supplemental Table S9).

951

# 952 Fixed-cell microscopy – general acquisition parameters

953 Immunofluorescently labelled Hela or iNeurons (antibodies indicated in figures and figure 954 legends and details in Materials Table, (Supplemental Table S9)) were imaged at room 955 temperature using a Yokogawa CSU-W1 spinning disk confocal on a Nikon Eclipse Ti-E 956 motorized microscope equipped with a Nikon Plan Apochromat 40×/0.40 N.A air-objective 957 lens, Nikon Apochromat 60×/1.42 N.A oil-objective lens and a Plan Apochromat 100×/1.45 N.A oil-objective lens. Signals of 405/488/568/647 fluorophores were excited in sequential 958 959 manner with a Nikon LUN-F XL solid state laser combiner ([laser line – laser power]: 405 -960 80mW, 488 - 80mW, 561 - 65mW, 640nm - 60mW]) using a Semrock Di01-961 T405/488/568/647 dichroic mirror. Fluorescence emissions were collected with 962 Chroma ET455/50m [405 nm], 488 Chroma ET525/50m [488 nm], 568 Chroma ET605/52m 963 [561 nm], 633 Chroma ET705/72m [640 nm] filters, respectively (Chroma Technologies). 964 Confocal images were acquired with a Hamamatsu ORCA-Fusion BT CMOS camera (6.5 965  $\mu m^2$  photodiode, 16-bit) camera and NIS-Elements image acquisition software. Consistent 966 laser intensity and exposure time were applied to all the samples, and brightness and 967 contrast were adjusted equally by applying the same minimum and maximum display values 968 in ImageJ/FiJi (Schindelin et al., 2012).

969

#### 970 Microscopy-based mitochondrial morphology measurements in iNeurons

971 For quantitative measurement of mitochondrial morphology in d12 iNeurons under fed or 972 mitophagy conditions, HeLa cells were seeded on 6-well 1.5 high performance glass bottom 973 plates (Cellvis, P06-1.5H-N) and mitophagy induced for the indicated time durations. 974 Alternatively, hESC were seeded on day 4 of differentiation into 6-well 1.5 high performance 975 glass bottom plates (Cellvis, P06-1.5H-N) and differentiated in the vessel into iNeurons and 976 mitophagy induced for the indicated time durations. Cells were fixed and stained as 977 described above. Z-stacks were acquired with a Nikon Plan Apo 100×/1.45 N.A oil-objective 978 lens and with the parameters stated above. Image quantification was performed in 979 ImageJ/FiJi using custom-written batch-macros.

Statistical analysis and plotting of microscopy data was performed in Prism (v9.1.0,
 GraphPad). Primary and secondary antibodies used in this study can be found in the
 Materials Table (Supplemental Table 9).

983

# 984 Microscopy-based mtDNA turnover measurements in HeLa and iNeurons

For quantitative measurement of mtDNA turnover after AO-induced mitophagy, HeLa cells were seeded on 6-well 1.5 high performance glass bottom plates (Cellvis, P06-1.5H-N) and

987 mitophagy induced for the indicated time durations. Alternatively, hESC were seeded on day 988 4 of differentiation into 6-well 1.5 high performance glass bottom plates (Cellvis, P06-1.5H-989 N) and differentiated in the vessel into iNeurons and mitophagy induced for the indicated 990 time durations. Cells were fixed and stained as described above, but primary antibody 991 incubation with aDNA (1:200) was performed overnight. Z-stacks were acquired with a Nikon 992 Plan Apo 100×/1.45 N.A oil-objective lens and with the parameters stated above. Image 993 guantification was performed in ImageJ/FiJi using custom-written batch-macros. In brief, 994 both aDNA and nuclear signals were converted to binary files and the nuclear signal 995 subtracted from the aDNA signal, resulting in an image stack containing only the mtDNA 996 intensities. The "Analyze Particles..." command (pixel size exclusion: 0.05-3, exclude edge 997 objects) was used to measure morphological features and results for each image-stack 998 saved as .csv files, together with the analyzed binary-mask overlay .tiff file for QC purposes. 999 Number of mtDNA signals were normalized to cell number found in the same image stack. 1000 Statistical analysis and plotting of microscopy data was performed in Prism (v9.1.0, 1001 GraphPad). Primary and secondary antibodies used in this study can be found in the 1002 Materials Table (Supplemental Table S9).

1003

# 1004 Microscopy-based measurements of p62 recruitment in HeLa

1005 For quantitative measurement of p62 recruitment to mitochondria, HeLa control and 1006 knockout cells were seed in 1.5 high performance glass bottom plates and doxycycline 1007 added over night to induce Parkin expression. Mitophagy was induced using Oligomycin A / 1008 Antimycin A for 16h in full DMEM in presence of doxycycline. Cells were fixed and stained 1009 as stated above and imaged using a Nikon Plan Apo  $60 \times /1.42$  N.A air-objective lens and 1010 with the parameters stated above. 8  $\mu$ M z-stacks were taken for each selected field of view.

1011 Image quantification was performed in ImageJ/FiJi using custom-written batch-1012 macros. The p62 channel was filtered (Gaussian Blue, sigma =2) and converted into binary 1013 files using the Intermodes thresholding method. p62 spots were counted using the "Analyze 1014 Particles..." function (pizel size exclusion: 0.1-30, exclude edge objects) and results for each 1015 image-stack saved as .csv files, together with the analyzed binary-mask overlay .tiff file for 1016 QC purposes. The DAPI channel was used to count nuclei for per cell normalization. 1017 Statistical analysis and plotting of microscopy data was performed in Prism (v9.1.0, 1018 GraphPad). Primary and secondary antibodies used in this study can be found in the 1019 Materials Table (Supplemental Table S9).

- 1020
- 1021

# Microscopy-based pUb-coverage measurements of mitochondria in iNeurons

For quantitative measurement of pUb coverage over mitochondria AO-induced mitophagy, hESC were seeded on day 4 of differentiation into 6-well 1.5 high performance glass bottom plates (Cellvis, P06-1.5H-N) and differentiated in the vessel into iNeurons and mitophagy induced for the indicated time durations. Cells were fixed and stained as described above, using anti-HSP60 to label mitochondria and anti-pUb (Ser65) to label pUb. 3-5 10 µm thick z-stacks per replicate per sample were acquired using a Nikon Plan Apo 60×/1.42 N.A airobjective lens and with the parameters stated above.

1029 Image quantification was performed in ImageJ/FiJi using custom-written batch-1030 macros. In brief, mitochondrial signal was filtered (Gaussian Blur, sigma=2), converted into 1031 binary files and holes in the resulting mask filled. pUb channel was thresholded into a binary 1032 file (Triangle method); these masks were measured using the "Analyze Particles..." 1033 command (pixel size exclusion: 0.5-∞, exclude edge objects) and results for each image-1034 stack saved as .csv files, together with the analyzed binary-mask overlay .tiff file for QC 1035 purposes. % of mitochondrial pUb coverage was calculated and normalized to [t]=6h AO. 1036 Statistical analysis and plotting of microscopy data was performed in Prism (v9.1.0, 1037 GraphPad). Primary and secondary antibodies used in this study can be found in the 1038 Materials Table (Supplemental Table S9).

1039

1040 Microscopy-based evaluation of Parkin translocation and mitophagy in FBXO7<sup>-/-</sup> cell lines

For guantitative measurement of Parkin translocation kinetics in FBXO7<sup>-/-</sup> cell lines, stable 1041 1042 HeLa cell lines expressing GFP-Parkin were created. WT and knockout cell lines were 1043 seeded into 12-well 1.5 high performance glass bottom plates (Cellvis, P12-1.5H-N) two 1044 days prior to experimental manipulation. Mitophagy was induced as described above using 1045 Antimycin A / Oligomycin A for 1h. Treated and control (fed) cells were fixed as described 1046 above, stained for mitochondria (HSP60), Parkin (GFP) and DNA (SpyDNA-555) and 8µm z-1047 stacks were acquired with a Nikon Plan Apo 60×/1.42 N.A oil-objective lens and with the 1048 parameters stated above. For guantitate single-cell analysis, a total of ~240 randomly xy-1049 marked stacks were acquired, and MIPs used for subsequent analysis. CellProfiler (Stirling 1050 et al., 2021) was used for the quantitative analysis of single-cells and mitochondrial objects 1051 (segmentation and analysis pipelines can be found on GitHub). Plotting of microscopy data 1052 was performed in Prism (v9.1.0, GraphPad) and R using the following libraries: tidyverse, 1053 dplyr, tibble, viridis, ggplot2, ggridges, ggsci. Primary and secondary antibodies used in this 1054 study can be found in the Materials Table (Supplemental Table S9).

1055

# 1056 Immunocytochemical sample preparation for 3D-SIM

1057 Sample preparation and SIM acquisition guidelines from (Kraus et al, 2017) were used as a 1058 basis for the super-resolution analysis of pUb spreading after mitophagy induction. iNeurons 1059 or HeLa cells were seeded on 18x18 mm Marienfeld Precision cover glasses thickness No. 1060 1.5H (tol. ± 5 µm). Cover glasses were coated, if necessary, for hESC / iNeuron culture as 1061 described above. After experimental manipulation, cells were fixed with warm 1062 paraformaldehyde 3% Glutaraldehyde 0.35% in 0.1M Sodium Cacodylate, pH 7.4 (Electron 1063 Microscopy Science) at 37°C for 30 min and permeabilized with 0.5% Triton X-100 in PBS 1064 for 15 minutes at room temperature. After three washes with 0.02% Tween20 in PBS 1065 (PBST), cells were blocked for 10 min in 3% BSA-1xPBS at room temperature and washed 1066 again three times in PBST. iNeurons were incubated overnight in primary antibodies in 3% 1067 BSA-1xPBS and washed three times with PBST. Secondary antibodies (Thermo Scientific, 1068 1:400 in 3% BSA-1xPBS) where applied for 1 h at room temperature. To stain nuclei, DAPI 1069 was added for 5 min to cells in PBST, washed three times for 5 min in 1xPBST before a 1070 post-fixation in 4% paraformaldehyde was performed. After 2 washes in PBST, coverslips 1071 were washed once in 1xPBS and mounted in Vectashield (Vector Laboratories, H-1000-10) 1072 on glass slides. Primary and secondary antibodies used in this study can be found in the 1073 Materials Table (Supplemental Table S9).

1074

# 1075 3D-SIM microscopy – acquisition parameters

1076 3D-SIM microscopy was performed on a DeltaVision OMX v4 using an Olympus 60x / 1.42 1077 Plan Apo oil objective (Olympus, Japan). The instrument is equipped with 405 nm, 445 nm, 1078 488 nm, 514 nm, 568 nm and 642 nm laser lines (all >= 100 mW) and images were recorded 1079 on a front-illuminated sCMOS (PCO Photonics, USA) in 512x512px image size mode, 1x 1080 binning, 125 nm z-stepping and with 15 raw images taken per z-plane (5 phase-shifts, 3 1081 angles). Raw image data was computationally reconstructed using CUDA-accelerated 3D-1082 SIM reconstruction code (https://github.com/scopetools/cudasirecon) based on (Gustafsson 1083 et al, 2008). Optimal optical transfer function (OTF) was determined via an in-house build 1084 developed by Talley Lambert from the NIC / CBMF software. (GitHub: 1085 https://github.com/tlambert03/otfsearch, all channels were registered to the 528nm output 1086 channel, Wiener filter: 0.002, background: 90).

1087

# 1088 3D-SIM microscopy – pUb – mitochondria 3D renderings & analysis

1089 3D renderings of 3D-SIM images were performed in UCSF Chimera X 1090 (https://www.cgl.ucsf.edu/chimerax/) using 32-bit .tiff image stacks. Image channels were 1091 sequentially imported and visualized as surfaces using the "Volume Viewer" tool. Surface 1092 representations were cleaned up using the "High dust" command, based on size filtering 1093 thresholding.

1094 3D-analysis of HeLa and iNeuron datasets from 3D-SIM datasets was performed 1095 using Imaris (Oxford Instruments, v9.7). After converting all multi-color .tiff to native .ims

1096 files, import into Imaris Arena and global background subtraction, mitochondrial and pUb 1097 objects were segmented from seeds (XY starting diameter:  $0.08 \,\mu\text{m} == \text{pixel size of images})$ , 1098 segmented based on automatic thresholding with local background subtraction and splitting 1099 of touching objects (0.4µm). Objects were piped into Imaris Vantage module for further 1100 analysis. In Vantage, nearest neighbour distances of pUb to pUb and between pUb and 1101 mitochondria, as well as volume of segmented objects were computed. This pipeline was 1102 tested on WT control cells and then applied for batch processing on all other genotype to 1103 allow for unbiased segmentation and analysis. On average, 10 cells were analysed per 1104 genotype per condition (121 total) for HeLa cells, and 12 cells (139 total) for iNeurons.

1105 Unless stated otherwise, all images depicted in figures are maximum-intensity 1106 projections.

1107

#### 1108 Flow cytometry-based measurement of mitophagic flux

1109 HeLa HFT/TO-PRKN cells of the indicated genotype (control, PINK1<sup>-/-,</sup> FBXO7 <sup>-/-</sup>) expressing 1110 mtKeima (Heo et al., 2015) were seeded into 12-well dishes. Upon reaching 60% 1111 confluency, 2 µg/ml doxycycline was added to induce Parkin expression for at least eight 1112 hours before depolarization of mitochondria. Following Parkin induction, cells were treated 1113 with Antimycin A (0.5  $\mu$ M) and Oligomycin (0.5  $\mu$ M) for the indicated number of hours. 1114 Bafilomycin A (25 nM, Sigma-Aldrich, B1793)-treated samples were treated for three hours. 1115 Control cells were fed three hours before experimental manipulation. To harvest cells for 1116 analysis, each well was washed with 1 mL PBS then treated with 100 µL of 0.25% trypsin for 1117 3 min at room temperature, resuspended in 300 µL of DMEM with 10% FBS. 200 µL of each 1118 sample was transferred to a flat-bottom 96-well plate for analysis by flow cytometry.

1119 hESCs were seeded at day 4 into coated 6-well dishes and differentiated. At day 12, 1120 mitophagy was induced with Antimycin A ( $0.5 \mu$ M) and Oligomycin ( $0.5 \mu$ M) or Bafilomycin A (25 nM, Sigma-Aldrich, B1793) for the indicated number of hours. Control cells were fed 2h 1122 before experimental manipulation. Cells were dissociated from the wells using Accutase and 1123 resuspended in 600 µL ND2 medium + 10 µM Y27632 and filtered through a cell strainer cap 1124 tube (Corning, 352235).

1125 Analysis of a population of at least 10,000 cells was performed on an Attune NxT 1126 (Thermo Fisher Scientific) detecting neutral mtKeima signal with excitation at 445 nm and 1127 emission 603 nm with a 48 nm bandpass and acidic mtKeima with 561 nm excitation and 1128 emission 620 nm and a 15 nm band pass. mtKeima ratio was analysed as previously 1129 described (An & Harper, 2018). Briefly, acidic:neutral Keima ratios were measured by gating 1130 a population of well-behaved single cells using forward and side scatter, followed by 1131 calculation of acidic:neutral mtKeima ratio on a per-cell basis in FlowJo Software (FlowJo, 1132 LLC).

1133

# 1134 QUANTIFICATION AND STATISTICAL ANALYSIS

Unless stated otherwise all quantitative experiments were performed in triplicate and
average with S.E.M. or S.D. as indicated in legends reported.

# 1138 **REFERENCES**

- 1139 Al Rawi S, Simpson L, McDonald NQ, Chernuha V, Elpeleg O, Zeviani M, Barker RA, Spiegel R,
- 1140 Laman H (2022) Study of an FBXO7 patient mutation reveals Fbxo7 and PI31 co-regulate
- 1141 proteasomes and mitochondria. *BioRxiv*
- 1142 An H, Harper JW (2018) Systematic analysis of ribophagy in human cells reveals bystander
- 1143 flux during selective autophagy. *Nat Cell Biol* 20: 135-143
- 1144 An H, Ordureau A, Korner M, Paulo JA, Harper JW (2020) Systematic quantitative analysis of
- 1145 ribosome inventory during nutrient stress. *Nature* 583: 303-309
- 1146 Antico O, Ordureau A, Stevens M, Singh F, Nirujogi RS, Gierlinski M, Barini E, Rickwood ML,
- 1147 Prescott A, Toth R et al (2021) Global ubiquitylation analysis of mitochondria in primary

- 1148 neurons identifies endogenous Parkin targets following activation of PINK1. *Sci Adv* 7:
- 1149 eabj0722
- 1150 Bader M, Benjamin S, Wapinski OL, Smith DM, Goldberg AL, Steller H (2011) A conserved F
- box regulatory complex controls proteasome activity in Drosophila. *Cell* 145: 371-382
- 1152 Benjamini Y, Hochberg Y (1995) Controlling the false discovery rate: a practical and powerful
- approach to multiple testing. *Journal of the Royal Statistical Society, Series B* 57: 289-300
- Bingol B, Tea JS, Phu L, Reichelt M, Bakalarski CE, Song Q, Foreman O, Kirkpatrick DS, Sheng
- 1155 M (2014) The mitochondrial deubiquitinase USP30 opposes parkin-mediated mitophagy.
- 1156 Nature 510: 370-375
- 1157 Burchell VS, Nelson DE, Sanchez-Martinez A, Delgado-Camprubi M, Ivatt RM, Pogson JH,
- 1158 Randle SJ, Wray S, Lewis PA, Houlden H et al (2013) The Parkinson's disease-linked proteins
- 1159 Fbxo7 and Parkin interact to mediate mitophagy. *Nat Neurosci* 16: 1257-1265
- 1160 Burman JL, Pickles S, Wang C, Sekine S, Vargas JNS, Zhang Z, Youle AM, Nezich CL, Wu X,
- Hammer JA *et al* (2017) Mitochondrial fission facilitates the selective mitophagy of protein
  aggregates. *J Cell Biol* 216: 3231-3247
- 1163 Chen G, Gulbranson DR, Hou Z, Bolin JM, Ruotti V, Probasco MD, Smuga-Otto K, Howden SE,
- 1164 Diol NR, Propson NE *et al* (2011) Chemically defined conditions for human iPSC derivation
- and culture. *Nat Methods* 8: 424-429
- 1166 Deutsch EW, Bandeira N, Sharma V, Perez-Riverol Y, Carver JJ, Kundu DJ, Garcia-Seisdedos
- 1167 D, Jarnuczak AF, Hewapathirana S, Pullman BS et al (2020) The ProteomeXchange
- consortium in 2020: enabling 'big data' approaches in proteomics. *Nucleic Acids Res* 48:D1145-D1152
- 1170 Di Fonzo A, Dekker MC, Montagna P, Baruzzi A, Yonova EH, Correia Guedes L, Szczerbinska
- 1171 A, Zhao T, Dubbel-Hulsman LO, Wouters CH *et al* (2009) FBXO7 mutations cause autosomal
- 1172 recessive, early-onset parkinsonian-pyramidal syndrome. *Neurology* 72: 240-245
- Dikic I, Elazar Z (2018) Mechanism and medical implications of mammalian autophagy. *Nat Rev Mol Cell Biol* 19: 349-364
- 1175 Elias JE, Gygi SP (2007) Target-decoy search strategy for increased confidence in large-scale
- 1176 protein identifications by mass spectrometry. *Nat Methods* 4: 207-214
- 1177 Eng JK, McCormack AL, Yates JR (1994) An approach to correlate tandem mass spectral data
- of peptides with amino acid sequences in a protein database. *J Am Soc Mass Spectrom* 5:976-989
- 1180 Erickson BK, Mintseris J, Schweppe DK, Navarrete-Perea J, Erickson AR, Nusinow DP, Paulo
- 1181 JA, Gygi SP (2019) Active Instrument Engagement Combined with a Real-Time Database
- Search for Improved Performance of Sample Multiplexing Workflows. *J Proteome Res* 18:1299-1306
- 1184 Evans CS, Holzbaur EL (2020) Degradation of engulfed mitochondria is rate-limiting in
- 1185 Optineurin-mediated mitophagy in neurons. *Elife* 9
- Gladkova C, Maslen SL, Skehel JM, Komander D (2018) Mechanism of parkin activation by
  PINK1. *Nature* 559: 410-414
- 1188 Goodall EA, Kraus F, Harper JW (2022) Mechanisms underlying ubiquitin-driven selective
- 1189 mitochondrial and bacterial autophagy. *Mol Cell* 82: 1501-1513
- 1190 Gustafsson MG, Shao L, Carlton PM, Wang CJ, Golubovskaya IN, Cande WZ, Agard DA, Sedat
- 1191 JW (2008) Three-dimensional resolution doubling in wide-field fluorescence microscopy by
- 1192 structured illumination. *Biophys J* 94: 4957-4970
- 1193 Harper JW, Ordureau A, Heo JM (2018) Building and decoding ubiquitin chains for
- 1194 mitophagy. Nat Rev Mol Cell Biol 19: 93-108

- 1195 Heo JM, Ordureau A, Paulo JA, Rinehart J, Harper JW (2015) The PINK1-PARKIN
- 1196 Mitochondrial Ubiquitylation Pathway Drives a Program of OPTN/NDP52 Recruitment and
- 1197 TBK1 Activation to Promote Mitophagy. *Mol Cell* 60: 7-20
- 1198 Houlden H, Singleton AB (2012) The genetics and neuropathology of Parkinson's disease.
- 1199 Acta Neuropathol 124: 325-338
- Huang T, Fang L, He R, Weng H, Chen X, Ye Q, Qu D (2020) Fbxo7 and Pink1 play a reciprocal role in regulating their protein levels. *Aging (Albany NY)* 13: 77-88
- 1202 Hur JK, Kim K, Been KW, Baek G, Ye S, Hur JW, Ryu SM, Lee YS, Kim JS (2016) Targeted
- mutagenesis in mice by electroporation of Cpf1 ribonucleoproteins. *Nat Biotechnol* 34: 807-808
- 1205 Huttlin EL, Bruckner RJ, Navarrete-Perea J, Cannon JR, Baltier K, Gebreab F, Gygi MP,
- 1206 Thornock A, Zarraga G, Tam S *et al* (2021) Dual proteome-scale networks reveal cell-specific 1207 remodeling of the human interactome. *Cell* 184: 3022-3040 e3028
- 1208 Huttlin EL, Jedrychowski MP, Elias JE, Goswami T, Rad R, Beausoleil SA, Villen J, Haas W,
- 1209 Sowa ME, Gygi SP (2010) A tissue-specific atlas of mouse protein phosphorylation and
- 1210 expression. *Cell* 143: 1174-1189
- 1211 Itzhak DN, Tyanova S, Cox J, Borner GH (2016) Global, quantitative and dynamic mapping of 1212 protein subcellular localization. *Elife* 5
- 1213 Jin J, Cardozo T, Lovering RC, Elledge SJ, Pagano M, Harper JW (2004) Systematic analysis
- 1214 and nomenclature of mammalian F-box proteins. *Genes Dev* 18: 2573-2580
- 1215 Jin SM, Lazarou M, Wang C, Kane LA, Narendra DP, Youle RJ (2010) Mitochondrial
- membrane potential regulates PINK1 import and proteolytic destabilization by PARL. *J Cell Biol* 191: 933-942
- 1218 Kane LA, Lazarou M, Fogel AI, Li Y, Yamano K, Sarraf SA, Banerjee S, Youle RJ (2014) PINK1
- phosphorylates ubiquitin to activate Parkin E3 ubiquitin ligase activity. *J Cell Biol* 205: 143153
- 1221 Katayama H, Kogure T, Mizushima N, Yoshimori T, Miyawaki A (2011) A sensitive and
- quantitative technique for detecting autophagic events based on lysosomal delivery. *ChemBiol* 18: 1042-1052
- 1224 Kazlauskaite A, Martinez-Torres RJ, Wilkie S, Kumar A, Peltier J, Gonzalez A, Johnson C,
- 1225 Zhang J, Hope AG, Peggie M *et al* (2015) Binding to serine 65-phosphorylated ubiquitin
- primes Parkin for optimal PINK1-dependent phosphorylation and activation. *EMBO Rep* 16:939-954
- 1228 Kirk R, Laman H, Knowles PP, Murray-Rust J, Lomonosov M, Meziane el K, McDonald NQ
- 1229 (2008) Structure of a conserved dimerization domain within the F-box protein Fbxo7 and the
- 1230 PI31 proteasome inhibitor. *J Biol Chem* 283: 22325-22335
- 1231 Koyano F, Okatsu K, Kosako H, Tamura Y, Go E, Kimura M, Kimura Y, Tsuchiya H, Yoshihara
- H, Hirokawa T *et al* (2014) Ubiquitin is phosphorylated by PINK1 to activate parkin. *Nature*510: 162-166
- 1234 Kraus F, Miron E, Demmerle J, Chitiashvili T, Budco A, Alle Q, Matsuda A, Leonhardt H,
- 1235 Schermelleh L, Markaki Y (2017) Quantitative 3D structured illumination microscopy of
- 1236 nuclear structures. *Nat Protoc* 12: 1011-1028
- 1237 Labun K, Montague TG, Krause M, Torres Cleuren YN, Tjeldnes H, Valen E (2019) CHOPCHOP
- 1238 v3: expanding the CRISPR web toolbox beyond genome editing. *Nucleic Acids Res* 47: W171-
- 1239 W174
- 1240 Lambert SA, Jolma A, Campitelli LF, Das PK, Yin Y, Albu M, Chen X, Taipale J, Hughes TR,
- 1241 Weirauch MT (2018) The Human Transcription Factors. *Cell* 172: 650-665

1242 Lazarou M, Jin SM, Kane LA, Youle RJ (2012) Role of PINK1 binding to the TOM complex and 1243 alternate intracellular membranes in recruitment and activation of the E3 ligase Parkin. Dev 1244 Cell 22: 320-333 1245 Lazarou M, Sliter DA, Kane LA, Sarraf SA, Wang C, Burman JL, Sideris DP, Fogel AI, Youle RJ 1246 (2015) The ubiquitin kinase PINK1 recruits autophagy receptors to induce mitophagy. 1247 Nature 524: 309-314 1248 Li J, Cai Z, Bomgarden RD, Pike I, Kuhn K, Rogers JC, Roberts TM, Gygi SP, Paulo JA (2021) 1249 TMTpro-18plex: The Expanded and Complete Set of TMTpro Reagents for Sample 1250 Multiplexing. J Proteome Res 20: 2964-2972 1251 Liu K, Jones S, Minis A, Rodriguez J, Molina H, Steller H (2019) PI31 Is an Adaptor Protein for 1252 Proteasome Transport in Axons and Required for Synaptic Development. Dev Cell 50: 509-1253 524 e510 1254 McAlister GC, Nusinow DP, Jedrychowski MP, Wuhr M, Huttlin EL, Erickson BK, Rad R, Haas 1255 W, Gygi SP (2014) MultiNotch MS3 enables accurate, sensitive, and multiplexed detection of 1256 differential expression across cancer cell line proteomes. Anal Chem 86: 7150-7158 1257 McWilliams TG, Mugit MM (2017) PINK1 and Parkin: emerging themes in mitochondrial 1258 homeostasis. Curr Opin Cell Biol 45: 83-91 1259 Narendra D, Tanaka A, Suen DF, Youle RJ (2008) Parkin is recruited selectively to impaired 1260 mitochondria and promotes their autophagy. J Cell Biol 183: 795-803 1261 Narendra DP, Jin SM, Tanaka A, Suen DF, Gautier CA, Shen J, Cookson MR, Youle RJ (2010) 1262 PINK1 is selectively stabilized on impaired mitochondria to activate Parkin. PLoS Biol 8: 1263 e1000298 1264 Ng MYW, Wai T, Simonsen A (2021) Quality control of the mitochondrion. Dev Cell 1265 Okatsu K, Koyano F, Kimura M, Kosako H, Saeki Y, Tanaka K, Matsuda N (2015) 1266 Phosphorylated ubiquitin chain is the genuine Parkin receptor. J Cell Biol 209: 111-128 1267 Ordureau A, Heo JM, Duda DM, Paulo JA, Olszewski JL, Yanishevski D, Rinehart J, Schulman 1268 BA, Harper JW (2015) Defining roles of PARKIN and ubiquitin phosphorylation by PINK1 in 1269 mitochondrial quality control using a ubiquitin replacement strategy. Proc Natl Acad Sci U S 1270 A 112: 6637-6642 1271 Ordureau A, Kraus F, Zhang J, An H, Park S, Ahfeldt T, Paulo JA, Harper JW (2021) Temporal 1272 proteomics during neurogenesis reveals large-scale proteome and organelle remodeling via 1273 selective autophagy. *Mol Cell* 1274 Ordureau A, Paulo JA, Zhang J, An H, Swatek KN, Cannon JR, Wan Q, Komander D, Harper JW 1275 (2020) Global Landscape and Dynamics of Parkin and USP30-Dependent Ubiguitylomes in 1276 iNeurons during Mitophagic Signaling. Mol Cell 77: 1124-1142 e1110 1277 Ordureau A, Paulo JA, Zhang W, Ahfeldt T, Zhang J, Cohn EF, Hou Z, Heo JM, Rubin LL, Sidhu 1278 SS et al (2018) Dynamics of PARKIN-Dependent Mitochondrial Ubiquitylation in Induced 1279 Neurons and Model Systems Revealed by Digital Snapshot Proteomics. Mol Cell 70: 211-227 1280 e218 1281 Ordureau A, Sarraf SA, Duda DM, Heo JM, Jedrychowski MP, Sviderskiy VO, Olszewski JL, 1282 Koerber JT, Xie T, Beausoleil SA et al (2014) Quantitative proteomics reveal a feedforward 1283 mechanism for mitochondrial PARKIN translocation and ubiquitin chain synthesis. Mol Cell 1284 56: 360-375 1285 Paisan-Ruiz C, Guevara R, Federoff M, Hanagasi H, Sina F, Elahi E, Schneider SA, 1286 Schwingenschuh P, Bajaj N, Emre M et al (2010) Early-onset L-dopa-responsive parkinsonism

- 1287 with pyramidal signs due to ATP13A2, PLA2G6, FBXO7 and spatacsin mutations. *Mov Disord*
- 1288 25: 1791-1800

1289 Paulo JA, O'Connell JD, Gygi SP (2016) A Triple Knockout (TKO) Proteomics Standard for

- 1290 Diagnosing Ion Interference in Isobaric Labeling Experiments. *J Am Soc Mass Spectrom* 27:1291 1620-1625
- 1292 Perez-Riverol Y, Bai J, Bandla C, Garcia-Seisdedos D, Hewapathirana S, Kamatchinathan S,
- 1293 Kundu DJ, Prakash A, Frericks-Zipper A, Eisenacher M et al (2022) The PRIDE database
- resources in 2022: a hub for mass spectrometry-based proteomics evidences. *Nucleic Acids Res* 50: D543-D552
- Pickrell AM, Youle RJ (2015) The roles of PINK1, parkin, and mitochondrial fidelity in
  Parkinson's disease. *Neuron* 85: 257-273
- 1298 Plubell DL, Wilmarth PA, Zhao Y, Fenton AM, Minnier J, Reddy AP, Klimek J, Yang X, David LL,
- 1299 Pamir N (2017) Extended Multiplexing of Tandem Mass Tags (TMT) Labeling Reveals Age and
- High Fat Diet Specific Proteome Changes in Mouse Epididymal Adipose Tissue. *Mol Cell Proteomics* 16: 873-890
- 1302 Rad R, Li J, Mintseris J, O'Connell J, Gygi SP, Schweppe DK (2021) Improved Monoisotopic
- 1303 Mass Estimation for Deeper Proteome Coverage. J Proteome Res 20: 591-598
- 1304 Rath S, Sharma R, Gupta R, Ast T, Chan C, Durham TJ, Goodman RP, Grabarek Z, Haas ME,
- 1305 Hung WHW et al (2021) MitoCarta3.0: an updated mitochondrial proteome now with sub-
- 1306 organelle localization and pathway annotations. *Nucleic Acids Res* 49: D1541-D1547
- 1307 Ravenhill BJ, Boyle KB, von Muhlinen N, Ellison CJ, Masson GR, Otten EG, Foeglein A,
- 1308 Williams R, Randow F (2019) The Cargo Receptor NDP52 Initiates Selective Autophagy by
- 1309 Recruiting the ULK Complex to Cytosol-Invading Bacteria. *Mol Cell* 74: 320-329 e326
- 1310 Sarraf SA, Raman M, Guarani-Pereira V, Sowa ME, Huttlin EL, Gygi SP, Harper JW (2013)
- 1311 Landscape of the PARKIN-dependent ubiquitylome in response to mitochondrial
- 1312 depolarization. *Nature* 496: 372-376
- 1313 Sauve V, Sung G, Soya N, Kozlov G, Blaimschein N, Miotto LS, Trempe JF, Lukacs GL, Gehring
- 1314 K (2018) Mechanism of parkin activation by phosphorylation. *Nat Struct Mol Biol* 25: 623-1315 630
- 1316 Savitski MM, Wilhelm M, Hahne H, Kuster B, Bantscheff M (2015) A Scalable Approach for
- 1317 Protein False Discovery Rate Estimation in Large Proteomic Data Sets. *Mol Cell Proteomics*
- 1318 14: 2394-2404
- 1319 Schindelin J, Arganda-Carreras I, Frise E, Kaynig V, Longair M, Pietzsch T, Preibisch S, Rueden
- 1320 C, Saalfeld S, Schmid B *et al* (2012) Fiji: an open-source platform for biological-image
- 1321 analysis. *Nat Methods* 9: 676-682
- 1322 Schweppe DK, Eng JK, Yu Q, Bailey D, Rad R, Navarrete-Perea J, Huttlin EL, Erickson BK, Paulo
- 1323 JA, Gygi SP (2020) Full-Featured, Real-Time Database Searching Platform Enables Fast and
- 1324 Accurate Multiplexed Quantitative Proteomics. *J Proteome Res* 19: 2026-2034
- 1325 Schweppe DK, Prasad S, Belford MW, Navarrete-Perea J, Bailey DJ, Huguet R, Jedrychowski
- 1326 MP, Rad R, McAlister G, Abbatiello SE et al (2019) Characterization and Optimization of
- 1327 Multiplexed Quantitative Analyses Using High-Field Asymmetric-Waveform Ion Mobility
- 1328 Mass Spectrometry. Anal Chem 91: 4010-4016
- 1329 Stirling DR, Swain-Bowden MJ, Lucas AM, Carpenter AE, Cimini BA, Goodman A (2021)
- 1330 CellProfiler 4: improvements in speed, utility and usability. *BMC Bioinformatics* 22: 433
- 1331 Stolz A, Ernst A, Dikic I (2014) Cargo recognition and trafficking in selective autophagy. Nat
- 1332 *Cell Biol* 16: 495-501
- 1333 Tai YC (2022) timecourse: Statistical Analysis for Developmental Microarray Time Course
- 1334 Data. R package version 1.68.0. <u>http://wwwbioconductororg</u>

- 1335 Tyanova S, Cox J (2018) Perseus: A Bioinformatics Platform for Integrative Analysis of
- 1336 Proteomics Data in Cancer Research. *Methods Mol Biol* 1711: 133-148
- 1337 Vargas JNS, Wang C, Bunker E, Hao L, Maric D, Schiavo G, Randow F, Youle RJ (2019)
- 1338 Spatiotemporal Control of ULK1 Activation by NDP52 and TBK1 during Selective Autophagy.
- 1339 *Mol Cell* 74: 347-362 e346
- 1340 Vingill S, Brockelt D, Lancelin C, Tatenhorst L, Dontcheva G, Preisinger C, Schwedhelm-
- 1341 Domeyer N, Joseph S, Mitkovski M, Goebbels S et al (2016) Loss of FBXO7 (PARK15) results
- in reduced proteasome activity and models a parkinsonism-like phenotype in mice. *EMBO J*35: 2008-2025
- 1344 Wang Y, Yang F, Gritsenko MA, Wang Y, Clauss T, Liu T, Shen Y, Monroe ME, Lopez-Ferrer D,
- 1345 Reno T *et al* (2011) Reversed-phase chromatography with multiple fraction concatenation
- 1346 strategy for proteome profiling of human MCF10A cells. *Proteomics* 11: 2019-2026
- 1347 Wauer T, Simicek M, Schubert A, Komander D (2015) Mechanism of phospho-ubiquitin-
- 1348 induced PARKIN activation. *Nature* 524: 370-374
- 1349 Wong YC, Holzbaur EL (2014) Optineurin is an autophagy receptor for damaged
- 1350 mitochondria in parkin-mediated mitophagy that is disrupted by an ALS-linked mutation.
- 1351 Proc Natl Acad Sci U S A 111: E4439-4448
- Yamano K, Youle RJ (2013) PINK1 is degraded through the N-end rule pathway. *Autophagy*9: 1758-1769
- 1354 Zhang Y, Pak C, Han Y, Ahlenius H, Zhang Z, Chanda S, Marro S, Patzke C, Acuna C, Covy J et
- 1355 *al* (2013) Rapid single-step induction of functional neurons from human pluripotent stem 1356 cells. *Neuron* 78: 785-798
- 1357 Zuris JA, Thompson DB, Shu Y, Guilinger JP, Bessen JL, Hu JH, Maeder ML, Joung JK, Chen ZY,
- 1358 Liu DR (2015) Cationic lipid-mediated delivery of proteins enables efficient protein-based
- 1359 genome editing in vitro and in vivo. Nat Biotechnol 33: 73-80
- 1360