1 Genetic Disruption of WASHC4 Drives Endo-lysosomal Dysfunction

2 and Cognitive-Movement Impairments in Mice and Humans

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24 ABSTRACT

25 Mutation of the WASH complex subunit, SWIP, is implicated in human intellectual 26 disability, but the cellular etiology of this association is unknown. We identify the neuronal 27 WASH complex proteome, revealing a network of endosomal proteins. To uncover how 28 dysfunction of endosomal SWIP leads to disease, we generate a mouse model of the 29 human WASHC4^{c.3056C>G} mutation. Quantitative spatial proteomics analysis of 30 SWIP^{P1019R} mouse brain reveals that this mutation destabilizes the WASH complex and 31 uncovers significant perturbations in both endosomal and lysosomal pathways. Cellular 32 and histological analyses confirm that SWIP^{P1019R} results in endo-lysosomal disruption 33 and uncover indicators of neurodegeneration. We find that SWIP^{P1019R} not only impacts 34 cognition, but also causes significant progressive motor deficits in mice. Remarkably, a retrospective analysis of SWIP^{P1019R} patients confirms motor deficits in humans. 35 36 Combined, these findings support the model that WASH complex destabilization, resulting from SWIP^{P1019R}, drives cognitive and motor impairments via endo-lysosomal 37 38 dysfunction in the brain.

39

41 **INTRODUCTION**

42 Neurons maintain precise control of their subcellular proteome using a 43 sophisticated network of vesicular trafficking pathways that shuttle cargo throughout their 44 elaborate processes. Endosomes function as a central hub in this vesicular relay system 45 by coordinating protein sorting between multiple cellular compartments, including surface 46 receptor endocytosis and recycling, as well as degradative shunting to the lysosome. How 47 endosomal trafficking is modulated in neurons remains a vital area of research due to the 48 unique degree of spatial segregation between organelles in neurons, and its strong 49 implication in neurodevelopmental and neurodegenerative diseases.

50 In non-neuronal cells, an evolutionarily conserved complex, the Wiskott-Aldrich 51 Syndrome protein and SCAR Homology (WASH) complex, coordinates endosomal 52 trafficking (Derivery and Gautreau, 2010; Linardopoulou et al., 2007). WASH is composed 53 of five core protein components: WASHC1 (aka WASH1), WASHC2 (aka FAM21), 54 WASHC3 (aka CCDC53), WASHC4 (aka SWIP), and WASHC5 (aka Strumpellin) 55 (encoded by genes Washc1-Washc5, respectively), which are broadly expressed in 56 multiple organ systems (Alekhina et al., 2017; Kustermann et al., 2018; McNally et al., 57 2017; Simonetti and Cullen, 2019; Thul et al., 2017). The WASH complex plays a central 58 role in non-neuronal endosomal trafficking by activating Arp2/3-dependent actin 59 branching at the outer surface of endosomes to influence cargo sorting and vesicular 60 scission (Gomez and Billadeau, 2009; Lee et al., 2016; Phillips-Krawczak et al., 2015; 61 Piotrowski et al., 2013; Simonetti and Cullen, 2019). WASH also interacts with at least 62 three main cargo adaptor complexes — the Retromer, Retriever. and COMMD/CCDC22/CCDC93 (CCC) complexes - all of which associate with distinct 63

64 sorting nexins to select specific cargo and enable their trafficking to other cellular 65 locations (Binda et al., 2019; Farfán et al., 2013; McNally et al., 2017; Phillips-Krawczak 66 et al., 2015; Seaman and Freeman, 2014; Singla et al., 2019). Loss of the WASH complex 67 in non-neuronal cells has detrimental effects on endosomal structure and function, as its 68 loss results in aberrant endosomal tubule elongation and cargo mislocalization (Bartuzi 69 et al., 2016; Derivery et al., 2009; Gomez et al., 2012; Gomez and Billadeau, 2009; 70 Phillips-Krawczak et al., 2015; Piotrowski et al., 2013). However, whether the WASH 71 complex performs an endosomal trafficking role in neurons remains an open question, as 72 no studies have addressed neuronal WASH function to date.

73 Consistent with the association between the endosomal trafficking system and 74 pathology, dominant missense mutations in WASHC5 (protein: Strumpellin) are 75 associated with hereditary spastic paraplegia (SPG8) (De Bot et al., 2013; Valdmanis et 76 al., 2007), and autosomal recessive point mutations in WASHC4 (protein: SWIP) and 77 WASHC5 are associated with syndromic and non-syndromic intellectual disabilities 78 (Assoum et al., 2020; Elliott et al., 2013; Ropers et al., 2011). In particular, an autosomal 79 recessive mutation in WASHC4 (c.3056C>G; p.Pro1019Arg) was identified in a cohort of 80 children with non-syndromic intellectual disability (Ropers et al., 2011). Cell lines derived 81 from these patients exhibited decreased abundance of WASH proteins, leading the authors to hypothesize that the observed cognitive deficits in SWIP^{P1019R} patients resulted 82 83 from disruption of neuronal WASH signaling (Ropers et al., 2011). However, whether this 84 mutation leads to perturbations in neuronal endosomal integrity, or how this might result 85 in cellular changes associated with disease, are unknown.

86 Here we report the analysis of neuronal WASH and its molecular role in disease 87 pathogenesis. We use in vivo proximity proteomics (iBioID) to uncover the neuronal 88 WASH proteome and demonstrate that it is highly enriched for components of endosomal 89 trafficking. We then generate a mouse model of the human WASHC4^{c.3056c>g} mutation 90 (SWIP^{P1019R}) (Ropers et al., 2011) to discover how this mutation may alter neuronal 91 trafficking pathways and test whether it leads to phenotypes congruent with human 92 patients. Using an adapted spatial proteomics approach (Geladaki et al., 2019), coupled 93 with a systems-level analysis of protein covariation networks, we find strong evidence for 94 substantial disruption of neuronal endosomal and lysosomal pathways in vivo. Cellular 95 analyses confirm a significant impact on neuronal endo-lysosomal trafficking in vitro and 96 in vivo, with evidence of lipofuscin accumulation and progressive apoptosis activation, 97 molecular phenotypes that are indicative of neurodegenerative pathology. Behavioral analyses of SWIP^{P1019R} mice at adolescence and adulthood confirm a role of WASH in 98 99 cognitive processes, and reveal profound, progressive motor dysfunction. Importantly, 100 retrospective examination of SWIP^{P1019R} patient data confirms motor dysfunction 101 coincident with cognitive impairments in humans. Our results establish that impaired 102 WASH complex function leads to altered neuronal endo-lysosomal function, which 103 manifests behaviorally as cognitive and movement impairments.

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105 **RESULTS**

Identification of the WASH complex proteome *in vivo* confirms a neuronal role in
 endosomal trafficking. While multiple mutations within the WASH complex have been
 identified in humans (Assoum et al., 2020; Elliott et al., 2013; Ropers et al., 2011;

109 Valdmanis et al., 2007), how these mutations lead to neurological dysfunction remains 110 unknown (Figure 1A). Given that previous work in non-neuronal cultured cells and non-111 mammalian organisms have established that the WASH complex functions in endosomal 112 trafficking, we first aimed to determine whether this role was conserved in the mouse 113 nervous system (Alekhina et al., 2017; Billadeau et al., 2010; Derivery et al., 2009; Gomez 114 et al., 2012; Gomez and Billadeau, 2009). To discover the likely molecular functions of 115 the neuronal WASH complex, we utilized an in vivo BioID (iBioID) paradigm developed in our laboratory to identify the WASH complex proteome from brain tissue (Uezu et al., 116 117 2016). BioID probes were generated by fusing a component of the WASH complex, 118 WASH1 (gene: Washc1), with the promiscuous biotin ligase, BioID2 (WASH1-BioID2, 119 Figure 1B), or by expressing BioID2 alone (negative control, solubleBioID2) under the 120 neuron-specific, human Synapsin-1 promoter (Kim et al., 2016). We injected 121 adenoviruses (AAV) expressing these constructs into the cortex of wild-type postnatal 122 day zero (P0) mice (Figure 1B). Two weeks post-injection, we administered daily 123 subcutaneous biotin for seven days to biotinylate in vivo substrates. The viruses displayed 124 efficient expression and activity in brain tissue, as evidenced by colocalization of the 125 WASH1-BioID2 viral epitope (HA) and biotinylated proteins (Streptavidin) (Figures 1C-F). 126 For label-free quantitative high-mass accuracy LC-MS/MS analyses, whole brain samples 127 were collected at P22, snap-frozen, and processed as previously described (Uezu et al., 128 2016). A total of 2,311 proteins were identified across all three experimental replicates, 129 which were further analyzed for those with significant enrichment in WASH1-BioID2 130 samples over solubleBioID2 negative controls (Table S1).

131 The resulting neuronal WASH proteome included 174 proteins that were 132 significantly enriched (Fold-change \geq 3.0, Benjamini-Hochberg P-Adjust < 0.1, Figure 133 1G). Of these proteins, we identified all five WASH complex components (Figure 1H), as 134 well as 13 previously reported WASH complex interactors (Figure 1I) (McNally et al., 135 2017; Phillips-Krawczak et al., 2015; Simonetti and Cullen, 2019; Singla et al., 2019), 136 which provided strong validity for our proteomic approach and analyses. Additional 137 bioinformatic analyses of the neuronal WASH proteome identified a network of proteins 138 implicated in vesicular trafficking, including 23 proteins enriched for endosomal functions 139 (Figure 1J) and 24 proteins enriched for endocytic functions (Figure 1K). Among these 140 endosomal and endocytic proteins were components of the recently identified endosomal 141 sorting complexes, CCC (CCDC93 and COMMD9) and Retriever (VPS35L) (Phillips-142 Krawczak et al., 2015; Singla et al., 2019), as well as multiple sorting nexins important for 143 recruitment of trafficking regulators to the endosome and cargo selection, such as SNX1-144 3, and SNX16 (Kvainickas et al., 2017; Maruzs et al., 2015; Simonetti et al., 2017). These 145 data demonstrated that the WASH complex interacts with many of the same proteins in 146 neurons as it does in yeast, amoebae, flies, and mammalian cell lines. Furthermore, there 147 were 32 proteins enriched for cytoskeletal regulatory functions (Figure 1L), including 148 actin-modulatory molecules such as the Arp2/3 complex subunit ARPC5, which is 149 consistent with WASH's role in activating this complex to stimulate actin polymerization 150 at endosomes for vesicular scission (Billadeau et al., 2010; Derivery et al., 2009). The 151 WASH1-BioID2 isolated complex also contained 28 proteins known to localize to the 152 excitatory post-synapse (Figure 1M). This included many core synaptic scaffolding 153 proteins, such as SHANK2-3 and DLGAP2-4 (Chen et al., 2011; Mao et al., 2015;

Monteiro and Feng, 2017; Wan et al., 2011), as well as modulators of synaptic receptors such as SYNGAP1 and SHISA6 (Barnett et al., 2006; Clement et al., 2012; Kim et al., 2003; Klaassen et al., 2016), which was consistent with the idea that vesicular trafficking plays an important part in synaptic function and regulation. Taken together, these results support a major endosomal trafficking role of the WASH complex in mouse brain.

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160 SWIP^{P1019R} does not incorporate into the WASH complex, reducing its stability and 161 levels in vivo. To determine how disruption of the WASH complex may lead to disease. 162 we generated a mouse model of a human missense mutation found in children with 163 intellectual disability, WASHC4^{c.3056c>g} (protein: SWIP^{P1019R}) (Ropers et al., 2011). Due to 164 the sequence homology of human and mouse Washc4 genes, we were able to introduce 165 the same point mutation in exon 29 of murine Washc4 using CRISPR (Derivery and 166 Gautreau, 2010; Ropers et al., 2011). This C>G point mutation results in a 167 Proline>Arginine substitution at position 1019 of SWIP's amino acid sequence (Figure 168 2A), a region thought to be critical for its binding to the WASH component, Strumpellin 169 (Jia et al., 2010; Ropers et al., 2011). Western blot analysis of brain lysate from adult homozygous SWIP^{P1019R} mutant mice (referred to from here on as MUT mice) displayed 170 171 significantly decreased abundance of two WASH complex members, Strumpellin and 172 WASH1 (Figure 2B). These results phenocopied data from the human patients (Ropers 173 et al., 2011) and suggested that the WASH complex is unstable in the presence of this 174 SWIP point mutation *in vivo*. To test whether this mutation disrupted interactions between 175 WASH complex subunits, we compared the ability of wild-type SWIP (WT) and SWIP^{P1019R} (MUT) to co-immunoprecipitate with Strumpellin and WASH1 in HEK cells. 176

177 Compared to WT, MUT SWIP co-immunoprecipitated significantly less Strumpellin and 178 WASH1 (IP: 54.8% and 41.4% of WT SWIP, respectively), suggesting that the SWIP^{P1019R} 179 mutation hinders WASH complex formation (Figure 2-figure supplement 1). Together 180 these data support the notion that SWIP^{P1019R} is a damaging mutation that not only 181 impairs its function, but also results in significant reductions of the WASH complex as a 182 whole.

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184 Spatial proteomics and unbiased network covariation analysis reveal significant disruptions in the endo-lysosomal pathway of SWIP^{P1019R} mutant mouse brain. 185 186 Next, we aimed to understand the impact of the SWIP^{P1019R} mutation on the subcellular 187 organization of the mouse brain proteome. We performed spatial proteomics by following 188 the protocol established by Geladaki et al., with modifications for homogenization of brain 189 tissue (Geladaki et al., 2019; Hallett et al., 2008). We isolated seven subcellular fractions 190 from brain tissue and quantified proteins in these samples using 16-plex TMT proteomics. 191 Using this spatial proteomics dataset, we developed a data-driven clustering approach to 192 classify proteins into subcellular compartments. This approach, which differs from the 193 support vector machine learning algorithm employed by Geladaki et al. (2019), was 194 motivated by the lack of a large corpus of brain-specific protein subcellular localization 195 information, and the greater complexity of brain tissue compared to cultured cells. In 196 addition to evaluating differential protein abundance between WT and SWIPP1019R MUT 197 brain, we utilized this spatial proteomics dataset to analyze network-level changes in 198 groups of covarying proteins to better understand WASH's function and explore the 199 cellular mechanisms by which SWIP^{P1019R} causes disease.

200 Brains from 10-month-old mice were gently homogenized to release intact 201 organelles, followed by successive centrifugation steps to enrich subcellular 202 compartments into different fractions based on their density (Figure 2C) (Geladaki et al., 203 2019). Seven WT and seven MUT fractions (each prepared from one brain, 14 samples 204 total) were labeled with unique isobaric tandem-mass tags and concatenated. We also 205 included two sample pooled quality controls (SPQCs), which allowed us to assess 206 experimental variability and perform normalization between experiments. By performing 207 this experiment in triplicate, deep coverage of the mouse brain proteome was obtained-208 across all 48 samples we quantified 86,551 peptides, corresponding to 7,488 proteins. 209 After data pre-processing, normalization, and filtering we retained 5,897 reproducibly 210 quantified proteins in the final dataset (Table S2).

211 We used generalized linear models (GLMs) to assess differential protein 212 abundance for intra-fraction comparisons between WT and MUT genotypes, and for 213 overall comparisons between WT and MUT groups, adjusted for baseline differences in 214 subcellular fraction. In the first analysis, there were 85 proteins with significantly altered 215 abundance in at least one of the 7 subcellular fractions (Benjamini-Hochberg P-Adjust < 216 0.1, Table S2 and Figure 2-figure supplement 2). Five proteins were differentially 217 abundant between WT and MUT in all 7 fractions, including four WASH proteins and 218 RAB21A—a known WASH interactor that functions in early endosomal trafficking 219 (WASHC1, WASHC2, WASHC4, WASHC5, Figure 2E) (Del Olmo et al., 2019; Simpson 220 et al., 2004). The abundance of the remaining WASH complex protein, WASHC3, was 221 found to be very low and was not retained in the final dataset due to its sparse quantification. These data affirm that the SWIPP1019R mutation destabilizes the WASH 222

complex. Next, to evaluate global differences between WT and MUT brain, we analyzed
the average effect of genotype on protein abundance across all fractions. At this level,
there were 687 differentially abundant proteins between WT and MUT brain (Bonferroni
P-Adjust < 0.05) (Table S2). We then aimed to place these differentially abundant proteins
into a more meaningful biological context using a systems-based approach.

228 For network-based analyses, we clustered the protein covariation network defined 229 by pairwise correlations between all 5,897 proteins. Our data-driven, quality-based 230 approach used Network Enhancement (Wang et al., 2018) to remove biological noise 231 from the covariation network and employed the Leiden algorithm (Traag et al., 2019) to 232 identify optimal partitions of the graph. We enforced module guality by permutation testing 233 (Ritchie et al., 2016) to ensure that identified modules exhibited a non-random topology. 234 Clustering of the protein covariation graph identified 255 modules of proteins that strongly 235 covaried together (see Methods for complete description of clustering approach).

236 To test for module-level differences between WT and MUT brain, we summarized 237 modules for each biological replicate (a single subcellular fraction prepared from either a 238 WT or MUT mouse) as the sum of their proteins, and extended our GLM framework to 239 identify changes in module abundance (adjusted for fraction differences) between 240 genotypes. 37 of the 255 modules exhibited significant differences in WT versus MUT 241 brain (Bonferroni P-Adjust < 0.05; Table S3). Of note, the module containing the WASH 242 complex, M19, was predicted to have endosomal function by annotation of protein 243 function, and was enriched for proteins identified by WASH1-BioID2 (hypergeometric test 244 P-Adjust < 0.05, bold node edges, Figure 2D). Similar to the WASH iBioID proteome 245 (Figure 1), M19 contained components of the CCC (CCDC22, CCDC93, COMMD1-3,

246 COMMD6-7, and COMMD9) and Retriever sorting complexes (VPS26C and VPS35L), 247 but not the Retromer sorting complex, suggesting that in the brain, the WASH complex 248 may not interact as closely with Retromer as it does in other cells (Figure 2D). Across all 249 fractions, the abundance of M19 was significantly lower in MUT brain compared to WT, 250 providing evidence that the SWIP^{P1019R} mutation reduces the stability of this protein 251 subnetwork and impairs its function (Figure 2F-G).

252 In contrast to the decreased abundance of the WASH complex/endosome module, 253 M19, we observed three modules (M2, M159, and M213) which were enriched for 254 lysosomal protein components (Geladaki et al., 2019), and exhibited increased 255 abundance in MUT brain (Figure 3). M159 (Figure 3B) contained the lysosomal protease 256 Cathepsin A (CTSA), while M213 (Figure 3D) contained Cathepsin B (CTSB), as well as 257 two key lysosomal hydrolases GLB1 and MAN2B2, and M2 (Figure 3C) contained two 258 Cathepsins (CTSS and CTSL) and several lysosomal hydrolases (e.g. GNS, GLA, and 259 MAN2B1) (Eng and Desnick, 1994; Mayor et al., 1993; Mok et al., 2003; Moon et al., 260 2016; Patel et al., 2018; Regier and Tifft, 1993; Rosenbaum et al., 2014). Notably, M2 261 also contained the lysosomal glycoprotein progranulin (GRN), which is integral to proper 262 lysosome function and whose loss is widely linked with neurodegenerative pathologies 263 (Baker et al., 2006; Pottier et al., 2016; Tanaka et al., 2017; Zhou et al., 2018). In addition, 264 M2 contained the hydrolase IDS, whose loss causes a lysosomal storage disorder that 265 can present with neurological symptoms (Hopwood et al., 1993; Schröder et al., 1994). 266 The overall increase in abundance of modules M2, M159, and M213, and these key 267 lysosomal proteins (Figure 3E-G), may therefore reflect an increase in flux through degradative lysosomal pathways in SWIP^{P1019R} brain. 268

269 Furthermore, Module 2 (Figure 3C) included multiple membrane proteins and 270 extracellular proteins, such as ITGA5 (an integrin shown to be upregulated and 271 redistributed upon loss of WASH1), ATP13A2 (a cation transporter whose loss causes a 272 Parkinsonian syndrome), and MMP17 (an extracellular metalloprotease), suggesting a 273 link between these proteins and lysosomal enzymatic function (English et al., 2000; 274 Ramirez et al., 2006; Zech et al., 2011). Increased abundance of these M2 proteins in 275 MUT brain may indicate that WASH complex disruption alters their cellular localization. 276 Taken together, these changes appear to reflect a pathological condition characterized 277 by distorted lysosomal metabolism and altered cellular trafficking.

278 In addition to these endo-lysosomal changes, network alterations were evident for 279 an endoplasmic reticulum (ER) module (M83), supporting a shift in the proteostasis of 280 mutant neurons (Figure 2-figure supplement 3B). Notably, within the ER module, M83, 281 there was increased abundance of chaperones (e.g. HSPA5, PDIA3, PDIA4, PDIA6, and 282 DNAJC3) that are commonly engaged in presence of misfolded proteins (Bartels et al., 283 2019; Kim et al., 2020; Montibeller and de Belleroche, 2018; Synofzik et al., 2014; Wang 284 et al., 2016). This elevation of ER stress modulators can be indicative of 285 neurodegenerative states, in which the unfolded protein response (UPR) is activated to 286 resolve misfolded species (Garcia-Huerta et al., 2016; Hetz and Saxena, 2017). These 287 data demonstrate that loss of WASH function not only alters endo-lysosomal trafficking, 288 but also causes increased stress on cellular homeostasis.

Finally, besides these endo-lysosomal and homeostatic changes, we also observed two synaptic modules (M35 and M248) that were reduced in MUT brain (Figure 291 2-figure supplement 3C-D). These included mostly excitatory post-synaptic proteins such

as HOMER2 and DLG4 (also identified in WASH1-BioID, Figure 1), consistent with
 endosomal WASH influencing synaptic regulation. Decreased abundance of these
 modules indicates that loss of the WASH complex may result in failure of these proteins
 to be properly trafficked to the synapse.

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297 Mutant neurons display structural abnormalities in endo-lysosomal compartments 298 in vitro. Combined, the proteomics data strongly suggested that endo-lysosomal 299 pathways are altered in adult SWIP^{P1019R} mutant mouse brain. Next, we analyzed whether 300 structural changes in this system were evident in primary neurons. Cortical neurons from 301 littermate WT and MUT P0 pups were cultured for 15 days in vitro (DIV15, Figure 4A), 302 then fixed and stained for established markers of early endosomes (Early Endosome 303 Antigen 1, EEA1; Figures 4B and 4C) and lysosomes (Cathepsin D, CathD; Figures 4D 304 and 4E). Reconstructed three-dimensional volumes of EEA1 and Cathepsin D puncta 305 revealed that MUT neurons display larger EEA1+ somatic puncta than WT neurons 306 (Figures 4G and 4J), but no difference in the total number of EEA1+ puncta (Figure 4F). 307 This finding is consistent with a loss-of-function mutation, as loss of WASH activity 308 prevents cargo scission from endosomes and leads to cargo accumulation (Bartuzi et al., 309 2016; Gomez et al., 2012). Conversely, MUT neurons exhibited significantly less 310 Cathepsin D+ puncta than WT neurons (Figure 4H), but the remaining puncta were 311 significantly larger than those of WT neurons (Figures 4I and 4K). These data support the 312 finding that the SWIP^{P1019R} mutation results in both molecular and morphological 313 abnormalities in the endo-lysosomal pathway.

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315 SWIP^{P1019R} mutant brains exhibit markers of abnormal endo-lysosomal structures

316 and cell death in vivo. As there is strong evidence that dysfunctional endo-lysosomal 317 trafficking and elevated ER stress are associated with neurodegenerative disorders, 318 adolescent (P42) and adult (10 month-old, 10mo) WT and MUT brain tissue were 319 analyzed for the presence of cleaved caspase-3, a marker of apoptotic pathway 320 activation, in four brain regions (Boatright and Salvesen, 2003; Porter and Jänicke, 1999). 321 Very little cleaved caspase-3 staining was present in WT and MUT mice at adolescence 322 (Figures 5A, 5B, and Figure 5-figure supplement 1). However, at 10mo, the MUT motor 323 cortices displayed significantly greater cleaved caspsase-3 staining compared to age-324 matched WT littermate controls (Figures 5D, 5E, and 5H). Furthermore, this difference 325 appeared to be selective for the motor cortex, as we did not observe significant 326 differences in cleaved caspase-3 staining at either age for hippocampal, striatal, or 327 cerebellar regions (Figure 5-figure supplement 1). These data suggested that neurons of the motor cortex were particularly susceptible to disruption of endo-lysosomal pathways 328 329 downstream of SWIP^{P109R}, perhaps because long-range corticospinal projections require 330 high fidelity of trafficking pathways (Blackstone et al., 2011; Slosarek et al., 2018; Wang 331 et al., 2014).

To further examine the morphology of primary motor cortex neurons at a subcellular resolution, samples from age-matched 7-month-old WT and MUT mice (7mo, animals each) were imaged by transmission electron microscopy (TEM). Strikingly, we observed large electron-dense inclusions in the cell bodies of MUT neurons (arrows, Figure 5L; pseudo-colored region, 5N). These dense structures were associated electronlucent lipid-like inclusions (asterisk, Figure 5N), and were visually consistent with

338 lipofuscin accumulation at lysosomal residual bodies (Poët et al., 2006; Valdez et al., 339 2017; Yoshikawa et al., 2002). Lipofuscin is a by-product of lysosomal breakdown of 340 lipids, proteins, and carbohydrates, which naturally accumulates over time in non-dividing 341 cells such as neurons (Höhn and Grune, 2013; Moreno-García et al., 2018; Terman and 342 Brunk, 1998). However, excessive lipofuscin accumulation is thought to be detrimental to 343 cellular homeostasis by inhibiting lysosomal function and promoting oxidative stress, 344 often leading to cell death (Brunk and Terman, 2002; Powell et al., 2005). As a result, 345 elevated lipofuscin is considered a biomarker of neurodegenerative disorders, including 346 Alzheimer's disease, Parkinson's disease, and Neuronal Ceroid Lipofuscinoses (Moreno-347 García et al., 2018). Therefore, the marked increase in lipofuscin area and number seen 348 in MUT electron micrographs (Figures 50 and 5P, respectively) is consistent with the 349 increased abundance of lysosomal pathways observed by proteomics, and likely reflects 350 an increase in lysosomal breakdown of cellular material. Together these data indicate that SWIPP1019R results in pathological lysosomal function that could lead to 351 352 neurodegeneration.

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SWIP^{P1019R} mutant mice display persistent deficits in cued fear memory recall. To observe the functional consequences of the SWIP^{P1019R} mutation, we next studied WT and MUT mouse behavior. Given that children with homozygous SWIP^{P1019R} point mutations display intellectual disability (Ropers et al., 2011) and SWIP^{P1019R} mutant mice exhibit endo-lysosomal disruptions implicated in neurodegenerative processes, behavior was assessed at two ages: adolescence (P40-50), and mid-late adulthood (5.5-6.5 mo). Interestingly, MUT mice performed equivalently to WT mice in episodic and working

361 memory paradigms, including novel object recognition and Y-maze alternations (Figure 362 6-figure supplement 1). However, in a fear conditioning task, MUT mice displayed a 363 significant deficit in cued fear memory (Figure 6). This task tests the ability of a mouse to 364 associate an aversive event (a mild electric footshock) with a paired tone (Figure 6A). 365 Freezing behavior of mice during tone presentation is attributed to hippocampal or 366 amygdala-based fear memory processes (Goosens and Maren, 2001; Maren and Holt, 367 2000; Vazdarjanova and McGaugh, 1998). Forty-eight hours after exposure to the paired 368 tone and footshock, MUT mice showed a significant decrease in conditioned freezing to 369 tone presentation compared to their WT littermates (Figures 6B and 6C). To ensure that 370 this difference was not due to altered sensory capacities of MUT mice, we measured the 371 startle response of mice to both electric foot shock and presented tones. In line with intact 372 sensation, MUT mice responded comparably to WT mice in these tests (Figure 6-figure 373 supplement 2). These data demonstrate that although MUT mice perceive footshock 374 sensations and auditory cues, it is their memory of these paired events that is significantly 375 impaired. Additionally, this deficit in fear response was evident at both adolescence and 376 adulthood (top panels, and bottom panels, respectively, Figures 6B and 6C). These changes are consistent with the hypothesis that SWIP^{P109R} is the cause of cognitive 377 378 impairments in humans.

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380 SWIP^{P1019R} mutant mice exhibit surprising motor deficits that are confirmed in 381 human patients. Because SWIP^{P1019R} results in endo-lysosomal pathology consistent 382 with neurodegenerative disorders in the motor cortex, we next analyzed motor function of 383 the mice over time. First, we tested the ability of WT and MUT mice to remain on a rotating

rod for five minutes (Rotarod, Figures 7A-7C). At both adolescence and adulthood, MUT mice performed markedly worse than WT littermate controls (Fig 7C). Mouse performance was not significantly different across trials, which suggested that this difference in retention time was not due to progressive fatigue, but more likely due to an overall difference in motor control (Mann and Chesselet, 2015).

389 To study the animals' movement at a finer scale, the gait of WT and MUT mice 390 was also analyzed using a TreadScan system containing a high-speed camera coupled 391 to a transparent treadmill (Figure 7D) (Beare et al., 2009). Interestingly, while the gait 392 parameters of mice were largely indistinguishable across genotypes at adolescence, a 393 striking difference was seen when the same mice were aged to adulthood (Figures 7E-394 7G). In particular, MUT mice took slower (Figure 7E), longer strides (Figure 7F), stepping 395 closer to the midline of their body (track width, Figure 7- figure supplement 1), and their 396 gait symmetry was altered so that their strides were no longer perfectly out of phase (out 397 of phase=0.5, Figure 7G). While these differences were most pronounced in the rear 398 limbs (as depicted in Figure 7E-7G), the same trends were present in front limbs (Figure 399 7-figure supplement 1). These findings demonstrate that SWIP^{P1019R} results in 400 progressive motor function decline that was detectable by the rotarod task at 401 adolescence, but which became more prominent with age, as both gait and strength 402 functions deteriorated.

These marked motor findings prompted us to re-evaluate the original reports of human SWIP^{P1019R} patients (Ropers et al., 2011). While developmental delay or learning difficulties were the primary impetus for medical evaluation, all patients also exhibited motor symptoms (mean age = 10.4 years old, Figure 7H). The patients' movements were

407 described as "clumsy" with notable fine motor difficulties, dysmetria, dysdiadochokinesia, 408 and mild dysarthria on clinical exam (Figure 7H). Recent communication with the parents 409 of these patients, who are now an average of 21 years old, revealed no notable symptom exacerbation. It is therefore possible that the SWIP^{P1019R} mouse model either exhibits 410 411 differences from human patients or may predict future disease progression for these 412 individuals, given that we observed significant worsening at 5-6 months old in mice (which 413 is thought to be equivalent to ~30-35 years old in humans) (Dutta and Sengupta, 2016; 414 Zhang et al., 2019).

415

416 **DISSCUSSION**

417 Taken together, the data presented here support a mechanistic model whereby SWIP^{P1019R} causes a loss of WASH complex function, resulting in endo-lysosomal 418 419 disruption and accumulation of neurodegenerative markers, such as upregulation of 420 unfolded protein response modulators and lysosomal enzymes, as well as build-up of 421 lipofuscin and cleaved caspase-3 over time. To our knowledge, this study provides the 422 first mechanistic evidence of WASH complex impairment having direct and indirect 423 organellar effects that lead to cognitive deficits and progressive motor impairments 424 (Figure 8).

Using *in vivo* proximity-based proteomics in wild-type mouse brain, we identify that the WASH complex interacts with the CCC (COMMD9 and CCDC93) and Retriever (VPS35L) cargo selective complexes (Bartuzi et al., 2016; Singla et al., 2019). Interestingly, we did not find significant enrichment of the Retromer sorting complex, a well-known WASH interactor, suggesting that it may play a minor role in neuronal WASH-

430 mediated cargo sorting (Figure 1). These data are supported by our TMT proteomics and covariation network analyses of SWIP^{P1019R} mutant brain, which clustered the WASH, 431 432 CCC, and Retriever complexes together in M19, but not the Retromer complex, which 433 was found in endosomal module M14 (Figure 2 and Figure 2-figure supplement 3A). 434 Systems-level protein covariation analyses also revealed that disruption of these WASH-435 CCC-Retriever interactions may have multiple downstream effects on the endosomal 436 machinery, since endosomal modules displayed significant changes in SWIP^{P1019R} brain 437 (including both M19, Figure 2, as well as M14, Figure 2-figure supplement 3A), with 438 corresponding decreases in the abundance of endosomal proteins including Retromer 439 subunits (VPS29 and VPS35), associated sorting nexins (e.g. SNX17 and SNX27), 440 known WASH interactors (e.g. RAB21 and FKBP15), and cargos (e.g. LRP1 and ITGA3) 441 (Figure 2-figure supplements 2 and 3) (Del Olmo et al., 2019; Farfán et al., 2013; 442 Fedoseienko et al., 2018; Halff et al., 2019; Harbour et al., 2012; McNally et al., 2017; Pan et al., 2010; Ye et al., 2020; Zimprich et al., 2011). While previous studies have 443 444 indicated that Retromer and CCC influence endosomal localization of WASH (Harbour et 445 al., 2012; Phillips-Krawczak et al., 2015; Singla et al., 2019), our findings of altered 446 endosomal networks containing decreased Retromer, Retriever, and CCC protein levels in SWIPP1019R mutant brain point to a possible feedback mechanism wherein WASH 447 448 impacts the protein abundance and/or stability of these interactors. Future studies 449 defining the hierarchical interplay between the WASH, Retromer, Retriever, and CCC 450 complexes in neurons could provide clarity on how these mechanisms are organized.

451 In addition to highlighting the neuronal roles of WASH in CCC- and Retriever-452 mediated endosomal sorting, our proteomics approach also identified protein modules

with increased abundance in SWIP^{P1019R} mutant brain. The proteins in these modules fell 453 454 into two interesting categories: lysosomal enzymes and proteins involved in the 455 endoplasmic reticulum (ER) stress response. Of note, some of the lysosomal enzymes 456 with elevated levels in MUT brain (GRN, M2; IDS, M2; and GNS, M213; Figure 3) are 457 also implicated in lysosomal storage disorders, where they generally have decreased, 458 rather than increased, function or expression (Hopwood et al., 1993; Mok et al., 2003; 459 Schröder et al., 1994; Ward et al., 2017). We speculate that loss of WASH function in our 460 mutant mouse model may lead to increased accumulation of cargo and associated 461 machinery at early endosomes (as seen in Figure 4, enlarged EEA1⁺ puncta), eventually 462 overburdening early endosomal vesicles and triggering transition to late endosomes for 463 subsequent fusion with degradative lysosomes (Figure 8). This would effectively increase 464 delivery of endosomal substrates to the lysosome compared to baseline, resulting in 465 enlarged, overloaded lysosomal structures, and elevated demand for degradative 466 enzymes. For example, since mutant neurons display increased lysosomal module 467 protein abundance (Figure 3), and larger lysosomal structures (Figures 4 and 5), they 468 may require higher quantities of progranulin (GRN, M2; Figure 3) for sufficient lysosomal 469 acidification (Tanaka et al., 2017).

Our findings that SWIP^{P1019R} results in reduced WASH complex stability and function, which may ultimately drive lysosomal dysfunction, are supported by studies in non-mammalian cells. For example, expression of a dominant-negative form of WASH1 in amoebae impairs recycling of lysosomal V-ATPases (Carnell et al., 2011) and loss of WASH in *Drosophila* plasmocytes affects lysosomal acidification (Gomez et al., 2012; Nagel et al., 2017; Zech et al., 2011). Moreover, mouse embryonic fibroblasts lacking

WASH1 display abnormal lysosomal morphologies, akin to the structures we observed in
cultured SWIP^{P1019R} MUT neurons (Gomez et al., 2012).

478 In addition to lysosomal dysfunction, endoplasmic reticulum (ER) stress is 479 commonly observed in neurodegenerative states, where accumulation of misfolded 480 proteins disrupts cellular proteostasis (Cai et al., 2016; Hetz and Saxena, 2017; 481 Montibeller and de Belleroche, 2018). This cellular strain triggers the adaptive unfolded 482 protein response (UPR), which attempts to restore cellular homeostasis by increasing the 483 cell's capacity to retain misfolded proteins within the ER, remedy misfolded substrates, 484 and trigger degradation of persistently misfolded species. Involved in this process are ER chaperones that we identified as increased in SWIPP1019R mutant brain including BiP 485 486 (HSPA5), calreticulin (CALR), calnexin (CANX), and the protein disulfide isomerase family members (PDIA1, PDIA4, PDIA6) (M83; Figure 2-supplement 3B) (Garcia-Huerta 487 488 et al., 2016). Many of these proteins were identified in the ER protein module found to be 489 significantly altered in MUT mouse brain (M83), supporting a network-level change in the 490 ER stress response (Figure 2-supplement 3B). One notable exception to this trend was 491 endoplasmin (HSP90B1, M136), which exhibited significantly decreased abundance in 492 SWIP^{P1019R} mutant brain (Table S2). This is surprising given that endoplasmin has been 493 shown to coordinate with BiP in protein folding (Sun et al., 2019), however it may highlight 494 a possible compensatory mechanism. Additionally, prolonged UPR can stimulate 495 autophagic pathways in neurons, where misfolded substrates are delivered to the 496 lysosome for degradation (Cai et al., 2016). These data highlight a relationship between 497 ER and endo-lysosomal disturbances as an exciting avenue for future research.

498 Strikingly, we observed modules enriched for resident proteins corresponding to 499 all 10 of the major subcellular compartments mapped by Geladaki et al. (2019; nucleus, 500 mitochondria, golgi, ER, peroxisome, proteasome, plasma membrane, lysosome, 501 cytoplasm, and ribosome; Supplementary File 1). The greatest dysregulations we 502 observed were in lysosomal, endosomal, ER, and synaptic modules, supporting the 503 hypothesis that SWIP^{P1019R} primarily results in disrupted endo-lysosomal trafficking. While 504 analysis of these dysregulated modules informs the pathobiology of SWIP^{P1019R}, our 505 spatial proteomics approach also identified numerous biologically cohesive modules, 506 which remained unaltered (Supplementary File 1). Given that many of these modules 507 contained proteins of unknown function, we anticipate that future analyses of these 508 modules and their protein constituents have great potential to inform our understanding 509 of protein networks and their influence on neuronal cell biology.

510 It has become clear that preservation of the endo-lysosomal system is critical to 511 neuronal function, as mutations in mediators of this process are implicated in neurological 512 diseases such as Parkinson's disease, Huntington's disease, Alzheimer's disease, 513 Frontotemporal Dementia, Neuronal Ceroid Lipofuscinoses (NCLs), and Hereditary 514 Spastic Paraplegia (Baker et al., 2006; Connor-Robson et al., 2019; Edvardson et al., 515 2012; Follett et al., 2019; Harold et al., 2009; Mukherjee et al., 2019; Pal et al., 2006; 516 Quadri et al., 2013; Seshadri et al., 2010; Tachibana et al., 2019; Valdmanis et al., 2007). 517 These genetic links to predominantly neurodegenerative conditions have supported the 518 proposition that loss of endo-lysosomal integrity can have compounding effects over time 519 and contribute to progressive disease pathologies. In particular, NCLs are lysosomal 520 storage disorders primarily found in children, with heterogenous presentations and

521 multigenic causations (Mukherjee et al., 2019). The majority of genes implicated in NCLs 522 affect lysosomal enzymatic function or transport of proteins to the lysosome (Mukherjee 523 et al., 2019; Poët et al., 2006; Ramirez-Montealegre and Pearce, 2005; Yoshikawa et al., 524 2002). Most patients present with marked neurological impairments, such as learning 525 disabilities, motor abnormalities, vision loss, and seizures, and have the unifying feature 526 of lysosomal lipofuscin accumulation upon pathological examination (Mukherjee et al., 527 2019). While the human SWIP^{P1019R} mutation has not been classified as an NCL (Ropers 528 et al., 2011), findings from our mutant mouse model suggest that loss of WASH complex 529 function leads to phenotypes bearing strong resemblance to NCLs, including lipofuscin 530 accumulation (Figures 4-7). As a result, our mouse model could provide the opportunity 531 to study these pathologies at a mechanistic level, while also enabling preclinical 532 development of treatments for their human counterparts.

533 Currently there is an urgent need for greater mechanistic investigations of 534 neurodegenerative disorders, particularly in the domain of endo-lysosomal trafficking. 535 Despite the continual increase in identification of human disease-associated genes, our 536 molecular understanding of how their protein equivalents function and contribute to 537 pathogenesis remains limited. Here we employ a systems-level analysis of proteomic 538 datasets to uncover biological perturbations linked to SWIP^{P1019R}. We demonstrate the power of combining in vivo proteomics and systems network analyses with in vitro and in 539 540 vivo functional studies to uncover relationships between genetic mutations and molecular 541 disease pathologies. Applying this platform to study organellar dysfunction in other 542 neurodegenerative and neurodevelopmental disorders may facilitate the identification of 543 convergent disease pathways driving brain disorders.





WASH interactors (n=13)

WASH complex (n=5)

J Endosomal trafficking (n=23)



Κ Endocytic function (n=24)



L Cytoskeletal regulation (n=32)



Μ Synaptic regulation (n=28)



G Neuronal WASH proteome (n=174) DLGAI NECTIN смт:

RÚVB UVBI VASHC5 (BP ASHC ГІР1 WASHC1 тмл ΤN PP1R SNX HSPE1 XNI **P**2 WASH iBioID Network Attributes

🔵 Node size: fold enrichment **BioID2** interaction ---- Protein-protein interaction

Figure 1. Identification of the WASH complex proteome *in vivo* confirms a neuronal role in endosomal trafficking

(A) The WASH complex is composed of five subunits, *Washc1* (WASH1), *Washc2* (FAM21), *Washc3* (CCDC53), *Washc4* (SWIP), and *Washc5* (Strumpellin). Human mutations in these components are associated with spastic paraplegia (De Bot et al., 2013; Jahic et al., 2015; Valdmanis et al., 2007), Ritscher-Schinzel Syndrome (Elliott et al., 2013), and intellectual disability (Assoum et al., 2020; Ropers et al., 2011).

(B) A BioID2 probe was attached to the c-terminus of WASH1 and expressed under the human synapsin-1 (hSyn1) promoter in an AAV construct for in vivo BioID (iBioID). iBioID probes (WASH1-BioID2-HA, or negative control solubleBioID2-HA) were injected into wild-type mouse brain at P0 and allowed to express for two weeks. Subcutaneous biotin injections (24 mg/kg) were administered over seven days for biotinylation, and then brains were harvested for isolation and purification of biotinylated proteins. LC-MS/MS identified proteins significantly enriched in all three replicates of WASH1-BioID2 samples over soluble-BioID2 controls.

(C) Representative image of WASH1-BioID2-HA expression in a mouse coronal brain section (Cx = cortex, Hipp = hippocampus, Thal = thalmus). Scale bar, 1 mm.

(D) Representative image of WASH1-BioID2-HA expression in mouse cortex (inset from C). Individual panels show nuclei (DAPI, blue), AAV construct HA epitope (green), and biotinylated proteins

(Streptavidin, red). Merged image shows colocalization of HA and Streptavidin (yellow). Scale bar, 50 μm. (E) Representative image of WASH1-BioID2-HA expression in mouse hippocampus (inset from C). Scale bar, 50 μm.

(F) Representative image of WASH1-BioID2-HA expression in mouse thalamus (inset from C). Scale bar, 50 μm.

(G) iBioID identified known and unknown proteins interactors of the WASH complex in murine neurons. Nodes size represents protein abundance fold-enrichment over negative control (range: 3 to 181.7), solid grey edges delineate iBioID interactions between the WASHC1 probe (seen in yellow at the center) and identified proteins, dashed edges indicate known protein-protein interactions from HitPredict database (López et al., 2015). (H-I) Clustergrams of:

(H) All five WASH complex proteins identified by iBioID.

(I) Previously reported WASH interactors (13/174), including the CCC and Retriever complexes.

(J) Endosomal trafficking proteins (23/174 proteins).

(K) Endocytic proteins (24/174).

(L) Proteins involved in cytoskeletal regulation (32/174), including Arp2/3 subunit ARPC5.

(M)Synaptic proteins (28/174). Clustergrams were annotated by hand and cross-referenced with Metascape (Zhou et al., 2019) GO enrichment of WASH1 proteome constituents over all proteins identified in the BioID experiment.





Figure 2. Spatial proteomics and network covariation analysis reveal significant disruptions to the WASH complex and an endosomal module in SWIP^{P1019R} mutant mouse brain

(A) Mouse model of the human SWIP^{P1019R} missense mutation created using CRISPR. A C>G point mutation was introduced into exon29 of murine $Y \approx @$, leading to a P1019R amino acid substitution. We hypothesize (H₁) that this mutation causes instability of the WASH complex.

(B) Representative western blot and quantification of WASH components, Strumpellin and WASH1 (predicted sizes in kDa: 134 and 72, respectively), as well as loading control β -Tubulin (55kDa) from whole adult whole brain lysate prepared from SWIP WT (Y **æ @** $I^{C/C}$) and SWIP homozygous MUT (Y **æ @** $I^{G/G}$) mice. Bar plots show quantification of band intensities relative to WT (n=3 mice per genotype). Strumpellin (WT 100.0 ± 5.2%, MUT 3.5 ± 0.7%, t_{2.1}=18.44, p=0.0024) and WASH1 (WT 100.0 ± 3.8%, MUT 1.1 ± 0.4%, t_{2.1}=25.92, p=0.0013) were significantly decreased. Equivalent amounts of protein were analyzed in each condition (β -Tubulin: WT 100.0 ± 8.2%, MUT 94.1 ± 4.1%, U=4, p>0.99).

(C) Spatial TMT proteomics experimental design. 7 subcellular fractions were prepared from one WT and one MUT mouse (10mo). These samples, as well as two pooled quality control (QC) samples, were labeled with unique TMT tags and concatenated for simultaneous LC-MS/MS analysis. This experiment was repeated three times (3 WT and 3 MUT brains total). To detect network-level changes, proteins were clustered into modules, and general linearized models (GLMs) were used to identify differences in module abundance between WT and MUT samples. The network shows an overview of the spatial proteomics graph in which the 37 differentially abundant modules are indicated by colored nodes.

(D) Protein module 19 (M19) contains subunits of the WASH, CCC, and Retriever complexes. Node size denotes its weighted degree centrality (~importance in module); purple node color indicates proteins with altered abundance in MUT brain relative to WT; black node border denotes proteins identified in the WASH1-BioID proteome (Figure 1); red, yellow, and green borders highlight protein components of the CCC, Retriever, and WASH complexes; black edges indicate known protein-protein interactions; and grey-red edges denote the relative strength of protein covariation within a module (gray = weak, red = strong). P-adjust values represent enrichment of proteins identified in the CORUM database adjusted for multiple comparisons (Giurgiu et al., 2019).

(E) Difference in normalized protein abundance for four WASH proteins found in M19 (SWIP: WT 6.28 \pm 0.41, MUT 4.89 \pm 0.28, p=3.98x10⁻²⁸; WASH1: WT 6.41 \pm 0.47, MUT 4.65 \pm 0.62, p=7.92x10⁻¹⁸; FAM21: WT 6.29 \pm 0.48, MUT 5.29 \pm 0.49, p=3.27x10⁻¹⁶; Strumpellin: WT 7.85 \pm 0.52, MUT 6.59 \pm 0.53, p=9.06x10⁻²⁵) and one control (Tubulin 4a: WT 8.57 \pm 0.52, MUT 8.52 \pm 0.58, p>0.99) across all three experimental replicates (n=3 independent experiments), presented as log₂(adjusted protein intensities). (F) Normalized average intensities for every protein within M19 across all seven subcellular fractions analyzed. Teal lines delineate protein levels in WT samples, purple lines delineate protein levels in MUT samples (averaged across three experimental replicates). Bolded lines demarcate the fitted intensity values for WT and MUT proteins (n=3 independent experiments).

(G) Difference in M19 abundance, adjusted for fraction differences and presented as log_2 (adjusted module abundance) (WT 13.21 ± 0.003, MUT 12.99 ± 0.003, p=0.0007; n=3 independent experiments).

Figure 3



Figure 3. Disruption of lysosomal protein networks in SWIP^{P1019R} mutant brain

(A) Simplified schematic of the endo-lysosomal pathway in neurons. Inset depicts representative lysosomal enzymes, such as proteases (CTSA, CTSB, CTSL), glycosidases (GLA, GLB1, MAN2B1), and sulfatases (GNS, IDS).

(B) Network graph of module 159 (M159). All proteins in M159 exhibit altered abundance in MUT brain, including lysosomal proteins, CTSA, PLA2G15, and GM2A.

(C) Module 2 (M2) contains multiple lysosomal proteins with increased abundance in MUT brain compared to WT, including CTSS, CTSL, GRN, IDS, MAN2B1.

(D) Module 213 (M213) contains multiple proteins with increased abundance in MUT brain, including lysosomal proteins, GLB1, GNS, CTSB, MAN2B2, and PLBD2. Network attributes (B-D): Node size denotes its weighted degree centrality (~importance in module), node color indicates proteins with altered abundance in MUT brain relative to WT, purple outline highlight proteins identified as lysosomal in (Geladaki et al., 2019), black edges indicate known protein-protein interactions, and grey-red edges denote the relative strength of protein covariation within a module (gray = weak, red = strong). P-Adjust values represent enrichment of proteins identified as lysosomal in Geladaki et al., 2019.

(E) The overall effect of genotype on M159 module abundance (WT 10.83 \pm 0.002, MUT 10.94 \pm 0.002, p=0.031).

(F) The overall effect of genotype on M2 module abundance (WT 13.74 \pm 0.001, MUT 13.85 \pm 0.0009, p=0.0006).

(G) The overall effect of genotype on M213 abundance (WT 12.17 \pm 0.002, MUT 12.33 \pm 0.002,

p=0.0037). Data reported as mean ± SEM, error bars are SEM. *p<0.05, ** p<0.01, ***p<0.001, empirical Bayes quasi-likelihood F-test with Bonferroni correction (E-G).

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Figure 4. SWIP^{P1019R} mutant neurons display structural abnormalities in endo-lysosomal compartments *in vitro*

(A) Experimental design. Cortices were dissected from P0 pups, and neurons were dissociated and cultured on glass coverslips for 15 days. Cultures were fixed, stained, and imaged using confocal microscopy. 3D puncta volumes were reconstructed from z-stack images using Imaris software.
(B-C) Representative 3D reconstructions of WT and MUT DIV15 neurons (respectively) stained for EEA1 (yellow) and MAP2 (magenta).

(D-E) Representative 3D reconstructions of WT and MUT DIV15 neurons (respectively) stained for Cathepsin D (cyan) and MAP2 (magenta).

(F) Graph of the average number of EEA1+ volumes per soma in each image (WT 95.0 \pm 5.5, n=24 neurons; MUT 103.7 \pm 3.7, n=24 neurons; t_{40.2}=1.314, p=0.1961).

(G) Graph of the average EEA1+ volume size per soma shows larger EEA1+ volumes in MUT neurons (WT 0.15 \pm 0.01 µm³, n=24 neurons; MUT 0.30 \pm 0.02 µm³, n=24 neurons; U=50, p<0.0001). (H) Graph of the average number of Cathepsin D+ volumes per soma illustrates less Cathepsin D+ volumes in MUT neurons (WT 30.4 \pm 1.4, n=42; MUT 17.2 \pm 0.9, n=42; t₇₁=7.943, p<0.0001). (I) Graph of the average Cathepsin D+ volume size per soma demonstrates larger Cathepsin D+ volumes in MUT neurons (WT 0.54 \pm 0.02 µm³, n=42; MUT 0.69 \pm 0.04 µm³, n=42; t₆₃=3.701, p=0.0005). (J) Histogram of EEA1+ volumes illustrate differences in size distributions between MUT and WT neurons. (K) Histogram of CathD+ volumes show differences in size distributions between MUT and WT neurons. Analyses included at least three separate culture preparations. Scale bars, 5 µm (B-E). Data reported as mean \pm SEM, error bars are SEM. ***p<0.001, ****p<0.0001, two-tailed t-tests or Mann-Whitney U test (G).



Figure 5. SWIP^{P1019R} mutant brains exhibit markers of abnormal endo-lysosomal structures and cell death *in vivo*

(A-B) Representative images of adolescent (P42) WT and MUT motor cortex stained with cleaved caspase-3 (CC3, green).

(C) Anatomical representation of mouse brain with motor cortex highlighted in red, adapted from the Allen Brain Atlas (Oh et al., 2014).

(D-E) Representative image of adult (10 mo) WT and MUT motor cortex stained with CC3 (green).

(F, G, I, and J) DAPI co-stained images for (A, B, D, and E, respectively). Scale bar for (A-J), 15 µm.

(H) Graph depicting the normalized percentage of DAPI+ nuclei that are positive for CC3 per image. No difference is seen at P42, but the amount of CC3+ nuclei is significantly higher in aged MUT mice (P42 WT 6.97 \pm 0.80%, P42 MUT 5.26 \pm 0.90%, 10mo WT 25.38 \pm 2.05%, 10mo MUT 44.01 \pm 1.90%,

n=24 images per genotype taken from 4 different mice, H=74.12, p<0.0001). We observed no difference in number of nuclei per image between genotypes.

(K) Representative transmission electron microscopy (TEM) image taken of soma from adult (7mo) WT motor cortex. Arrowheads delineate electron-dense lipofuscin material, Nuc = nucleus.

(L) Representative transmission electron microscopy (TEM) image taken of soma from adult (7mo) MUT motor cortex.

(M) Inset from (K) highlights lysosomal structure in WT soma. Pseudo-colored region depicts lipofuscin area, demarcated as L.

(N) Inset from (L) highlights large lipofuscin deposit in MUT soma (L, pseudo-colored region) with electrondense and electron-lucent lipid-like (asterisk) components.

(O) Graph of areas of electron-dense regions of interest (ROI) shows increased ROI size in MUT neurons (WT $2.4 \times 10^5 \pm 2.8 \times 10^4 \text{ nm}^2$, n=50 ROIs; MUT $8.2 \times 10^5 \pm 9.7 \times 10^4 \text{ nm}^2$, n=75 ROIs; U=636, p<0.0001).

(P) Graph of the average number of presumptive lysosomes with associated electron-dense material reveals increased number in MUT samples (WT 3.14 ± 0.72 ROIs, n=14 images; MUT 10.86 ± 1.42 ROIs, n=14 images; U=17, p<0.0001). For (N) and (O), images were taken from multiple TEM grids, prepared from n=3 animals per genotype. Scale bar for all TEM images, 1 µm. Data reported as mean ± SEM, error bars are SEM. ***p<0.001, ****p<0.0001, Kruskal-Wallis test (F), Mann-Whitney U test (O-P).



Figure 6. SWIPP1019R mutant mice display persistent deficits in cued fear memory recall

(A) Experimental fear conditioning paradigm. After acclimation to a conditioning chamber, mice received a mild aversive 0.4mA footshock paired with a 2900Hz tone. 48 hours later, the mice were placed in a chamber with different tactile and visual cues. The mice acclimated for two minutes and then the 2900Hz tone was played (no footshock) and freezing behavior was assessed.

(B) Line graphs of WT and MUT freezing response during cued tone memory recall. Data represented as average freezing per genotype in 30 s time bins. The tone is presented after t = 120 s, and remains on for 120 seconds (Tone ON). Two different cohorts of mice were used for age groups P42 (top) and 6.5mo (bottom). Two-way ANOVA analysis of average freezing during Pre-Tone and Tone periods reveal a Genotype x Time effect at P42 (WT n=10, MUT n=10, $F_{1,18}$ =4.944, p=0.0392) and 6.5mo (WT n=13, MUT n=11, $F_{1,22}$ = 13.61, p=0.0013).

(C) Graphs showing the average %time freezing per animal before and during tone presentation. Top: freezing is reduced by 20% in MUT adolescent mice compared to WT littermates (Pre-tone WT 16.5 \pm 2.2%, n=10; Pre-tone MUT 13.0 \pm 1.8%, n=10; t₃₆=0.8569, p=0.6366; Tone WT 52.8 \pm 3.8%, n=10; Tone MUT 38.0 \pm 3.6%, n=10; t₃₆=3.539, p=0.0023), Bottom: freezing is reduced by over 30% in MUT adult mice compared to WT littermates (Pre-tone WT 21.1 \pm 2.7%, n=13; Pre-tone MUT 23.7 \pm 3.8%, n=11; t₄₄=0.4675, p=0.8721; Tone WT 69.7 \pm 4.3%, n=13; Tone MUT 53.1 \pm 5.2%, n=11; t₄₄=2.921, p=0.0109). Data reported as mean \pm SEM, error bars are SEM. *p<0.05, **p<0.01, two-way ANOVAs (B) and Sidak's post-hoc analyses (C).



Child (mean = 10.4yo)

7

4

F

Figure 7. SWIP^{P1019R} **mutant mice exhibit surprising motor deficits that are confirmed in human patients** (A) Rotarod experimental setup. Mice walked atop a rod rotating at 32rpm for 5 minutes, and the duration of time they remained on the rod before falling was recorded.

+

+

(B) Line graph of average duration animals remained on the rod per genotype across four trials, with an intertrial interval of 40 minutes. The same cohort of animals was tested at two different ages, P45 (top) and 5.5

+

+
months (bottom). Genotype had a significant effect on task performance at both ages (top, P45: genotype effect, $F_{1,25}$ =7.821, p=0.0098. bottom, 5.5mo: genotype effect, $F_{1,23}$ = 7.573, p=0.0114).

(C) Graphs showing the average duration each animal remained on the rod across trials. At both ages, the MUT mice exhibited an almost 50% reduction in their ability to remain on the rod (Top, P45: WT 169.9 \pm 25.7 s, MUT 83.8 \pm 15.9 s, U=35, p=0.0054. Bottom, 5.5mo: WT 135.9 \pm 20.9 s, MUT 66.7 \pm 9.5 s, t₁₈=3.011, p=0.0075).

(D) TreadScan task. Mice walked on a treadmill for 20 s while their gate was captured with a high-speed camera. Diagrams of gait parameters measured in (E-G) are shown below the TreadScan apparatus.

(E) Average swing time per stride for hindlimbs. At P45 (top), there is no significant difference in rear swing time (WT 156.2 \pm 22.4 ms, MUT 132.3 \pm 19.6 ms, U=83, p=0.7203). At 5.5mo (bottom), MUT mice display

significantly longer rear swing time (WT 140 \pm 6.2 ms, MUT 252.0 \pm 21.6 ms, t₁₂=4.988, p=0.0003).

(F) Average stride length for hindlimbs. At P45 (top), there is no significant difference in stride length (WT 62.3 \pm 2.0 mm, MUT 60.5 \pm 2.1 mm, U=75, p=0.4583). At 5.5mo (bottom), MUT mice take significantly longer strides with their hindlimbs (WT 60.8 \pm 0.8 mm, MUT 73.6 \pm 2.7 mm, t_{11.7}=4.547, p=0.0007).

(G) Average homologous coupling for front and rear limbs. Homologous coupling is 0.5 when the left and right feet are completely out of phase. At P45 (top), WT and MUT mice exhibit normal homologous coupling (WT 0.48 \pm 0.005, MUT 0.48 \pm 0.004, U=76.5, p=0.4920). At 5.5 mo (bottom), MUT mice display decreased homologous coupling, suggestive of abnormal gait symmetry (WT 0.48 \pm 0.003, MUT 0.46 \pm 0.004, t_{18.8}=3.715, p=0.0015). At P45: n=14 WT, n=13 MUT; At 5.5mo: n=14 WT, n=11 MUT.

(H) Table of motor findings in clinical exam of human patients with the homozygous SWIP^{P1019R} mutation (n=7). All patients exhibit motor dysfunction (+ = symptom present). Data reported as mean \pm SEM, error bars are SEM.*p<0.05, **p<0.01, ***p<0.001, ****p<0.0001, two-way repeated measure ANOVAs (B), Mann-Whitney U tests and two-tailed t-tests (C-G).



Figure 8. Model of neuronal endo-lysosomal pathology in SWIP^{P1019R} mutant mice

(A) Wild-type WASH function in mouse brain. Under normal conditions, the WASH complex interacts with many endosomal proteins and cytoskeletal regulators, such as the Arp2/3 complex. These interactions enable restructuring of the endosome surface (actin in gray) and allow for cargo segregation and scission of vesicles. Substrates are transported to the late endosome for lysosomal degradation, to the Golgi network for modification, or to the cell surface for recycling.

(B) Loss of WASH function leads to increased lysosomal degradation in mouse brain. Destabilization of the WASH complex leads to enlarged endosomes and lysosomes, with increased substrate accumulation at the lysosome. This suggests an increase in flux through the endo-lysosomal pathway, possibly as a result of mis-localized endosomal substrates.

(C) Wild-type mice exhibit normal motor function.

(D) SWIP^{P1019R} mutant mice display progressive motor dysfunction in association with these subcellular alterations.



Figure 2- figure supplement 1. Overexpression of SWIP^{P1019R} decreases WASH complex binding in cultured cells; related to Figure 2

(A) Schematic showing overexpression of WT or MUT SWIP^{P1019R} in HEK293T cells followed by immunoprecipitation.

(B) Western blots of input (5%, left) and immunoprecipitated (IP, right) protein. Two samples per condition were run on two separate gels, n=4 biological replicates from separate experiments.

(C) Quantification of B normalized to WT. Strumpellin (WT 100.0 \pm 6.8%, MUT 54.8 \pm 8.0%, t_{5.9}=4.290, p=0.0054), WASH1 (WT 100.0 \pm 7.3%, MUT 41.4 \pm 4.4%, t_{4.9}=6.902, p=0.0011), HA (WT 100.0 \pm 4.1%, MUT 107.8 \pm 4.1%, t_{6.0}=1.344, p=0.2275). Data reported as mean \pm SEM, error bars are SEM. **p<0.01, two-tailed t-tests.



Figure 2- figure supplement 2. SWIP^{P1019R} MUT brain displays significant alterations in protein abundance compared to WT; related to Figures 2 and 3

(A) Interactome of altered proteins. Nodes reflect protein name, light gray lines delineate proteins identified as different in MUT compared to WT (ΔWASHC4 in center), dark dashed lines indicate known protein-protein interactions from HitPredict database (López et al., 2015). Color reflects cellular function seen in B. Nodes with red borders delineate WASH complex proteins.

(B) Cellular function of proteins in A, as reflected in published literature. % reflects the percentage of proteins in a category out of the total 85 altered proteins.

- (C-G) Clustergrams of:
- (C) Proteins with increased (red) or decreased (blue) abundance in MUT brains compared to WT.
- (D) Known protein components of the WASH complex and their previously reported interactors.
- (E) Proteins with lysosomal function.
- (F) Proteins identified in the WASH1-BioID2 proteome (Figure 1).
- (G) Proteins with links to intellectual disability (I.D.) or neurodegeneration (Degen.).





С













Figure 2- figure supplement 3. Multiple protein networks display significant alterations in MUT brain compared to WT; related to Figures 2 and 3

(A) Module 14 (M14) containing endosomal proteins. Two of the three retromer sorting complex subunits are highlighted with red borders, with enrichment calculated relative to the CORUM database (Giurgiu et al., 2019).

(B) Module 83 (M83) containing endoplasmic reticulum (ER) stress response proteins. Seven of the 185 proteins identified to have ER function by LOPIT-DC spatial proteomics are highlighted with orange borders (Geladaki et al., 2019).

(C-D) Modules 248 and 35 (M248, M35, respectively) containing synaptic proteins. Network attributes for graphs in B-E: Node size denotes its weighted degree centrality (~importance in module), colored node indicates altered abundance in MUT brain relative to WT, black node border denotes proteins identified in the WASH1-BioID proteome (Figure 1), colored node border denotes proteins identified in other datasets, black edges indicate known protein-protein interactions, and grey-red edges denote the relative strength of protein covariation within a module (gray = weak, red = strong). P-adjust enrichment values are calculated relative to the CORUM database (B), Lopit-DC dataset (C), or excitatory postsynapse proteome (DLG4-BioID, D-E) (Geladaki et al., 2019; Giurgiu et al., 2019; Uezu et al., 2016). (E) Summary box plots of module abundance for all seven brain fractions analyzed presented as log2(adjusted module intensities). All modules display significant differences between WT and MUT groups (M14: WT 13.94 ± 0.002 MUT 13.82 ± 0.002, p=0.0207; M83: WT 14.46 ± 0.004, MUT 14.70 ± 0.003, p=0.0088; M248: WT 13.83 ± 0.001, MUT 13.77 ± 0.001, p=0.00134; M35: WT 14.19 ± 0.001, MUT 14.10 ± 0.001, p=0.0030).

(F) Normalized protein abundance for all proteins in each module across the seven subcellular fractions analyzed. Plots correspond to modules seen in F, and data colors reflect genotypes in F. Thick lines represent the mean protein intensity per genotype. Number of proteins in each module are indicated at the top of each graph. Data reported as mean \pm SEM, error bars are SEM. *p<0.05, **p<0.01, ***p<0.001, GLM (model: 0 ~ Fraction + Module) and empirical Bayes quasi-likelihood F-test with Bonferroni correction (F).



Figure 5- figure supplement 1. There is no significant difference in striatal, cerebellar, or hippocampal cell death between WT and MUT mice; related to Figure 5

(A) Representative image of adolescent (P42) WT striatum stained with cleaved caspase-3 (CC3, green).

(B) Representative image of adolescent (P42) MUT striatum stained with cleaved caspase-3 (CC3, green). (C and D) DAPI co-stained images of A and B, respectively.

(E) Anatomical representation of mouse brain with striatum highlighted in red, adapted from the Allen Brain Atlas (Oh et al., 2014).

(F) Representative image of adult (10 mo) WT striatum stained with CC3 (green).

(G) Representative image of adult (10 mo) MUT striatum stained with CC3 (green).

(H and I) DAPI co-stained images of F and G, respectively. Scale bars for A-I are 15 $\mu m.$

(J) Graph depicting the normalized % of DAPI+ nuclei that are positive for CC3 per image. No difference is seen between genotypes at either age (P42 WT $3.70 \pm 0.99\%$, P42 MUT $1.95 \pm 0.49\%$, 10mo WT $16.77 \pm 2.09\%$, 10mo MUT 24.86 $\pm 2.17\%$, H=61.87, p<0.0001).

(K) Representative image of adolescent (P42) WT cerebellum stained with cleaved caspase-3 (CC3, green).

(L) Representative image of adolescent (P42) MUT cerebellum stained with cleaved caspase-3 (CC3, green).

(M and N) DAPI and Calbindin co-stained images of K and L, respectively.

(O) Anatomical representation of mouse cerebellum, adapted from the Allen Brain Atlas (Oh et al., 2014). Red region highlights area used for imaging.

(P) Representative image of adult (10 mo) WT cerebellum stained with CC3 (green).

(Q) Representative image of adult (10 mo) MUT cerebellum stained with CC3 (green). No significant CC3 staining is observed at either age.

(R and S) DAPI and Calbindin co-stained images of P and Q, respectively. Scale bars for K-S are 50 μ m. (T) Graph depicting the number of Calbindin+ somas per image, a marker for Purkinje cells. No difference is seen between genotypes at either age (P42 WT 20.50 ± 0.53, P42 MUT 20.67 ± 0.59, 10mo WT 21.42 ± 0.85, 10mo MUT 22.63 ± 0.74, H=4.891, p=0.1799).

(U) Representative image of adolescent (P42) WT cerebellum stained with cleaved caspase-3 (CC3, green).

(V) Representative image of adolescent (P42) MUT cerebellum stained with cleaved caspase-3 (CC3, green).

(W and X) DAPI and Calbindin co-stained images of U and V, respectively.

(Y) Anatomical representation of mouse hippocampus CA1, adapted from the Allen Brain Atlas (Oh et al., 2014). Red region highlights area used for imaging.

(Z) Representative image of adolescent (P42) WT cerebellum stained with cleaved caspase-3 (CC3, green).

(AA) Representative image of adolescent (P42) MUT cerebellum stained with cleaved caspase-3 (CC3, green).

(BB and CC) DAPI and Calbindin co-stained images of U and V, respectively.

(DD) Graph of the normalized % of DAPI+ nuclei that are positive for CC3 per image. Very little CC3 staining is seen at either age, regardless of genotype (P42 WT $0.21 \pm 0.11\%$, P42 MUT $0.15 \pm 0.11\%$, 10mo WT $0.81 \pm 0.28\%$, 10mo MUT $4.13 \pm 0.96\%$, H=20.27, p=0.0001). Data obtained from four animals per condition, and reported as mean ± standard error of the mean (SEM), with error bars as SEM. Kruskal-Wallis test (J,T, and DD).



Figure 6- figure supplement 1. SWIP^{P1019R} mutant mice do not display deficits in spatial working memory or novel object recognition; related to Figure 6

(A) Y-maze paradigm. Mice were placed in the center of the maze and allowed to explore all three arms freely for five minutes. Each arm had distinct visual cues.

(B) Graph depicting the percent alternations achieved for each mouse at adolescence (WT 51.48 \pm 2.47%, MUT 50.81 \pm 2.19%, t₂₄=0.2036, p=0.8404).

(C) Graph depicting the percent alternations achieved for each mouse at adulthood (WT 56.12 \pm 1.53%, MUT 57.93 \pm 2.56%, t₁₈=0.6074, p=0.5511).

(D) Graphs of the number of direct revisits mice made to the arm they just explored reveal no difference between genotypes at adolescence (WT 2.64 \pm 0.50, MUT 2.42 \pm 0.42, t₂₄=0.3482, p=0.7308).

(E) Graphs of the number of direct revisits mice made at adulthood (WT 1.36 \pm 0.33, MUT 1.42 \pm 0.36, t₂₄=0.1231, p=0.9031).

(F) Similar to D-E, there were no differences in the number of indirect revisits (ex: arm A \rightarrow arm B \rightarrow arm

A) between genotypes at adolescence (WT 10.21 ± 1.03, MUT 12.00 ± 1.48, U=69, p=0.4515)

(G) Indirect revisits at adulthood (WT 12.86 \pm 1.26, MUT 15.17 \pm 1.43, t₂₃=1.211, p=0.2380).

(H) There were no significant differences in total distance travelled at adolescence, suggesting that motor function did not affect Y-maze performance (WT 2401 \pm 98.9 cm, MUT 2406 \pm 121.0 cm, t₂₂=0.03281, p=0.9741).

(I) No difference in distanced travelled at adulthood (WT 2761 \pm 111.6 cm, MUT 3124 \pm 191.1 cm, t₁₈=1.638, p=0.1189).

(J) Novel object task. Mice first performed a 5-minute trial in which they were placed in an arena with two identical objects and allowed to explore freely, while their behavior was tracked with video software. Mice were returned to their home cage, and then re-introduced to the arena a half an hour later, where one of the objects had been replaced with a novel object, and their behavior was again tracked. Twenty-four hours later the same test was performed, but the novel object was replaced with another new object.

(K) Graph depicting adolescent animals' preference for the novel object during the three phases of the task, training (Train), short-term memory (STM), and long-term memory (LTM). No significant difference in object preference is seen between genotypes for any phase (Train WT -0.092 \pm 0.065, Train MUT -0.123 \pm 0.072, STM WT 0.488 \pm 0.076, STM MUT 0.315 \pm 0.094, LTM WT 0.479 \pm 0.046, LTM MUT 0.373 \pm 0.076, F_{1,22}=1.840, p=0.1887).

(L) Graph depicting adult animals' preference for the novel object. (Train WT -0.066 \pm 0.053, Train MUT -0.130 \pm 0.089, STM WT 0.416 \pm 0.060, STM MUT 0.274 \pm 0.096, LTM WT 0.316 \pm 0.059, LTM MUT 0.306 \pm 0.050, F_{1.22}=0.9735, p=0.3345).

(M) Graph depicting the total amount of time (in seconds) adolescent animals spent exploring both objects in each phase of the task. No significant difference in exploration time was observed across genotypes, suggesting that genotype does not hinder object exploration (Train P44 WT 38.10 ± 2.45 s, Train P44 MUT 50.04 ± 5.44 s, STM P44 WT 55.02 ± 4.31 s, STM P44 MUT 63.60 ± 4.50 s, LTM P45 WT 46.75 ± 3.34 s, LTM P45 MUT 68.08 ± 7.54 s, $F_{1,22}$ =7.373, p=0.0126).

(N) Graph depicting the total time adult animals spent exploring both objects in each the task (Train WT $39.10 \pm 3.62 \text{ s}$, Train MUT $51.15 \pm 4.91 \text{ s}$, STM WT $44.75 \pm 4.87 \text{ s}$, STM MUT $49.56 \pm 6.16 \text{ s}$, LTM WT $39.02 \pm 5.29 \text{ s}$, LTM MUT $55.15 \pm 7.34 \text{ s}$, F_{1,22}=2.936, p=0.1007). For all Y-maze measures, WT n=14, MUT n=12. For all novel object measures, WT n=13, MUT n=11. Data reported as mean \pm SEM, error bars are SEM. Two-tailed t-tests or Mann-Whitney U tests (B-I), two-way ANOVAs (K-N).



Figure 6- figure supplement 2. SWIP^{P1019R} mutant mice do not have significant deficits in **contextual fear memory recall, auditory perception, or tactile sensation; related to Figure 6** (A) Experimental fear conditioning scheme. After acclimation, mice received a mild aversive 0.4mA footshock paired with a 2900Hz tone in a conditioning chamber. 24 hours later, the mice were placed back in the same chamber to assess freezing behavior (without footshock or tone).

(B) Experimental startle response setup used to assess hearing and somatosensation. Mice were placed in a plexiglass tube atop a load cell that measured startle movements in response to stimuli. (C) Line graphs of WT and MUT freezing response during the contextual memory recall task. Data represented as average freezing per genotype in 30 second time bins. The task was performed with two different cohorts for the different ages, P42 (top) and 6.5mo (bottom). Top: no significant difference in freezing at P42 (Two-way repeated measure ANOVA, Genotype effect, $F_{1,18}$ =0.088, p=0.7698. Sidak's post-hoc analysis, 30 s p=0.8388, 60 s p=0.9990, 90 s p=0.9964, 120 s p=0.3281), Bottom: no significant difference in freezing at 6.5mo (Two-way repeated measure ANOVA, Genotype effect, $F_{1,22}$ = 3.723, p=0.0667. Sidak's post-hoc analysis, 30 s p=0.8977, 60 s p=0.1636, 90 s p=0.9979, 120 s p=0.0037).

(D) Graphs showing the average total freezing time per animal during context exposure. Top: no significant difference is seen between WT and MUT mice at P42 (WT 34.01 \pm 6.32%, MUT 36.99 \pm 7.81%, t₁₇=2.985, p=0.7699). Bottom: no significant different is seen between genotypes at 6.5mo (WT 43.94 \pm 6.00%, MUT 28.73 \pm 4.80%, t_{21.6}=1.980, p=0.0606).

(E) Graphs of individual animals' startle response to a 2900Hz tone played at 80dB. MUT mice were not significantly more reactive to the tone than WT at P50 (WT 25.96 \pm 4.95, MUT 40.68 \pm 5.05, U=35, p=0.2799), or at 6.5mo (WT 14.07 \pm 3.27, MUT 14.85 \pm 1.49, U=47, p=0.2768).

(F) Graphs of individual animals' startle response to a 0.4mA footshock. No significant difference observed between genotypes at either age (P55 WT 1527 \pm 215.7, P55 MUT 1996 \pm 51.0, U=28.50, p=0.0542; 6.5mo WT 1545 \pm 179.5, 6.5mo MUT 1817 \pm 119.1, U=47, p=0.2360). Startle response reported in arbitrary units (A.U.). For all adolescent measures: WT n=10, MUT n=10. For adult freezing measures: WT n=13, MUT n=11. For adult startle responses: WT n=13, MUT n=10. Data reported as mean \pm SEM, error bars are SEM.*p<0.05, **p<0.01, two-way repeated measure ANOVAs (C), two-tailed t-tests (D), and Mann-Whitney U tests (E-F).



Figure 7- figure supplement 1. Progressive gait changes in SWIP^{P1019R} mutant mice are not restricted to rear limbs; related to Figure 7

(A) Graph of average swing time per stride for front limbs. At P45 (top), there is no significant difference in front swing time (WT 177.5 \pm 10.9 ms, MUT 175.3 \pm 8.7 ms, t₂₄=0.1569, p=0.8766). At 5.5 mo (bottom), MUT mice take significantly longer to swing their forelimbs (WT 178.6 \pm 6.2 ms, MUT 206.3 \pm 7.2 ms, t_{21.4}=2.927, p=0.0079).

(B) Graph of average stride length for front limbs. At P45 (top), there is no difference in WT and MUT stride length (WT 57.0 \pm 0.9 mm, MUT 59.2 \pm 1.4 mm, U=68, p=0.2800). At 5.5mo (bottom), MUT mice take significantly longer strides with their forelimbs (WT 60.0 \pm 0.6 mm, MUT 63.7 \pm 0.9 mm, t₁₇=3.545, p=0.0024).

(C) Graph of average homolateral coupling, the fraction of a reference foot's stride when its ipsilateral foot starts its stride. At P45, there is no significant difference in homolateral coupling (WT 0.48 \pm 0.005, MUT 0.48 \pm 0.004, U=90, p=0.9713), but at 5.5mo, MUT mice display decreased homolateral coupling (WT 0.48 \pm 0.002, MUT 0.45 \pm 0.005, t_{13.5}=3.469, p=0.0039).

(D) Graph of average track width between front limbs. At P45 (top), there is a significantly narrower front track width in MUT compared to WT (WT 16.99 ± 0.15 mm, MUT 15.12 ± 0.33 mm, t₁₇=5.192, p<0.0001). This difference persists into adulthood at 5.5mo (WT 19.36 ± 0.23 mm, MUT 16.74 ± 0.46 mm, t₁₅=5.055, p=0.0001).

(E) Graph of average track width between rear limbs. At P45 (top), there is no difference in WT and MUT rear track widths (WT 29.58 \pm 0.51 mm, MUT 28.77 \pm 0.36 mm, t₂₃=1.292, p=0.2091). At 5.5mo (bottom), mutants display significantly narrower rear track widths (WT 32.59 \pm 0.34 mm, MUT 30.01 \pm 0.46 mm, t_{19.4}=4.502, p=0.0002). For P45 measures: WT n=14, MUT n=13; for 5.5mo measures: WT n=14, MUT n=11. Data reported as mean \pm SEM, error bars are SEM. **p<0.01, ***p<0.001, ***p<0.001, two-tailed t-tests or Mann-Whitney U tests.

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558

559 AUTHOR CONTRIBUTIONS

Conceptualization: JLC, TWAB, IHK, and SHS; Methodology: JLC, TWAB, IHK, and SHS;
Investigation: JLC, TWAB, IHK, GW, TH, RV, ES, and AR; Resources: ES, RV, and SHS;
Writing—Original Draft: JLC and SHS; Writing—Editing: JLC, TWAB, IHK, and SHS;
Visualization: JLC; Funding Acquisition: JLC, TWAB, IHK, and SHS. All authors
discussed the results and commented on the manuscript.

565

566 **DECLARATION OF INTERESTS**

567 The authors declare no competing interests.

568 **RESOURCE AVAILABILITY**

- 569 Lead Contact
- 570 Further information and requests for resources and reagents should be directed to and
- 571 will be fulfilled by the Lead Contact, Scott Soderling (scott.soderling@duke.edu).
- 572 Materials Availability
- 573 Plasmids generated by this study are available upon request from corresponding author
- 574 Scott H. Soderling (scott.soderling@duke.edu).

575 Data and Code Availability

- 576 The data and source code used in this study are available online at 577 https://github.com/twesleyb/SwipProteomics.
- 578

579 MATERIALS AND METHODS

580 Animals

581 We generated *Washc4* mutant (SWIP^{P1019R}) mice in collaboration with the Duke 582 Transgenic Core Facility to mimic the de novo human variant at amino acid 1019 of 583 human WASHc4. A CRISPR-induced CCT>CGT point mutation was introduced into exon 584 29 of Washc4. 50ng/µl of the sgRNA (5'-ttgagaatactcacaagaggagg-3'), 100ng/µl Cas9 585 mRNA, and 100ng/µl of a repair oligonucleotide containing the C>G mutation were 586 injected into the cytoplasm of B6SJLF1/J mouse embryos (Jax #100012) (See Table S4 587 for the sequence of the repair oligonucleotide). Mice with germline transmission were then 588 backcrossed into a C57BL/ 6J background (Jax #000664). At least 5 backcrosses were obtained before animals were used for behavior. We bred heterozygous SWIP^{P1019R} mice 589

590 together to obtain age-matched mutant and wild-type genotypes for cell culture and 591 behavioral experiments. Genetic sequencing was used to screen for germline 592 transmission of the C>G point mutation (FOR: 5'-tgcttgtagatgtttttcct-3', REV: 5'-593 gttaacatgatcctatggcg-3'). All mice were housed in the Duke University's Division of 594 Laboratory Animal Resources or Behavioral Core facilities at 2-5 animals/cage on a 595 14:10h light:dark cycle. All experiments were conducted with a protocol approved by the 596 Duke University Institutional Animal Care and Use Committee in accordance with NIH 597 quidelines.

598 Human Subjects

599 We retrospectively analyzed clinical findings from seven children with homozygous 600 *WASHC4*^{c.3056C>G} mutations (obtained by Dr. Rajab in 2010 at the Royal Hospital, Muscat, 601 Oman). The original report of these human subjects and parental consent for data use 602 can be found in (Ropers et al., 2011).

603 Cell Lines

604 HEK293T cells (ATCC #CRL-11268) were purchased from the Duke Cell Culture 605 facility. and were tested for mycoplasma contamination. HEK239T cells were used for co-606 immunoprecipitation experiments and preparation of AAV viruses.

607 **Primary Neuronal Culture**

608 Primary neuronal cultures were prepared from P0 mouse cortex. P0 mouse pups 609 were rapidly decapitated and cortices were dissected and kept individually in 5ml 610 Hibernate A (Thermo #A1247501) supplemented with 2% B27(Thermo #17504044) at 611 4°C overnight to allow for individual animal genotyping before plating. Neurons were then 612 treated with Papain (Worthington #LS003120) and DNAse (VWR #V0335)-supplemented Hibernate A for 18min at 37°C and washed twice with plating medium (plating medium:
Neurobasal A (Thermo #10888022) supplemented with 10% horse serum, 2% B-27, and
1% GlutaMAX (Thermo #35050061)), and triturated before plating at 250,000 cells/well
on poly-L-lysine-treated coverslips (Sigma #P2636) in 24-well plates. Plating medium was
replaced with growth medium (Neurobasal A, 2% B-27, 1% GlutaMAX) 2 hours later. Cell
media was supplemented and treated with AraC at DIV5 (5uM final concentration/well).
Half-media changes were then performed every 4 days.

620 Plasmid DNA Constructs

621 For immunoprecipitation experiments, a pmCAG-SWIP-WT-HA construct was 622 generated by PCR amplification of the human WASHC4 sequence, which was then 623 inserted between Nhel and Sall restriction sites of a pmCAG-HA backbone generated in 624 our lab. Site-directed mutagenesis (Agilent #200517) was used to introduce a C>G point 625 mutation into this pmCAG-SWIP-WT-HA construct for generation of a pmCAG-SWIP-626 MUT-HA construct (FOR: 5'-ctacaaagttgagggtcagacggggaacaattatatagaaa-3', REV: 5'-627 tttctatataattgttccccgtctgaccctcaactttgtag-3'). For iBioID experiments, an AAV construct 628 expressing hSyn1-WASH1-BioID2-HA was generated by cloning a Washc1 insert 629 between Sall and HindIII sites of a pAAV-hSyn1-Actin Chromobody-Linker-BioID2-pA 630 construct (replacing Actin Chromobody) generated in our lab. This backbone included a 631 25nm GS linker-BioID2-HA fragment from Addgene #80899, generated by Kim et al. (Kim 632 et al., 2016). An hSyn1-solubleBioID2-HA construct was created similarly, by removing 633 Actin Chromobody from the above construct. Oligonucleotide sequences are reported in 634 Table S4. Sequences of the plasmid DNA constructs are available online (see Key 635 Resources Table).

636 **AAV Viral Preparation**

637 AAV preparations were performed as described previously(Uezu et al., 2016). The 638 day before transfection, HEK293T cells were plated at a density of 1.5x10⁷ cells per 639 15cm² plate in DMEM media with 10% fetal bovine serum and 1% Pen/Strep (Thermo 640 #11965-092, Sigma #F4135, Thermo #15140-122). Six HEK293T 15cm² plates were 641 used per viral preparation. The next day, 30µg of pAd-DeltaF6 helper plasmid, 15µg of 642 AAV2/9 plasmid, and 15µg of an AAV plasmid carrying the transgene of interest were mixed in OptiMEM with PEI-MAX (final concentration 80µg/ml, Polysciences #24765). 643 644 2ml of this solution were then added dropwise to each of the 6 HEK293T 15cm² plates. 645 Eight hours later, the media was replaced with 20ml DMEM+10%FBS. 72 hours post-646 transfection, cells were scraped and collected in the media, pooled, and centrifuged at 647 1,500 rpm for 5 min at RT. The final pellet from the 6 cell plates was resuspended in 5 ml 648 of cell lysis buffer (15 mM NaCl, 5 mM Tris-HCl, pH 8.5), and freeze-thawed three times 649 using an ethanol/dry ice bath. The lysate was then treated with 50U/ml of Benzonase 650 (Novagen #70664), for 30min in a 37°C water bath, vortexed, and then centrifuged at 651 4,500 rpm for 30 min at 4°C. The resulting supernatant containing AAV particles was added 652 to the top of an iodixanol gradient (15%, 25%, 40%, 60% top to bottom) in an Optiseal 653 tube (Beckman Coulter #361625). The gradient was then centrifuged using a Beckman 654 Ti-70 rotor in a Beckman XL-90 ultracentrifuge at 67,000rpm for 70min, 18°C. The purified 655 viral solution was extracted from the 40%/60% iodixanol interface using a syringe, and 656 placed into an Amicon 100kDa filter unit (#UFC910024). The viral solution was washed 657 in this filter 3 times with 1X ice-cold PBS by adding 5ml of PBS and centrifuging at 4,900rpm for 45min at 4°C to obtain a final volume of approximately 200µl of concentrated
virus that was aliquoted into 5-10µl aliquots and stored at -80°C until use.

660 Immunocytochemistry

Primary antibodies: Rabbit anti-EEA1 (Cell Signaling Technology #C45B10,
1:500), Rat anti-CathepsinD (Novus #204712, 1:250), Guinea Pig anti-MAP2 (Synaptic
Systems #188004, 1:500)

Secondary antibodies: Goat anti-Rabbit Alexa Fluor 568 (Invitrogen #A11036,
1:1000), Goat anti-Guinea Pig Alexa Fluor 488 (Invitrogen #A11073, 1:1000), Goat antiRat Alexa Fluor 488 (Invitrogen #A11006, 1:1000), Goat anti-Guinea Pig Alexa Fluor 555
(Invitrogen #A21435, 1:1000)

668 At DIV15, neurons were fixed for 15 minutes using ice-cold 4%PFA/4% sucrose in 669 1X PBS, pH 7.4 (for EEA1 staining), or 30 minutes with 50% Bouin's solution/4% sucrose 670 (for CathepsinD staining, Sigma #HT10132), pH 7.4(Cheng et al., 2018). Fixed neurons were washed with 1X PBS, then permeabilized with 0.25% TritonX-100 in PBS for 8 671 672 minutes at room temperature (RT), and blocked with 5% normal goat serum/0.2% Triton-673 X100 in PBS (blocking buffer) for 1 hour at RT with gentle rocking. For EEA1/MAP2 674 staining, samples were incubated with primary antibodies diluted in blocking buffer at RT 675 for 1 hour. For CathepsinD/MAP2 staining, samples were incubated with primary 676 antibodies diluted in blocking buffer overnight at 4°C. For both conditions, samples were 677 washed three times with 1X PBS, and incubated for 30min at RT with secondary 678 antibodies, protected from light. After secondary antibody staining, coverslips were 679 washed three times with 1X PBS, and mounted with FluoroSave mounting solution

680 (Sigma #345789). See antibody section for primary and secondary antibody681 concentrations.

682 Immunohistochemistry

Primary antibodies: Rabbit anti-Cleaved Caspase-3 (Cell Signaling Technology
#9661, 1:2000), Mouse anti-Calbindin (Sigma #C9848, 1:2000), Rat anti-HA 3F10 (Sigma
#12158167001, 1:500)

Secondary antibodies: Donkey anti-Rabbit Alexa Fluor 488 (Invitrogen #A21206,
1:2000), Goat anti-Mouse Alexa Fluor 594 (Invitrogen #A11032, 1:2000), Goat anti-Rat
Alexa Fluor 488 (Invitrogen #A11006, 1:5000), Streptavidin Alexa Fluor 594 conjugate
(Invitrogen #S32356, 1:5000), 4',6-diamidino-2-phenylindole (DAPI, Sigma #D9542,
1:1000 for 10min at RT)

691 Mice were deeply anesthetized with isoflurane and then transcardially perfused 692 with ice-cold heparinized PBS (25U/ml) by gravity flow. After clearing of liver and lungs 693 (~2min), perfusate was switched to ice-cold 4% PFA in 1X PBS (pH 7.4) for 15 minutes. 694 Brains were dissected, post-fixed in 4%PFA overnight at 4°C, and then cryoprotected in 695 30% sucrose/1X PBS for 48hr at 4°C. Fixed brains were then mounted in OTC (Sakura 696 TissueTek #4583) and stored at -20°C until cryosectioning. Every third sagittal section 697 (30 µm thickness) was collected from the motor cortex and striatal regions. Free-floating 698 sections were then permeabilized with 1%TritonX-100 in 1X PBS at RT for 2 hr, and 699 blocked in 1X blocking solution (Abcam #126587) diluted in 0.2%TritonX-100 in 1X PBS 700 for 1hr at RT. Sections were then incubated in primary antibodies diluted in the 1X 701 blocking solution for two overnights at 4°C. After three washes with 0.2%TritonX-100 in 702 1X PBS, the sections were then incubated in secondary antibodies diluted in 1X blocking buffer for one overnight at 4°C. Sections were then washed four times with 0.2%TritonX100 in 1X PBS at RT, and mounted onto coverslips with FluoroSave mounting solution
(Sigma #345789).

706 Western Blotting

Primary antibodies: Rabbit anti-Strumpellin (Santa Cruz #sc-87442, 1:500), Rabbit
anti-WASH1 c-terminal (Sigma #SAB4200373, 1:500), Mouse anti-Beta Tubulin III
(Sigma #T8660, 1:10,000), Mouse anti-HA (BioLegend #MMS-101P, 1:5000)

Secondary antibodies: Donkey anti-Rabbit-HRP (GE Life Sciences #NA934,
1:5,000), Goat anti-mouse-HRP (GE Life Sciences #NA931, 1:5000)

712 Ten micrograms of each sample were electrophoresed through a 12-well, 4-20% SDS-PAGE gel (Bio-Rad #4561096) at 100V for 1hr at RT, transferred onto a 713 714 nitrocellulose membrane (GE Life Sciences #GE10600002) at 100V for 70min at RT on 715 ice, and blocked with 5% nonfat dry milk in TRIS-buffered saline containing 0.05% Tween-716 20 (TBST, pH 7.4). Gels were saved for Coomassie staining at RT for 30 min. Membranes 717 were probed with one primary antibody at a time for 24hr at 4°C, then washed four times 718 with TBST at RT before incubating with the corresponding species-specific secondary 719 antibody at RT for 1hr. Membranes were washed with TBST, and then enhanced 720 chemiluminescence (ECL) substrate was added (Thermo Fischer #32109). Membranes 721 were exposed to autoradiography films and scanned with an Epson 1670 at 600dpi. We 722 probed each membrane with one antibody at a time, then stripped the membrane with 723 stripping buffer (Thermo Fischer #21059) for 10min at RT, and then blocked for 1hr at RT 724 before probing with the next antibody. Order of probes: Strumpellin, then β -Tubulin, then 725 WASH1. We determined the optical density of the bands using Image J software (NIH).

Data obtained from three independent experiments were plotted and statistically analyzed
 using GraphPad Prism (version 8) software.

728 Immunoprecipitation

729 HEK293T cells were transfected with pmCAG-SWIP-WT-HA or pmCAG-SWIP-730 MUT-HA constructs for three days, as previously described (Mason et al., 2011). Cells 731 were lysed with lysis buffer (25mM HEPES, 150mM NaCl, 1mM EDTA, 1% NonidetP-40, 732 pH 7.4) containing protease inhibitors (5mM NaF, 1mM orthovanadate, 1mM AEBSF, and 733 2 µg/mL leupeptin/pepstatin) and centrifuged at 1,700g for 5 min. Collected supernatant 734 was incubated with 30µl of pre-washed anti-HA agarose beads (Sigma #A2095) on a 735 sample rotator (15 rpm) for 2 hrs at 4°C. Beads were then washed 3 times with lysis 736 buffer, and sample buffer was added before subjecting to immunoblotting as described 737 above. The protein-transferred membrane was probed individually for WASH1, 738 Strumpellin, and HA. Data were collected from four separate preparations of WT and MUT conditions. 739

740 Electron Microscopy

741 Adult (7mo) WT and MUT SWIP^{P1019R} mice were deeply anesthetized with 742 isoflurane and then transcardially perfused with warmed heparinized saline (25U/ml 743 heparin) for 4 minutes, followed by ice-cold 0.15M cacodylate buffer pH 7.4 containing 744 2.5% glutaraldehyde (Electron Microscopy Sciences #16320), 3% paraformaldehyde, 745 and 2mM CaCl₂ for 15 minutes. Brain samples were dissected and stored on ice in the 746 same fixative for 2 hours before washing in 0.1M sodium cacodylate buffer (3 changes 747 for 15 minutes each). Samples were then post-fixed in 1.0% OsO₄ in 0.1 M Sodium 748 cacodylate buffer for 1 hour on a rotator. Samples were then washed in 3, 15-minute

749 changes of 0.1M sodium cacodylate. Samples were then placed into *en bloc* stain (1%) 750 uranyl acetate) overnight at 4°C. Subsequently, samples were dehydrated in a series of 751 ascending acetone concentrations including 50%, 70%, 95%, and 100% for three cycles 752 with 15 minutes incubation at each concentration change. Samples were then placed in 753 a 50:50 mixture of epoxy resin (Epon) and acetone overnight on a rotator. This solution 754 was then replaced twice with 100% fresh Epon for at least 2 hours at room temperature 755 on a rotator. Samples were embedded with 100% Epon resin in BEEM capsules (Ted 756 Pella) for 48 hours at 60°C. Samples were ultrathin sectioned to 60-70nm on a Reichert 757 Ultracut E ultramicrotome. Harvested grids were then stained with 2% uranyl acetate in 758 50% ethanol for 30 minutes and Sato's lead stain for 1 min. Micrographs were acquired 759 using a Phillips CM12 electron microscope operating at 80Kv, at 1700x magnification. 760 Micrographs were analyzed in Adobe Photoshop 2019, using the "magic wand" tool to 761 demarcate and measure the area of electron-dense and electron-lucent regions of 762 interest (ROIs). Statistical analyses of ROI measurements were performed in GraphPad 763 Prism (version 8) software. The experimenter was blinded to genotype for image 764 acquisition and analysis.

765 iBioID Sample Preparation

AAV2/9 viral probes, hSyn1-WASH1-BioID2-HA or hSyn1-solubleBioID2-HA, were injected into wild-type CD1 mouse brains using a Hamilton syringe (#7635-01) at age P0-P1 to ensure viral spread throughout the forebrain(Glascock et al., 2011). 15 days postviral injection, biotin was subcutaneously administered at 24mg/kg for seven consecutive days for biotinylation of proteins in proximity to BioID2 probes. Whole brains were extracted on the final day of biotin injections, snap frozen, and stored in liquid nitrogen

until protein purification. Seven brains were used for protein purification of each probe,
and each purification was performed three times independently (21 brains total for
WASH1-BioID2, 21 for solubleBioID2).

775 We performed all homogenization and protein purification on ice. A 2ml Dounce 776 homogenizer was used to individually homogenize each brain in a 1:1 solution of Lysis-777 R:2X-RIPA buffer solution with protease inhibitors (Roche cOmplete tablets 778 #11836153001). Each sample was sonicated three times for 7 seconds and then 779 centrifuged at 5000g for 5min at 4°C. Samples were transferred to Beckman Coulter 1.5ml 780 tubes (#344059), and then spun at 45,000rpm in a Beckman Coulter tabletop 781 ultracentrifuge (TLA-55 rotor) for 1hr at 4°C. SDS was added to supernatants (final 1%) 782 and samples were then boiled for 5min at 95°C. We next combined supernatants from 783 the same condition together (WASH1-BioID2 vs. solubleBioID2) in 15ml conical tubes to 784 rotate with 30µl high-capacity NeutrAvidin beads overnight at 4°C (Thermo #29204).

785 The following day, all steps were performed under a hood with keratin-free 786 reagents. Samples were spun down at 6000rpm, 4°C for 5min to pellet the beads and 787 remove supernatant. The pelleted beads then went through a series of washes, each for 788 10 min at RT with 500ul of solvent, and then spun down on a tabletop centrifuge to pellet 789 the beads for the next wash. The washes were as follows: 2% SDS twice, 1% TritonX100-790 1%deoxycholate-25mM LiCl2 once, 1M NaCL twice, 50mM Ammonium Bicarbonate 791 (Ambic) five times. Beads were then mixed 1:1 with a 2X Laemmli sample buffer that 792 contained 3mM biotin/50mM Ambic, boiled for 5 mins at 95°C, vortexed three times, and 793 then biotinylated protein supernatants were stored at -80°C until LC-MS/MS.

794 LC-MS/MS for iBioID

795 We gave the Duke Proteomics and Metabolomics Shared Resource (DPMSR) six 796 eluents from streptavidin resins (3 x WASH1-BioID2, 3 x solubleBioID2), stored on dry 797 ice. Samples were reduced with 10 mM dithiolthreitol for 30 min at 80°C and alkylated 798 with 20 mM iodoacetamide for 30 min at room temperature. Next, samples were 799 supplemented with a final concentration of 1.2% phosphoric acid and 256 µL of S-Trap 800 (Protifi) binding buffer (90% MeOH/100mM TEAB). Proteins were trapped on the S-Trap, 801 digested using 20 ng/µl sequencing grade trypsin (Promega) for 1 hr at 47°C, and eluted 802 using 50 mM TEAB, followed by 0.2% FA, and lastly using 50% ACN/0.2% FA. All 803 samples were then lyophilized to dryness and resuspended in 20 µL 1%TFA/2% 804 acetonitrile containing 25 fmol/µL yeast alcohol dehydrogenase (UniProtKB P00330; 805 ADH YEAST). From each sample, 3 µL was removed to create a pooled QC sample 806 (SPQC) which was run analyzed in technical triplicate throughout the acquisition period.

807 Quantitative LC/MS/MS was performed on 2 µL of each sample, using a 808 nanoAcquity UPLC system (Waters) coupled to a Thermo QExactive HF-X high resolution 809 accurate mass tandem mass spectrometer (Thermo) via a nanoelectrospray ionization 810 source. Briefly, the sample was first trapped on a Symmetry C18 20 mm × 180 µm 811 trapping column (5 µl/min at 99.9/0.1 v/v water/acetonitrile), after which the analytical 812 separation was performed using a 1.8 µm Acquity HSS T3 C18 75 µm × 250 mm column 813 (Waters) with a 90-min linear gradient of 5 to 30% acetonitrile with 0.1% formic acid at a 814 flow rate of 400 nanoliters/minute (nL/min) with a column temperature of 55°C. Data 815 collection on the QExactive HF-X mass spectrometer was performed in a data-dependent 816 acquisition (DDA) mode of acquisition with a r=120,000 (@ m/z 200) full MS scan from m/z 375 – 1600 with a target AGC value of 3e⁶ ions followed by 30 MS/MS scans at 817

r=15,000 (@ m/z 200) at a target AGC value of 5e⁴ ions and 45 ms. A 20s dynamic
exclusion was employed to increase depth of coverage. The total analysis cycle time for
each sample injection was approximately 2 hours.

821 LOPIT-DC Subcellular Fractionation

822 We performed three independent fractionation experiments with one adult SWIP 823 mutant brain and one WT mouse brain fractionated in each experiment. Each mouse was 824 sacrificed by isoflurane inhalation and its brain was immediately extracted and placed into 825 a 2ml Dounce homogenizer on ice with 1ml isotonic TEVP homogenization buffer (320mM 826 sucrose, 10mM Tris base, 1mM EDTA, 1mM EGTA, 5mM NaF, pH7.4 (Hallett et al., 827 2008)). A cOmplete mini protease inhibitor cocktail tablet (Sigma #11836170001) was 828 added to a 50ml TEVP buffer aliquot immediately before use. Brains were homogenized 829 for 15 passes with a Dounce homogenizer to break the tissue, and then this lysate was 830 brought up to a 5ml volume with additional TEVP buffer. Lysates were then passed 831 through a 0.5ml ball-bearing homogenizer for two passes (14 µm ball, Isobiotec) to 832 release organelles. Final brain lysate volumes were approximately 7.5ml each. Lysates 833 were then divided into replicate microfuge tubes (Beckman Coulter #357448) to perform 834 differential centrifugation, following Geladaki et. al's LOPIT-DC protocol(Geladaki et al., 835 2019). Centrifugation was carried out at 4°C in a tabletop Eppendorf 5424 centrifuge for 836 spins at 200g, 1,000g, 3,000g, 5,000g, 9,000g, 12,000g, and 15,000g. To isolate the final 837 three fractions, a tabletop Beckman TLA-100 ultracentrifuge with a TLA-55 rotor was used 838 at 4°C with speeds of: 30,000g, 79,000g, and 120,000g, respectively. Samples were kept 839 on ice at all times and pellets were stored at -80°C. Pellets from seven fractions (5,000g-840 120,000g) were used for proteomic analyses.

841 **16-plex TMT LC-MS/MS**

842 The Duke Proteomics and Metabolomics Shared Resource (DPMSR) processed 843 and prepared fraction pellets from all 42 frozen samples simultaneously (7 fractions per 844 brain from 3 WT and 3 MUT brains). Due to volume constraints, each sample was split 845 into 3 tubes, for a total of 126 samples, which were processed in the following manner: 846 100µL of 8M Urea was added to the first aliguot then probe sonicated for 5 seconds with 847 an energy setting of 30%. This volume was then transferred to the second and then third 848 aliquot after sonication in the same manner. All tubes were centrifuged at 10,000g and 849 any residual volume from tubes 1 and 2 were added to tube 3. Protein concentrations 850 were determined by BCA on the supernatant in duplicate (5 µL each assay). Total protein 851 concentrations for each replicate ranged from 1.1 mg/mL to 7.8 mg/mL with total protein 852 quantities ranging from 108.3 to 740.81 µg. 60 µg of each sample was removed and 853 normalized to 52.6µL with 8M Urea and 14.6µL 20% SDS. Samples were reduced with 854 10 mM dithiolthreitol for 30 min at 80°C and alkylated with 20 mM iodoacetamide for 30 855 min at room temperature. Next, they were supplemented with 7.4 µL of 12% phosphoric 856 acid, and 574 µL of S-Trap (Protifi) binding buffer (90% MeOH/100mM TEAB). Proteins 857 were trapped on the S-Trap, digested using 20 ng/ μ l sequencing grade trypsin (Promega) 858 for 1 hr at 47°C, and eluted using 50 mM TEAB, followed by 0.2% FA, and lastly using 859 50% ACN/0.2% FA. All samples were then lyophilized to dryness.

Each sample was resuspended in 120 μL 200 mM triethylammonium bicarbonate,
pH 8.0 (TEAB). From each sample, 20μL was removed and combined to form a pooled
quality control sample (SPQC). Fresh TMTPro reagent (0.5 mg for each 16-plex reagent)
was resuspended in 20 μL 100% acetonitrile (ACN) and was added to each sample.

Samples were incubated for 1 hour at RT. After the 1-hour reaction, 5 µL of 5% hydroxylamine was added and incubated for 15 minutes at room temperature to quench the reaction. Each 16-plex TMT experiment consisted of the WT and MUT fractions from one mouse, as well as the 2 SPQC samples. Samples corresponding to each experiment were concatenated and lyophilized to dryness.

869 Samples were resuspended in 800µL 0.1% formic acid. 400µg was fractionated 870 into 48 unique high pH reversed-phase fractions using pH 9.0 20 mM Ammonium formate 871 as mobile phase A and neat acetonitrile as mobile phase B. The column used was a 2.1 872 mm x 50 mm XBridge C18 (Waters) and fractionation was performed on an Agilent 1100 873 HPLC with G1364C fraction collector. Throughout the method, the flow rate was 0.4 874 mL/min and the column temperature was 55°C. The gradient method was set as follows: 875 0 min, 3%B; 1 min, 7% B; 50 min, 50%B; 51 min, 90% B; 55 min, 90% B; 56 min, 3% B; 876 70 min, 3% B. 48 fractions were collected in equal time segments from 0 to 52 minutes, 877 then concatenated into 12 unique samples using every 12th fraction. For instance, 878 fraction 1, 13, 25, and 37 were combined, fraction 2, 14, 26, and 38 were combined, etc. 879 Fractions were frozen and lyophilized overnight. Samples were resuspended in 66 µL 880 1%TFA/2% acetonitrile prior to LC-MS analysis.

Quantitative LC/MS/MS was performed on 2 μ L (1 μ g) of each sample, using a nanoAcquity UPLC system (Waters) coupled to a Thermo Orbitrap Fusion Lumos high resolution accurate mass tandem mass spectrometer (Thermo) equipped with a FAIMS Pro ion-mobility device via a nanoelectrospray ionization source. Briefly, the sample was first trapped on a Symmetry C18 20 mm × 180 μ m trapping column (5 μ l/min at 99.9/0.1 v/v water/acetonitrile), after which the analytical separation was performed using a 1.8

887 µm Acquity HSS T3 C18 75 µm × 250 mm column (Waters) with a 90-min linear gradient 888 of 5 to 30% acetonitrile with 0.1% formic acid at a flow rate of 400 nanoliters/minute 889 (nL/min) with a column temperature of 55°C. Data collection on the Fusion Lumos mass 890 spectrometer was performed for three different compensation voltages (CV: -40v, -60v, -891 80v). Within each CV, a data-dependent acquisition (DDA) mode of acquisition with a 892 r=120,000 (@ m/z 200) full MS scan from m/z 375 - 1600 with a target AGC value of $4e^5$ 893 ions was performed. MS/MS scans were acquired in the Orbitrap at r=50,000 (@ m/z 200) 894 from m/z 100 with a target AGC value of 1e⁵ and max fill time of 105 ms. The total cycle 895 time for each CV was 1s, with total cycle times of 3 sec between like full MS scans. A 45s 896 dynamic exclusion was employed to increase depth of coverage. The total analysis cycle 897 time for each sample injection was approximately 2 hours.

898 Following UPLC-MS/MS analyses, data were imported into Proteome Discoverer 899 2.4 (Thermo Scientific). The MS/MS data were searched against a SwissProt Mouse 900 database (downloaded November 2019) plus additional common contaminant proteins, 901 including yeast alcohol dehydrogenase (ADH), bovine casein, bovine serum albumin, as 902 well as an equal number of reversed-sequence "decoys" for FDR determination. Mascot 903 Distiller and Mascot Server (v 2.5, Matrix Sciences) were utilized to produce fragment ion 904 spectra and to perform the database searches. Database search parameters included 905 fixed modification on Cys (carbamidomethyl) and variable modification on Met (oxidation), 906 Asn/Gln (deamindation), Lys (TMTPro) and peptide N-termini (TMTPro). Data were 907 searched at 5 ppm precursor and 0.02 product mass accuracy with full trypsin enzyme 908 rules. Reporter ion intensities were calculated using the Reporter lons Quantifier

algorithm in Proteome Discoverer. Percolator node in Proteome Discoverer was used toannotate the data at a maximum 1% protein FDR.

911 Mouse Behavioral Assays

912 Behavioral tests were performed on age-matched WT and homozygous 913 SWIP^{P1019R} mutant littermates. Male and female mice were used in all experiments. 914 Testing was performed at two time points: P42-55 days old as a young adult age, and 5.5 915 months old as mid-adulthood, so that we could compare disease progression in this 916 mouse model to human patients (Ropers et al., 2011). The sequence of behavioral testing 917 was: Y-maze (to measure working memory), object novelty recognition (to measure short-918 term and long-term object recognition memory), TreadScan (to assess gait), and steady-919 speed rotarod (to assess motor control and strength) for 40-55 day old mice. Testing was 920 performed over 1.5 weeks, interspersed with rest days for acclimation. This sequence 921 was repeated with the same cohort at 5.5-6 months old, with three additional measures 922 added to the end of testing: fear conditioning (to assess associative fear memory), a 923 hearing test (to measure tone response), and a shock threshold test (to assess 924 somatosensation). Of note, a separate, second cohort of mice was evaluated for fear 925 conditioning, hearing, and shock threshold testing at adolescence. After each trial, 926 equipment was cleaned with Labsan to remove residual odors. The experimenter was 927 blinded to genotype for all behavioral analyses.

928 **Y-maze**

Working memory was evaluated by measuring spontaneous alternations in a 3arm Y-maze under indirect illumination (80-90 lux). A mouse was placed in the center of the maze and allowed to freely explore all arms, each of which had different visual cues

for spatial recognition. Trials were 5 min in length, with video data and analyses captured by EthoVision XT 11.0 software (Noldus Information Technology). Entry to an arm was define as the mouse being >1 body length into a given arm. An alternation was defined as three successive entries into each of the different arms. Total % alternation was calculated as the total number of alternations/the total number of arm entries minus 2 x100.

938 Novel Object Recognition

939 One hour before testing, mice were individually exposed to the testing arena (a 48 940 x 22 x 18cm white opague arena) for 10min under 80-100lux illumination without any 941 objects. The test consisted of three phases: training (day 1), short-term memory test 942 (STM, day 1), and long-term memory test (LTM, day 2). For the training phase, two 943 identical objects were placed 10 cm apart, against opposing walls of the arena. A mouse 944 was placed in the center of the arena and given full access to explore both objects for 5 min and then returned to its home cage. For STM testing, one of the training objects 945 946 remained (the now familiar object), and a novel object replaced one of the training objects 947 (similar in size, different shape). The mouse was returned to the arena 30 minutes after 948 the training task and allowed to explore freely for 5 mins. For LTM testing, the novel object 949 was replaced with another object, and the familiar object remained unchanged. The LTM 950 test was also 5 min in duration, conducted 24hr after the training task. Behavior was 951 scored using Ethovision 11.0 XT software (Noldus) and analyzed by a blind observer. 952 Object contact was defined as the mouse's nose within 1 cm of the object. We analyzed 953 both number of nose contacts with each object and duration of contacts. Preference 954 scores were calculated as (duration contact_{novel} - duration contact_{familiar}) / total duration

contact_{novel+familiar}. Positive scores signified a preference for the novel object; whereas,
negative scores denoted a preference for the familiar object, and scores approaching
zero indicated no preference.

958 TreadScan

959 A TreadScan forced locomotion treadmill system (CleversSys Inc, Reston, 960 Virginia) was used for gait recording and analysis. Each mouse was recorded walking on 961 a transparent treadmill at 45 days old, and again at 5.5 months old. Mice were acclimated 962 to the treadmill chamber for 1 minute before the start of recording to eliminate exploratory 963 behavior confounding normal gait. Trials were 20 seconds in length, with mice walking at 964 speeds between 13.83 and 16.53 cm/sec (P45 WT average 15.74 cm/s; P45 MUT 965 average 15.80 cm/s; 5.5mo WT average 15.77 cm/s; 5.5mo MUT average 15.85 cm/s). 966 A high-speed digital camera attached to the treadmill captured limb movement at a frame 967 rate of 100 frames/second. We used TreadScan software (CleversSys) and 968 representative WT and MUT videos to generate footprint templates, which were then used 969 to identify individual paw profiles for each limb. Parameters such as stance time, swing 970 time, step length, track width, and limb coupling were recorded for the entire 20 sec 971 duration for each animal. Output gait tracking was verified manually by a blinded 972 experimenter to ensure consistent limb tracking throughout the duration of each video.

973 Steady Speed Rotarod

A 5-lane rotarod (Med Associates, St. Albans, VT) was used for steady-speed motor analysis. The rod was run at a steady speed of 32rpm for four, 5-minute trials, with a 40-minute inter-trial interval. We recorded mouse latency to fall by infrared beam break,

977 or manually for any mouse that completed two or more rotations on the rod without978 walking. Mice were randomized across lanes for each trial.

979 Fear Conditioning

980 Animals were examined in contextual and cued fear conditioning as described by 981 Rodriguiz and Wetsel (Rodriguiz and Wetsel, 2006). Two separate cohorts of mice were 982 used testing the two age groups. A three-day testing paradigm was used to assess 983 memory: conditioning on day 1, context testing 24-hr post-conditioning on day 2, and cued 984 tone testing 48hr post-conditioning on day 3. All testing was conducted in fear 985 conditioning chambers (Med Associates). In the conditioning phase, mice were first 986 acclimated to the test chamber for two minutes under ~100 lux illumination. Then a 987 2900Hz, 80dB tone (conditioned stimulus, CS) played for 30 sec, which terminated with 988 a paired 0.4mA, 2 sec scrambled foot shock (unconditioned stimulus, US). Mice were 989 removed from the chamber and returned to their home cage 30 sec later. In the context 990 testing phase, mice were placed in the same conditioning chamber and monitored for 991 freezing behavior for a 5 min trial period, in the absence of the CS and US. For cued tone 992 testing, the chambers were modified to different dimensions and shapes, contained 993 different floor and wall textures, and lighting was adjusted to 50 lux. Mice acclimated to 994 the chamber for 2 min, and then the CS was presented continuously for 3 min. Contextual 995 and cued fear memory was assessed by freezing behavior, captured by automated video 996 software (CleversSys).

997 Hearing Test

We tested mouse hearing using a startle platform (Med Associates) connected to
Startle Pro Software in a sound-proof chamber. Mice were placed in a ventilated restraint

1000 cylinder connected to the startle response detection system to measure startle to each 1001 acoustic stimulus. After two minutes of acclimation, mice were assessed for an acoustic 1002 startle response to seven different tone frequencies, 2kHz, 3kHz, 4kHz, 8kHz, 12kHz, 1003 16kHz, and 20kHz that were randomly presented three times each at four different 1004 decibels, 80, 100, 105, and 110dB, for a total of 84 trials. A random inter-trial interval of 1005 15-60 seconds (average 30sec) was used to prevent anticipation of a stimulus. An 1006 animal's reaction to the tone was recorded as startle reactivity in the first 100msec of the 1007 stimulus presentation, which was transduced through the platform's load cell and 1008 expressed in arbitrary units (AU).

1009 Startle Response (Somatosensation)

1010 Mouse somatosensation was tested by placing mice in a startle chamber (Med 1011 Associates) connected to Startle Pro Software. Mice were placed atop a multi-bar cradle 1012 within a ventilated plexiglass restraint cylinder, which allows for horizontal movement within the chamber, but not upright rearing. After two minutes of acclimation, each mouse 1013 1014 was exposed to 10 different scrambled shock intensities, ranging from 0 to 0.6mA with 1015 randomized inter-trial intervals of 20-90 seconds. Each animal's startle reactivity during 1016 the first 100 msec of the shock was transduced through the platform's load cell and 1017 recorded as area under the curve (AUC) in arbitrary units (AU).

1018

1019 QUANTIFICATION AND STATISTICAL ANALYSIS

1020 Experimental conditions, number of replicates, and statistical tests used are stated 1021 in each figure legend. Each experiment was replicated at least three times (or on at least 1022 3 separate animals) to assure rigor and reproducibility. Both male and female age-
1023 matched mice were used for all experiments, with data pooled from both sexes. Data 1024 compilation and statistical analyses for all non-proteomic data were performed using 1025 GraphPad Prism (version 8, GraphPad Software, CA), using a significance level of 1026 alpha=0.05. Prism provides exact p values unless p<0.0001. All data are reported as 1027 mean ± SEM. Each data set was tested for normal distribution using a D'Agostino-Person 1028 normality test to determine whether parametric (unpaired Student's t-test, one-way 1029 ANOVA, two-way ANOVA) or non-parametric (Mann-Whitney, Kruskal-Wallis) tests 1030 should be used. Parametric assumptions were confirmed with the Shapiro-Wilk test 1031 (normality) and Levine's test (error variance homogeneity) for ANOVA with repeated 1032 measures testing. The analysis of iBioID and TMT proteomics data are described below. 1033 All proteomic data and analysis scripts are available online (see Resource Availability).

1034 Imaris 3D reconstruction

1035 For EEA1+ and CathepsinD+ puncta analyses, coverslips were imaged on a Zeiss 1036 LSM 710 confocal microscope. Images were sampled at a resolution of 1024 x 1024 1037 pixels with a dwell time of 0.45 usec using a 63 x/1.4 oil immersion objective, a 2.0 times 1038 digital zoom, and a z-step size of 0.37 µm. Images were saved as ".lsm" formatted files, 1039 and quantification was performed on a POGO Velocity workstation in the Duke Light 1040 Microscopy Core Facility using Imaris 9.2.0 software (Bitplane, South Windsor, CT). For 1041 analyses, we first used the "surface" tool to make a solid fill surface of the MAP2-stained 1042 neuronal soma and dendrites, with the background subtraction option enabled. We 1043 selected a threshold that demarcated the neuron structure accurately while excluding 1044 background. For EEA1 puncta analyses, a 600 x 800 µm selection box was placed around 1045 the soma in each image and surfaces were created for EEA1 puncta within the selection

box. Similarly, for CathepsinD puncta analyses, a 600 x 600 µm selection box was placed around the soma(s) in each image for surface creation. The same threshold settings were used across all images, and individual surface data from each soma were exported for aggregate analyses. The experimenter was blinded to sample conditions for both image acquisition and analysis.

1051 Cleaved Caspase-3 Image Analysis

1052 Z-stack images were acquired on a Zeiss 710 LSM confocal microscope. Images 1053 were sampled at a resolution of 1024 x 1024 pixels with a dwell time of 1.58 using 1054 a 63x/1.4 oil immersion objective (for cortex, striatum, and hippocampus) or 20x/0.8 dry 1055 objective (cerebellum), a 1.0 times digital zoom, and a z-step size of 0.67 µm. Images 1056 were saved as ".lsm" formatted files, and then converted into maximum intensity 1057 projections (MIP) using Zen 2.3 SP1 software. Quantification of CC3 colocalization with 1058 DAPI was performed on the MIPs using the Particle Analyzer function in FIJI ImageJ 1059 software. The experimenter was blind to sample conditions for both image acquisition and 1060 analysis.

1061 iBioID Quantitative Analysis

Following UPLC-MS/MS analyses, data was imported into Proteome Discoverer 2.2 (Thermo Scientific Inc.), and aligned based on the accurate mass and retention time of detected ions ("features") using Minora Feature Detector algorithm in Proteome Discoverer. Relative peptide abundance was calculated based on area-under-the-curve (AUC) of the selected ion chromatograms of the aligned features across all runs. The MS/MS data was searched against the SwissProt *Mus musculus* database (downloaded in April 2018) with additional proteins, including yeast ADH1, bovine serum albumin, as 1069 well as an equal number of reversed-sequence "decoys" for false discovery rate (FDR) 1070 determination. Mascot Distiller and Mascot Server (v 2.5, Matrix Sciences) were utilized 1071 to produce fragment ion spectra and to perform the database searches. Database search 1072 parameters included fixed modification on Cys (carbamidomethyl), variable modifications 1073 on Meth (oxidation) and Asn and Gln (deamidation), and were searched at 5 ppm 1074 precursor and 0.02 Da product mass accuracy with full trypsin enzymatic rules. Peptide 1075 Validator and Protein FDR Validator nodes in Proteome Discoverer were used to annotate 1076 the data at a maximum 1% protein FDR.

1077 Protein intensities were exported from Proteome Discoverer and processed using 1078 custom R scripts. Carboxylases and keratins, as well as 315 mitochondrial proteins(Calvo 1079 et al., 2016), were removed from the identified proteins as known contaminants. Next, we 1080 performed sample loading normalization to account for technical variation between the 9 1081 individual MS runs. This is done by multiplying intensities from each MS run by a scaling 1082 factor, such that the average of all total run intensities are equal. As QC samples were 1083 created by pooling equivalent aliquots of peptides from each biological replicate, the 1084 average of all biological replicates should be equal to the average of all technical SPQC 1085 replicates. We performed sample pool normalization to SPQC samples to standardize 1086 protein measurements across all samples and correct for batch effects between MS 1087 analyses. Sample pool normalization adjusts the protein-wise mean of all biological 1088 replicates to be equal to the mean of all SPQC replicates. Finally, proteins that were 1089 identified by a single peptide, and/or identified in less than 50% of samples were removed. 1090 Any remaining missing values were inferred to be missing not at random due to the left 1091 shifted distribution of proteins with missing values and imputed using the k-nearest

1092 neighbors algorithm using the impute.knn function in the R package impute 1093 (impute::impute.knn). Normalized protein data was analyzed using edgeR, an R package 1094 for the analysis of differential expression/abundance that models count data using a 1095 binomial distribution methodology. Differential enrichment of proteins in the WASH1-1096 BioID2 pull-down relative to the solubleBioID2 control pull-down were evaluated with an 1097 exact test as implemented by the edgeR::exactTest function. To consider a protein 1098 enriched in the WASH interactome, we required that a protein exhibit a fold change greater than 3 over the negative control with an exact test Benjamini Hochberg adjusted 1099 1100 p-value (FDR) less than 0.1. With these criteria, 174 proteins were identified as WASH1 1101 interactome proteins. Raw peptide and final normalized protein data as well as the 1102 statistical results can be found in Table S1.

1103 Proteins that function together often interact directly. We compiled experimentally-1104 determined protein-protein interactions (PPIs) among the WASH1 interactome from the 1105 HitPredict database(López et al., 2015) using a custom R package, getPPIs, (available 1106 online at twesleyb/getPPIs). We report PPIs among the WASH1 interactome in Table S1. 1107 Bioinformatic GO analysis was conducted by manual annotation of identified 1108 proteins and confirmed with Metascape analysis(Zhou et al., 2019) of WASH1-BioID2 1109 enriched proteins using the 2,311 proteins identified in the mass spec analysis as 1110 background.

1111 Raw peptide intensities were exported from Proteome Discover for downstream 1112 analysis and processing in R. Following database searching, protein scoring using the 1113 Protein FDR Validator algorithm, and removal of contaminant species, the dataset

retained 86,551 peptides corresponding to the identification of 7,488 unique proteins.

1115 These data, as well as statistical results can be found in Table S2.

1116 **TMT Proteomics Quantitative Analysis**

1117 Peptide level data from the spatial proteomics analysis of SWIP^{P1019R} MUT and 1118 MUT brain were exported from Proteome Discoverer (version 2.4) and analyzed using 1119 custom R and Python scripts. Peptides from contaminant and non-mouse proteins were 1120 removed. First, we performed sample loading normalization, normalizing the total ion 1121 intensity for each TMT channel within an experiment to be equal. Sample loading 1122 normalization corrects for small differences in the amount of sample analyzed and 1123 labeling reaction efficiency differences between individual TMT channels within an 1124 experiment.

We found that in each TMT experiment there were a small number of missing values (mean percent missing = 1.6 +/- 0.17%). Missing values were inferred to be missing at random based on the overlapping distributions of peptides with missing values and peptides without missing values. We imputed these missing values using the knearest neighbor algorithm (impute::impute.knn). Missing values for SPQC samples were not imputed. Peptides with any missing SPQC data were removed.

Following sample loading normalization, SPQC replicates within each experiment should yield identical measurements. As peptides with irreproducible QC measurements are unlikely to be quantitatively robust, and their inclusion may bias downstream processing (see IRS normalization below), we sought to remove them. To assess intrabatch variability, we utilized the method described by Ping et al., 2019(Ping et al., 2018). Briefly, peptides were binned into 5 groups based on the average intensity of the two SPQC replicates. For each pair of SPQC measurements, the log ratio of SPQC intensities was calculated. To identify outlier QC peptides, we plotted the distribution of these log ratios for each bin. Peptides with ratios that were more than four standard deviations away from the mean of its intensity bin were considered outliers and removed (Total number of SPQC outlier peptides removed = 474).

1142 Proteins were summarized as the sum of all unique peptide intensities 1143 corresponding to a unique UniProtKB Accession identifier, and sample loading 1144 normalization was performed across all three experiments to account for inter-1145 experimental technical variability. In a TMT experiment, the peptides selected for MS2 1146 fragmentation for any given protein is partially random, especially at lower signal-to-noise 1147 peptides. This stochasticity means that proteins are typically quantified by different 1148 peptides in each experiment. Thus, although SPQC samples should yield identical protein 1149 measurements in each of the three experiments (as it is the same sample analyzed in 1150 each experiment), the observed protein measurements exhibit variability due to their 1151 quantification by different peptides. To account for this protein-level bias, we utilized the 1152 internal reference scaling (IRS) approach described by Plubell et al., 2017(Plubell et al., 1153 2017). IRS normalization scales the protein-wise geometric average of all SPQC 1154 measurements across all experiments to be equal, and simultaneously adjusts biological 1155 replicates. In brief, each protein is multiplied by a scaling factor which adjusts its intra-1156 experimental SPQC values to be equal to the geometric mean of all SPQC values for the 1157 normalization effectively standardizes three experiments. This step protein 1158 measurements between different mass spectrometry experiments.

1159 The final normalization step was to perform sample pool normalization using 1160 SPQC samples as a reference. This normalization step, sometimes referred to as global 1161 internal standard normalization, accounts for batch effects between experiments, and 1162 reflects the fact that after technical normalization, the mean of biological replicates should 1163 be equal to the mean of SPQC replicates.

1164 Before assessing protein differential abundance, we removed irreproducible 1165 proteins. This included proteins that were quantified in less than 50% of all samples, 1166 proteins that were identified by a single peptide, and proteins that had missing SPQC 1167 values. Across all 42 biological replicates, we observed that a small number of proteins 1168 had potential outlier measurements that were either several orders of magnitude greater 1169 or less than the mean of its replicates. In order to identify and remove these proteins, we 1170 assessed the reproducibility of protein measurements within a fraction in the same 1171 manner used to identify and filter SPQC outlier peptides. A small number of proteins were 1172 identified as outliers if the average log ratio of their 3 technical replicates was more than 1173 4 standard deviations away from the mean of its intensity bin (n=349). In total, we retained 1174 5,897 of the original 7,488 proteins in the final dataset.

Differential protein abundance was assessed using the final normalized protein data for intrafraction comparisons between WT and MUT groups using a general linear model as implemented by the edgeR::glmQLFit and edgeR::glmQLFTest functions(MD et al., 2009). Although this approach was originally devised for analysis of single-cell RNA-sequencing data, this approach is also appropriate for proteomics count data which is over-dispersed, negative binomially distributed, and often only includes a small number of replicates (for an example of edgeR's application to proteomics see Plubell et al.,

2017(Plubell et al., 2017))(McCarthy et al., 2012; MD et al., 2009). For intrafraction
comparisons, P-values were corrected using the Benjamini Hochberg procedure within
edgeR. An FDR threshold of 0.1 was set for significance for intrafraction comparisons.

We utilized edgeR's flexible GLM framework to test the hypothesis that the abundance of proteins in the WT group was significantly different from that in the MUT group irrespective of fraction differences (Table S2). For WT vs. MUT contrasts, we considered proteins with an FDR < 0.05 significant (n=687). For plotting, we adjusted normalized protein abundances for fraction differences by fitting the data with an additive linear model with fraction as a blocking factor, as implemented by the removeBatchEffect algorithm from the R limma package(Ritchie et al., 2015).

1192 To construct a protein covariation graph, we assessed the pairwise covariation 1193 (correlation) between all 5,897 proteins using the biweight midcorrelation 1194 (WGCNA::bicor) statistic(Seyfried et al., 2017), a robust alternative to Pearson's 1195 correlation. The resulting complete, signed, weighted, and symmetric adjacency matrix 1196 was then re-weighted using the 'Network Enhancement' approach. Network 1197 enhancement removes noise from the graph, and facilitates downstream community 1198 detection(Wang et al., 2018).

The enhanced adjacency matrix was clustered using the Leiden algorithm(Traag et al., 2019), a recent extension and improvement of the well-known Louvain algorithm(Mucha et al., 2010). The Leiden algorithm functions to optimize the partition of a graph into modules by maximizing a quality statistic. We utilized the 'Surprise' quality statistic(Traag et al., 2015) to identify optimal partitions of the protein covariation graph. To facilitate biological inferences drawn from the network's organization, we recursively

split 27 modules that contained more than 100 nodes and removed modules that were
smaller than 5 proteins. Initial clustering of the network resulted in the identification of 324
modules.

1208 To reduce the likelihood of identifying false positive modules, we enforced module 1209 guality using a permutation procedure (NetRep::modulePreservation)(Ritchie et al., 2016) 1210 and removed modules with any insignificant permutation statistics (Bonferroni P-Adjust > 1211 0.05). The following statistics were used to enforce module quality: 'avg.weight' (average 1212 edge weight), 'avg.cor' (average bicor correlation R²), and 'avg.contrib' (guantifies how 1213 similar an individual protein's abundance profile is to the summary of its module). Proteins 1214 which were assigned to modules with insignificant module guality statistics were not 1215 considered clustered as the observed quality of their module does not differ from random. 1216 After filtering, approximately 85% of all proteins were assigned a cluster. The median 1217 percent variance explained by the first principle component of a module (a measure of 1218 module cohesiveness) was high (59.8%). After removal of low-guality modules, the 1219 analysis retained 255 distinct modules of proteins that strongly covaried together (Table 1220 S3).

To evaluate modules that were changing between WT and MUT genotypes, we extended the GLM framework to test for protein differential abundance. Modules were summarized as the sum of their proteins and fit with a GLM, with fraction as a blocking factor. In this statistical design, we were interested in the average effect of genotype on all proteins in a module. For plotting, module abundance was adjusted for fraction differences using the removeBatchEffect function (package: limma). We utilized the

Bonferroni method to adjust P-values for 255 module level comparisons and considered modules with an adjusted P-value less than 0.05 were considered significant (n=37).

1229 Module Gene Set Enrichment Analysis

1230 Modules were analyzed for enrichment of the WASH interactome (this paper), 1231 Retriever complex (McNally et al., 2017), CORUM protein complexes (Giurgiu et al., 1232 2019), and subcellular predictions generated by Geladaki et al. (Geladaki et al., 2019) 1233 using the hypergeometric test with Bonferroni P-value correction for multiple comparisons. The union of all clustered and pathway proteins was used as background 1234 1235 for the hypergeometric test. In addition to analysis of these general cellular pathways, we 1236 analyzed modules for enrichment of neuron-specific subcellular compartments-this 1237 included the presynapse (Takamori et al., 2006), excitatory postsynapse (Uezu et al., 1238 2016), and inhibitory postsynapse (Uezu et al., 2016). These gene lists are available 1239 online https://github.com/twesleyb/geneLists. at

1240 Network Visualization

1241 Network graphs were visualized in Cytoscape (Version 3.7.2). We used the 1242 Perfuse Force Directed Layout (weight = edge weight). In this layout, strongly connected 1243 nodes tend to be positioned closer together. In some instances, node location was 1244 manually adjusted to visualize the module more compactly. Node size was set to be 1245 proportional to the weighted degree centrality of a node in its module subgraph. Node 1246 size thus reflects node importance in the module. Visualizing co-expression or co-1247 variation networks is challenging because every node is connected to every other node 1248 (the graph is complete). To aid visualization of module topology, we removed weak edges 1249 from the graphs. A threshold for each module was set to remove the maximal number of

1250	edges before the module subgraph split into multiple components. This strategy enables
1251	visualization of the strongest paths in a network.
1252	
1253	SUPPLEMENTARY FILES
1254	Supplementary File 1: Representative modules from major subcellular
1255	compartments.
1256	Supplementary File 2: Source data for Western blots.
1257	• Table S1: WASH iBioID raw and normalized data and the corresponding
1258	statistical results.
1259	• Table S2: SWIP ^{P1019R} TMT raw and normalized proteomics data and the
1260	corresponding statistical results.
1261	• Table S3: Module-level data and statistical results from network analysis of
1262	SWIP ^{P109R} proteomics.
1263	• Table S4: DNA oligonucleotides used in this study.
1264	• Table S5: Key resources.
1265	
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Supplementary File 1. Representative modules from major subcellular compartments.



Subcellular compartment with modules exhibiting significant changes in abundance between WT and SWIP MUT brain.

Subcellular compartments with no significant difference between WT and MUT brain.

Figure 1. Schematic overview of major subcellular compartments in neurons. SWIP^{P1019R} MUT mouse brain exhibited significant differences in endosomal, lysosomal, ER, and postsynaptic modules compared to WT. Each of the following figures contains (A) normalized protein abundance for all proteins in a module, (B) schematic of the predicted subcellular compartment, and (C) network subgraph vizualization of a module. These modules, which do not exhibit a significant difference between WT and SWIP^{P1019R} brain, support the hypothesis that the primary cellular deficit resulting from SWIP^{P1019R} is dysregulation of endo-lysosomal pathways.

Module 8: Golgi Apparatus



Figure 2. No significant difference in a predicted golgi apparatus module. (A) normalized protein abundance for all proteins in M8, which is predicted to represent the golgi apparatus based on its significant enrichment of golgi proteins from Geladaki *et al.*, (2019). (B) schematic representation of the golgi apparatus. (C) M8 subnetwork showing organization of M8 proteins including 10 out of the 13 golgi proteins quantified in this study. Note the presence of MAN1A2 a mannosidase which resides in the golgi and functions in protein glycoslyation.

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M11 network graph.



Figure 4. No significant difference in a predicted mitochondrial module.
(A) normalized protein abundance for all proteins in M27, predicted to have mitochondrial function by its enrichment of mitochondrial proteins (Geladaki *et al.*, 2019).
(B) schematic of the neuronal mitochondria found in the presynapse. (C) M27 network which contains 38 of the of the 261 mitochondrial proteins quantified in this study.


Figure 5. No significant difference in a predicted cytoplasmic module. (A) normalized protein abundance for all proteins in M45, predicted to have cytoplasmic localization. (B) schematic of the predicted cytoplasmic compartment, and (C) M45 network. M45 contained 9 of the 239 cytoplasmic proteins (Geladaki *et al.*, 2019) quantified in this study.



Figure 6. No significant difference in a predicted ER module.

(A) normalized protein abundance for all proteins in M81, predicted to represent the endoplasmic reticulum (ER) based on its enrichment of ER proteins (Geladaki et al., 2006). Note that unlike M83 (Fig. S3), there is no significant difference between MUT WT brain. (B) Schematic of the predicted ER compartment, and (C) M81 network. M81 contains 34 ER proteins incuding Erlin1/2 which function in a complex as a part of the ERAD pathway.



Figure 7. No significant difference in a predicted nuclear module. (A) normalized protein abundance for all proteins in M103, a predicted nuclear module. (B) schematic of the predicted nuclear compartment, and (C) M103 network. M103 contains 26 resident nuclear proteins, including LSM4 and LSM4, which function in spliceosome assembly.

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Figure 8. No significant difference in a predicted inhibitory postsynaptic density (iPSD) module. (A) normalized protein abundance for all proteins in M155, a predicted iPSD module. (B) schematic of the predicted iPSD compartment, and (C) M155 network. M155 contains 5 iPSD proteins including two GABA_AR subunits and one GABA_BR subunit identified by Uezu *et al.*, 2016.



Figure 9. No significant difference in a predicted ribosome module.

(A) normalized protein abundance for all proteins in M164, predicted to have ribosomal function by enrichment of ribosome proteins (Geladaki *et al.*, 2019).
(B) schematic of the predicted ribosomal compartment, and (C) M164 network.



Figure 10. No significant difference in a predicted proteosome module. (A) normalized protein abundance for all proteins in M184, predicted to represent the proteasome. (B) schematic of the predicted proteosomal compartment, and (C) M184 network. M184 contained 11 out of the 26 proteosomal proteins from Geladaki *et al.*, (2019) that were quantified in this study.



Figure 11. No significant difference in a predicted excitatory postsynapse (ePSD) module. (A) normalized protein abundance for all proteins in M206, predicted to have postsynaptic function by enrichment of proteins known to reside in the excitatory postsynapse. (B) schematic of the predicted ePSD compartment, and (C) M206 network. M206 contained 7 out of the 138 ePSD proteins reported by Uezu *et al.*, (2016) and is also enriched for proteins identified by WASH-BioID (this study). In contrast to modules M35 and M143, which exhibited decreased abundance in SWIP MUT brain, M206 does not exhibit a global difference between WT and MUT brain.



Figure 12. No significant difference in a predicted peroxisome module. (A) normalized protein abundance for all proteins in M209, predicted to represent the peroxisome based on its enrichment of peroxisomal proteins (Geladaki et al., 2019). (B) Schematic of the peroxisome and (C) network of the M209 module which contains 11 of the 22 peroxisomal proteins quantified in this study.