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Dissecting the Physiological Roles of ULK1/2 in the Mouse Brain

Bo Wang
University of Tennessee Health Science Center

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Dissecting the Physiological Roles of ULK1/2 in the Mouse Brain

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Dissecting the Physiological Roles of ULK1/2 in the Mouse Brain

A Dissertation
Presented for
The Graduate Studies Council
The University of Tennessee
Health Science Center

In Partial Fulfillment
Of the Requirements for the Degree
Doctor of Philosophy
From The University of Tennessee

By
Bo Wang
December 2016
ACKNOWLEDGEMENTS

The past 5 years has been an amazing adventure. As an international student, this experience of working towards doctoral degree in a foreign country is unforgettable and priceless. In retrospect, I am full of appreciation to those who have facilitated my study in the United States. My greatest and most sincere thanks go to my mentor, Dr. Mondira Kundu. She guided me through the past four and a half years, and built me in various different ways. As a mentor, she is extremely patient, encouraging, supportive, and instrumental. And because of that, I had the chance to explore many scientific issues that interested me. A student really couldn’t ask for more from a mentor. More importantly, her passion and meticulousness in science inspired me tremendously to pursue a career in academia in the past and in the future. Although I will soon leave her laboratory, the skills that I learned here will always stay with me, and for that, she deserves every single credit.

I have also had the privilege of working with all the colleagues in the lab. They are extremely patient with me and literally helped me a million times. Often, they prioritized my needs and carried me through hard times, without which I couldn’t fulfill my goals here. They are happy for my achievement and sorry for my loss, making me feel at least I am not alone here in the United States. We share countless memories and moments, and I will carry those wherever I will be in the future. You will be deeply missed.

I want to thank all the people along the ways. My committee members including: Dr. Kristin M. Hamre, Dr. Joseph T. Opferman, Dr. David J. Solecki, and Dr. Paul J. Taylor have provided significant insight into my research. They devoted their time and effort generously and guided me through all the years until now. Their guidance will be my lifelong treasure.

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ABSTRACT

Mammalian UNC-51–like kinases 1 and 2 (ULK1 and ULK2), Caenorhabditis elegans UNC-51 and Drosophila melanogaster Atg1 are redundant serine/threonine kinases that regulate flux through the autophagy pathway in response to various types of cellular stress. C. elegans UNC-51 and D. melanogaster Atg1 also promote axonal growth and defasciculation, and disruption of these genes results in defects in axon guidance in invertebrates. Germline Ulk1/2-deficient mice die perinatally. Therefore, we used a conditional-knockout approach to investigate the roles of ULK1/2 in the brain. Mice lacking Ulk1 and Ulk2 in their central nervous systems (CNS) showed defects in axonal pathfinding and defasciculation affecting the corpus callosum (CC), anterior commissure (AC), corticothalamic axons (CTAs) and thalamocortical axons (TCAs) and mossy fibers. These defects led to impaired midline crossing of callosal axons, anterior commissure hypoplasia and disorganization of the somatosensory cortex. The axon guidance defects observed in Ulk1/2 double knockout (dko) and in CNS-specific (Nestin-Cre) Ulk1/2 conditional double knockout (cdko) mice were not recapitulated in mice lacking other autophagy genes (i.e. Atg7 or Fip200), and was associated with abnormal localization of the axon guidance molecule, transient axonal glycoprotein-1 (TAG-1) in the distal CTAs. Approximately 40% of the Ulk1/2 cdko animals died shortly after birth; the remaining animals survived up to 4 months. Although the mice showed neuronal degeneration, specifically in the hippocampal CA1 region, the neurons showed no accumulation of P62+/ubiquitin+ inclusions or abnormal membranous structures, which are observed in mice lacking other autophagy genes, such as Atg7, and Fip200. Rather, neuronal death was associated with activation of the unfolded protein response (UPR) pathway. An unbiased proteomics approach identified SEC16A as a novel ULK1/2-interacting partner. ULK-mediated phosphorylation of SEC16A regulated the assembly of endoplasmic reticulum (ER) exit sites and ER-to-Golgi trafficking of specific cargo such as, the serotonin transporter SERT, and did not require other autophagy proteins (e.g. ATG13). The defect in ER-to-Golgi trafficking activated the UPR pathway in ULK-deficient cells; both processes were reversed upon expression of SEC16A with a phosphomimetic substitution. Thus, the regulation of ER-to-Golgi trafficking by ULK1/2 is essential for cellular homeostasis. Moreover, the defect in SERT trafficking may also contribute to the disrupted formation of the barrel cortex in the Ulk1/2 cdko mice. Together, these data highlight the autophagy-independent role of ULK1 and ULK2 in maintaining cellular homeostasis and regulating axon guidance in the mammalian brain.
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<th>Full Form</th>
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<tbody>
<tr>
<td>AC</td>
<td>Anterior commissure</td>
</tr>
<tr>
<td>AMPK</td>
<td>5' AMP-activated protein kinase</td>
</tr>
<tr>
<td>ATF6</td>
<td>Activating Transcription Factor 6</td>
</tr>
<tr>
<td>ATG5</td>
<td>Autophagy related 5</td>
</tr>
<tr>
<td>ATG7</td>
<td>Autophagy related 7</td>
</tr>
<tr>
<td>ATG13</td>
<td>Autophagy Related 13</td>
</tr>
<tr>
<td>ATG14</td>
<td>Autophagy Related 14</td>
</tr>
<tr>
<td>BiP</td>
<td>Binding immunoglobulin protein</td>
</tr>
<tr>
<td>BP</td>
<td>Blocking peptide</td>
</tr>
<tr>
<td>CA1</td>
<td>Region I of hippocampus proper</td>
</tr>
<tr>
<td>C. elegans</td>
<td>Caenorhabditis elegans</td>
</tr>
<tr>
<td>CC</td>
<td>Corpus callosum</td>
</tr>
<tr>
<td>cdko</td>
<td>Conditional double knockout</td>
</tr>
<tr>
<td>cGAMP</td>
<td>Cyclic guanosine monophosphate–adenosine monophosphate</td>
</tr>
<tr>
<td>cGAS</td>
<td>cGAMP synthase</td>
</tr>
<tr>
<td>CHOP</td>
<td>DNA damage-inducible transcript 3</td>
</tr>
<tr>
<td>CNS</td>
<td>Central nervous system</td>
</tr>
<tr>
<td>CO</td>
<td>Cytochrome oxidase</td>
</tr>
<tr>
<td>COPII</td>
<td>Coat protein complex II</td>
</tr>
<tr>
<td>CTA</td>
<td>Corticothalamic axon</td>
</tr>
<tr>
<td>Cvt pathway</td>
<td>Cytoplasm-to-vacuole targeting</td>
</tr>
<tr>
<td>DH</td>
<td>Dendate hilus</td>
</tr>
<tr>
<td>Abbreviation</td>
<td>Full Name</td>
</tr>
<tr>
<td>--------------</td>
<td>-----------</td>
</tr>
<tr>
<td>DiI</td>
<td>1,1’-dioctadecyl-3,3,38,38, tetramethylindocarbocyanine perchlorate</td>
</tr>
<tr>
<td>D. melanogaster</td>
<td><em>Drosophila melanogaster</em></td>
</tr>
<tr>
<td>dTh</td>
<td>Dorsal thalamus</td>
</tr>
<tr>
<td>EC</td>
<td>External capsule</td>
</tr>
<tr>
<td>eIF2α</td>
<td>Eukaryotic Initiation Factor 2α</td>
</tr>
<tr>
<td>Endo H</td>
<td>Endoglycosidase H</td>
</tr>
<tr>
<td>ER</td>
<td>Endoplasmic reticulum</td>
</tr>
<tr>
<td>ERES</td>
<td>ER exit sites</td>
</tr>
<tr>
<td>FIP200</td>
<td>RB1-inducible coiled-coil protein 1</td>
</tr>
<tr>
<td>GFAP</td>
<td>Glial fibrillary acidic protein</td>
</tr>
<tr>
<td>HC</td>
<td>Hippocampal commissure</td>
</tr>
<tr>
<td>H&amp;E</td>
<td>Hematoxylin and eosin</td>
</tr>
<tr>
<td>IBA1</td>
<td>Ionized calcium-binding adapter molecule 1</td>
</tr>
<tr>
<td>IC</td>
<td>Internal capsule</td>
</tr>
<tr>
<td>IFN</td>
<td>Interferon</td>
</tr>
<tr>
<td>ipb</td>
<td>Infrapyramidal bundle of mossy fibers</td>
</tr>
<tr>
<td>IPs</td>
<td>Immunoprecipitates</td>
</tr>
<tr>
<td>IRF3</td>
<td>Interferon regulatory factor 3</td>
</tr>
<tr>
<td>ISGs</td>
<td>Interferon stimulated genes</td>
</tr>
<tr>
<td>KI</td>
<td>Kinase inactive</td>
</tr>
<tr>
<td>L1</td>
<td>Neural adhesion molecule L1</td>
</tr>
<tr>
<td>MAPK</td>
<td>Mitogen-activated protein kinases</td>
</tr>
<tr>
<td>MEFs</td>
<td>Murine embryonic fibroblasts</td>
</tr>
<tr>
<td>Abbr</td>
<td>Description</td>
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<tr>
<td>-------</td>
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<tr>
<td>mTOR</td>
<td>Mechanistic target of rapamycin</td>
</tr>
<tr>
<td>NCAM</td>
<td>Neural cell adhesion molecule</td>
</tr>
<tr>
<td>NeuN</td>
<td>Neuronal nuclei</td>
</tr>
<tr>
<td>NF165</td>
<td>Neurofilament 165 kDa</td>
</tr>
<tr>
<td>NHS</td>
<td>N-hydroxysuccinimido</td>
</tr>
<tr>
<td>NSM</td>
<td>Neurosecretory motor</td>
</tr>
<tr>
<td>PAS</td>
<td>Preautophagosomal structure</td>
</tr>
<tr>
<td>P62/SQSTM1</td>
<td>Sequestosome 1</td>
</tr>
<tr>
<td>Pb</td>
<td>Probst bundle</td>
</tr>
<tr>
<td>PI3K</td>
<td>Phosphatidylinositol 3-kinase</td>
</tr>
<tr>
<td>PM</td>
<td>plasma membrane</td>
</tr>
<tr>
<td>PNGase</td>
<td>Peptide-N-glycosidase F</td>
</tr>
<tr>
<td>PARP1</td>
<td>Poly (ADP-ribose) polymerase 1</td>
</tr>
<tr>
<td>polyQ-Htt</td>
<td>Polyglutamine expanded huntingtin</td>
</tr>
<tr>
<td>PSPB</td>
<td>Pallial-subpallial boundary</td>
</tr>
<tr>
<td>Ser</td>
<td>Seronine</td>
</tr>
<tr>
<td>SERT</td>
<td>Serotonin transporter</td>
</tr>
<tr>
<td>SO</td>
<td>Stratum oriens;</td>
</tr>
<tr>
<td>Spb</td>
<td>Suprapyramidal bundle of mossy fibers</td>
</tr>
<tr>
<td>SR</td>
<td>Stratum radiatum</td>
</tr>
<tr>
<td>STING</td>
<td>Stimulator of interferon genes</td>
</tr>
<tr>
<td>TAG-1</td>
<td>Transient axonal glycoprotein-1</td>
</tr>
<tr>
<td>Tbr1</td>
<td>T-Box, Brain 1</td>
</tr>
<tr>
<td>Abbreviation</td>
<td>Description</td>
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<td>--------------</td>
<td>----------------------------------</td>
</tr>
<tr>
<td>TCA</td>
<td>Thalamocortical axon</td>
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<tr>
<td>ULK1</td>
<td>UNC-51–like kinase 1</td>
</tr>
<tr>
<td>ULK2</td>
<td>UNC-51–like kinase 2</td>
</tr>
<tr>
<td>UPR</td>
<td>Unfolded protein response</td>
</tr>
<tr>
<td>VPL</td>
<td>Ventral posterolateral nucleus</td>
</tr>
<tr>
<td>VPM</td>
<td>Ventral posteromedial nucleus</td>
</tr>
<tr>
<td>VSV-G</td>
<td>Vesicular stomatitis virus (VSV) G-protein</td>
</tr>
<tr>
<td>WGA</td>
<td>Wheat germ agglutinin</td>
</tr>
<tr>
<td>XBP1</td>
<td>X-Box Binding Protein 1</td>
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<tr>
<td>5-HT</td>
<td>Serotonin</td>
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CHAPTER 1. INTRODUCTION

Autophagy is a catabolic process that occurs under basal physiological and stressed conditions. Cargoes, including superfluous organelles or proteins, are enclosed in double membrane-bound autophagosomes and degraded in lysosomes (Levine and Kroemer, 2008; Mizushima et al., 2008). As an evolutionarily conserved mechanism, autophagy is required for cellular homeostasis from yeast to mammals (Levine and Kroemer, 2008; Mizushima et al., 2008). It is widely accepted that autophagy dysregulation contributes to a variety of human diseases, including neurodegeneration, cancer, and infectious diseases (Levine and Kroemer, 2008; Mizushima et al., 2008).

To date, more than 30 autophagy-related (ATG) genes have been identified in yeast—the majority of which have homologues or counterparts in mammals (Feng et al., 2014). ATG genes can be grouped into several functional units: the ULK/ATG1 complex, which activates the VPS34/PI3K complex and promotes AT9-mediated membrane recycling; the VPS34/PI3K complex, which is essential for membrane nucleation; the two ubiquitin-like conjugation systems (Atg12-Atg5 and Atg8/LC3); Atg2-Atg18 complex; and the transmembrane protein Atg9 (Feng et al., 2014).

Since ULK/Atg1 is the only kinase in the autophagy pathway, and kinases are well accepted druggable targets for therapeutic purposes, ULK/ATG1 has gained much attention in the past decade. While potent inhibitors of ULK kinases have been developed to target autophagy in the context of diseases (Egan et al., 2015; Petherick et al., 2015), how ULK/ATG1 drives the cellular responses to physiological vs. stressed conditions is not clear. A better understanding about the function of ULK/Atg1 will help to improve the specificity of the inhibitors and minimize potential side effects in therapeutic outcomes.

ULK/ATG1 Regulates the Cellular Response to Metabolic Stress

The Canonical Autophagy-Inducing ULK/ATG1 Kinase Complex

ULK/ATG1 is the only known kinase with a specific role in autophagy. In yeast, disruption of Atg1 function abrogates autophagy at an early stage, leading to the conclusion that Atg1 functions at the initial step of autophagy (Matsuura et al., 1997). Meanwhile, yeast also utilizes the cytoplasm-to-vacuole targeting (Cvt) pathway to transport two vacuolar enzymes, aminopeptidase I and a-mannosidase, to the vacuole—a process that is dependent on Atg1 (Harding et al., 1995). Atg1 is reported to interact with at least eight other autophagy genes, some of which are only required for autophagy regulation (Atg13, Atg17, Atg29, Atg31), while the others (Atg11, Atg20, Atg24 and Vac8) are essential specifically for the Cvt pathway (Mizushima, 2010). As Cvt proceeds constitutively under nutrient-rich conditions and the Atg1-mediated autophagy occurs after nutrient depletion, it is thought that Atg1 may function as a switch between
autophagy and the Cvt pathway through post-translational modifications or shifting its associations with other interacting partners (Mizushima, 2010). Atg1 does not interact with Atg13 or Atg17 under nutrient-rich conditions but can bind to Atg13 upon starvation, and the increased binding of Atg13 upregulates its kinase activity (Kamada et al., 2000). The mammalian homolog of Atg1, ULK1, was first shown to regulate autophagy in a kinase screening for novel regulators of autophagy in response to amino acid starvation. ULK2, a close homologue of ULK1, plays a partially redundant role in regulating autophagy (Cheong et al., 2011; Lee and Tournier, 2011). The mammalian ULK complex consists of ULK1 and ULK2, ATG13, FIP200 (the mammalian counterpart of yeast Atg17), and ATG101. ATG13 and FIP200 modulate proper localization, stability, and kinase activity of ULK1 and ULK2 (Ganley et al., 2009; Hosokawa et al., 2009; Jung et al., 2009). Nevertheless, unlike the yeast Atg1-Atg13-Atg17, the ULK complex exists constitutively, irrespective of nutritional conditions (Ganley et al., 2009; Hosokawa et al., 2009; Jung et al., 2009), suggesting differential regulation of autophagy in different organisms. In addition, there appears to be no equivalent of the yeast Cvt pathway in mammalian cells. It remains to be seen whether the mechanism by which Atg1 delivers vacuole enzymes is inherited by mammals during evolution.

**Signalings Lead to Activation of ULK/ATG1**

Much effort has been devoted to the search for novel ULK/ATG1 substrates and towards understanding how the ULK/ATG1 complex is regulated by upstream regulators in response to cellular nutrient status (Figure 1-1). Under fed conditions, yeast Atg13 is hyperphosphorylated by TOR (target of rapamycin) kinase but is rapidly dephosphorylated when TOR is inhibited by starvation (Kamada et al., 2000; Kamada et al., 2010). As a consequence, it promotes Atg1-Atg13-Atg17 complex formation and the kinase activity of Atg1 (Kamada et al., 2000; Kamada et al., 2010). It is not yet clear if TOR also phosphorylates Atg1 under basal condition (Papinski and Kraft, 2016). In a slightly different fashion, both mammalian ULK1 and ATG13 are phosphorylated by mTOR under normal conditions, and they are quickly de-phosphorylated upon starvation, thus activating the ULK complex (Alers et al., 2012; Ganley et al., 2009; Hosokawa et al., 2009; Jung et al., 2009; Kim et al., 2011). Meanwhile, two studies almost simultaneously demonstrated that following glucose or Earle's Balanced Salt Solution (EBSS) starvation, activated AMPK directly phosphorylates ULK1 at Ser467, Ser555, Thr574, and Ser637 (Egan et al., 2011; Kim et al., 2011). Although phosphorylation of ULK1 by AMPK has no impact on the kinase activity of ULK1, it is sufficient to modulate the function of ULK complex (Alers et al., 2012; Egan et al., 2011; Kim et al., 2011; Mack et al., 2012). Thus, mTOR and AMPK work mutually to sense the metabolic cues and relay the signals to ULK complex. Independent of AMPK, ULK1 can be activated by the GSK3-TIP60 signaling axis upon growth factor deprivation. It is shown that depletion of growth factors from the cells triggers deinhibition of GSK3, which then directly phosphorylates the acetyltransferase TIP60 at Ser86. The activated TIP60 subsequently acetylates ULK1 to induce autophagy (Lin et al., 2012). Although cAMP-dependent protein kinase A (PKA) has been reported to phosphorylate yeast Atg1
Figure 1-1. ULK/ATG1 drives cellular responses to metabolic stresses.

Multiple nutrient-sensing pathways such as AKT-GSK3 axis, mTOR-AMPK axis, and some other unknown molecules for extracellular or intracellular stimuli impinge on ULK/ATG1 to achieve different regulatory effects mainly through posttranslational modification. The inhibited AKT, due to growth factor depletion, leads to activation of GSK3 and subsequent activation of TIP60, which catalyzes acetylation of ULK1 to induce autophagy. ULK/ATG1 is normally inhibited by mTOR. The energy sensor AMPK can release the suppressive effects of mTOR on ULK complex to promote autophagy. Some unknown molecules may exist to detect intracellular protein aggregates and signal the proteomic stress to the ULK complex to enhance protein degradation. Once activated by upstream regulators, ULK/ATG1 phosphorylates different substrates to regulate autophagy. The established substrates of ULK/ATG1 include Beclin1, VPS34, and ATG14 of the VPS34 complex as well as Atg9, DENND3, FUNDC1, and P62. The phosphorylated substrates of ULK/ATG1 kinase have been implicated to function at different steps of autophagy, such as initiation, membrane recycling, and autophagosome trafficking, etc. See text for details.
(Budovskaya et al., 2005), the potential link between mammalian PKA and ULK1/2 has yet to be established.

**Autophagy-Related Substrates**

The autophagy-related substrates of ULK/ATG1 can be categorized into at least four different groups: (i) ULK complex, comprising of ULK1, ATG13, FIP200, and ATG101, (ii) VPS34 complex, including Beclin-1, VPS34, and ATG14, (iii) ATG9, and, (iv) other substrates that are selective for certain conditions. While mammalian ULK1 was found to autophosphorylate Thr180, located in the kinase activation loop essential for intact kinase activity of ULK1 (Bach et al., 2011), it has been reported to phosphorylate ATG13 at Ser318, a modification required for targeting ATG13 to depolarized mitochondria (Joo et al., 2011). There is some evidence that ULK1 phosphorylates human FIP200 at Ser934, Ser986, and Ser1323, though the functional significance of these phosphorylation events has not yet been determined (Egan et al., 2015). Ser11 and Ser203 of human ATG101 are reported to be phosphorylated in an ULK1-dependent manner (Egan et al., 2015). Similar to the case of FIP200, how the phosphorylation of ATG101 contributes to autophagy is not clear. It is noteworthy that not all the reported phosphorylation sites on the putative substrates of ULKs are conserved, indicating that ULK complex may be regulated in a species-specific manner. Independent laboratories have identified Beclin-1, VPS34, and ATG14 (components of the VPS34 complex) as bona fide substrates of ULK1 during amino acid (AA) starvation (Egan et al., 2015; Park et al., 2016; Russell et al., 2013). By phosphorylating human Beclin-1, VPS34, and ATG14 at Ser14, Ser249, and Ser29, respectively, ULK1 enhances the activity of VPS34 complex to promote autophagosome formation (Egan et al., 2015; Park et al., 2016; Russell et al., 2013). ATG1/ULK has been implicated in Atg9/ATG9 cycling in both yeast and mammalian cells. In yeast, Atg9 cycles between the preautophagosomal structure (PAS) and the peripheral sites, in an Atg1-dependent manner (Reggiori et al., 2004). Recent studies, however, have demonstrated that Atg1-mediated phosphorylation of Atg9 at multiple sites is required for recruitment of Atg8 and Atg18 to the PAS (Papinski et al., 2014). Nevertheless, phosphorylation of Atg9 by Atg1 is not necessary for its shuttling between PAS and cytosol, as phosphor-mutant Atg9 localizes to the PAS properly, in contrast to the abnormal accumulation of Atg9 in the PAS in yeast with Atg1 deletion, suggesting that Atg1 modulates Atg9 function by both kinase-dependent and -independent mechanisms (Papinski et al., 2014). Whether ULK1(2) governs ATG9 cycling in the mammals is not as clear. An earlier reports showed that knockdown of ULK1 abolished starvation-induced redistribution of ATG9 (Young et al., 2006), recent work suggests that independent of ULK1, ATG9 can be recruited to DFCP1-positive phagophores (Orsi et al., 2012). It is yet to be determined whether mammalian ATG9 is also a substrate of ULK1(2) and whether the mechanisms how Atg1 governs the Atg9 function are conserved.

In addition to targeting various components of the autophagy machinery, several proteins not typically regarded as autophagy molecules are also bona fide substrates of ULK/ATG1 under certain conditions. FUNDC1 is a mitochondrial outer-membrane
protein that acts as a cargo receptor protein in hypoxia-induced mitophagy. Phosphorylation of mouse FUNDC1 at Ser17 by ULK1 is required for efficient binding of LC3 to FUNDC1 and mitophagy progression (Wu et al., 2014). Following EBSS treatment, the guanine nucleotide exchange factor DENND3 is phosphorylated at Ser554 and Ser572 in an ULK1- and ULK2-dependent manner (Xu et al., 2015). The phosphorylation promotes activation of Rab12, a small GTPase that regulates autophagosome trafficking. The idea that ULK1 and ULK2 regulate autophagosome trafficking is intriguing. The increasing number of substrates of ULK/ATG1 that appear to work at different stages of the autophagy pathway strongly argues that ULK1 and ULK2 function at multiple steps during autophagy induction. However, to verify this concept can be technically difficult, unless distinct mutants of ULK/ATG1 can be created to precisely dissect the upstream and downstream functions of these enzymes. Alternatively, according to different metabolic cues, ULK/ATG1 can probably target different substrates to achieve differential responses. In fact, a recent study has demonstrated that in response to proteasome inhibition or expression of polyglutamine-expanded huntingtin (polyQ-Htt), ULK1 phosphorylates the autophagy receptor protein p62 at Ser409 to induce clearance of ubiquitinated protein. More importantly, the ULK1-mediated phosphorylation does not occur upon nutrient starvation (Lim et al., 2015). Future studies to systematically analyze phosphorylation of known ULK substrates under various stress conditions will provide significant insights into the regulatory mechanism underlying ULK-mediated autophagy.

The Historical View of ULK/ATG1 Kinase

Although the yeast Atg1 was the first known autophagy-defective strain initially identified in a global screen for autophagy loss-of-function strains (Tsukada and Ohsumi, 1993), the isolation and validation of the metazoan homologs was a twisted story.

_C. elegans_ UNC-51 kinase was originally cloned as a protein critical for axonal development because disrupting _unc-51_ expression results in uncoordinated movement of _C. elegans_ due to widespread axonal guidance and fasciculation abnormalities in the motor neurons (Ogura et al., 1994). Aberrant vesicles and membranous structures are oftentimes observed in these neurons (McIntire et al., 1992). Similar to _unc-51_ mutants, _D. melanogaster atg1_ mutants also exhibit abnormal axonal tracts in the ventral nerve cord, including premature truncation and defective midline crossing of longitudinal tracts at the embryonic stage (Toda et al., 2008). Later analysis also revealed abnormal defasciculation of the larval mushroom body axons in _D. melanogaster atg1_ mutants (Mochizuki et al., 2011). Mammalian _Ulk1_ and _Ulk2_ were first cloned based on their high degree of homology to the _C. elegans unc-51_ gene (Yan et al., 1998; Yan et al., 1999). And the neuronal function of ULK1 was subsequently demonstrated by independent studies. Dominant-negative expression of _Ulk1_ in mouse cultured cerebellar granule neurons inhibits neurite formation and extension (Tomoda et al., 1999). Meanwhile, knocking down the expression of _Ulk1, Ulk2_, or both, leads to extensive neurite branching and stalled axonal growth in mouse dorsal root ganglion neurons in culture (Zhou et al., 2007).
While ULK/ATG1 kinase plays evolutionary conserved role in axonal development, the C. elegans UNC-51 turned out to be a close homolog of yeast Atg1, sharing 52.7% similarity in their N-terminal kinase domain (Matsuura et al., 1997). The role of yeast Atg1 in autophagy regulation led to the finding that UNC-51 is also essential for proper localization of autophagosomal marker proteins and starvation-induced dauer development, an autophagy-dependent process (Melendez et al., 2003). In addition, as observed in the S. cerevisiae and C. elegans, the ATG1 has been shown to act in autophagy initiation, downstream of Drosophila TOR (dTOR) (Scott et al., 2007). Of note, overexpression of ATG1 is sufficient to induce autophagy (Scott et al., 2007). However, the function of ULK1 in autophagy regulation wasn’t discovered until about a decade ago when Tooze lab performed a siRNA screening for essential genes in amino acid starvation-induced autophagy (Chan et al., 2007). Since then, ULK1 and, to a lesser extent, ULK2 have been substantially studied in the context of autophagy regulation under various conditions (Figure 1-2).

**Closing Introductory Remarks**

Because of their diverse functions, ULK1 and ULK2 can be potentially linked to various human diseases including neurodegeneration. However, vast majority of studies of ULK1 and ULK2 are performed *in vitro*, and the physiological relevance of ULK1 and ULK2 is poorly characterized. On the one hand, one of the primary roles of autophagy is to maintain cellular homeostasis through degrading damaged organelles and protein aggregates. Neurons, as one of the few long-lived cell types, heavily rely on autophagy for survival (Nixon, 2013). Thus, it is not surprising that deleting key autophagy genes such as Atg5, Atg7, and Fip200, in the central nervous system (CNS) causes widespread cell death associated with accumulation of p62 and ubiquitin-positive inclusion bodies, a common pathological feature in a variety of human neurological diseases (Hara et al., 2006; Komatsu et al., 2006; Liang et al., 2010). Whether ULK1 and ULK2 are important for neuronal autophagy under physiological conditions is an important question that remains to be answered. On the other hand, the precise *in vivo* role of ULKs (in axonal development) has not yet been demonstrated unequivocally. Additionally, it is unclear whether autophagy is required for axonal elongation during mammalian neurodevelopment. Since no neuronal functions have been documented for other members of the ULK complex, including ATG13 and FIP200 (Kaizuka and Mizushima, 2015; Liang et al., 2010), it is likely that ULK1 and ULK2 play specialized roles in neurodevelopment, and this might involve a group of separate, novel binding partners. Although *Ulk1* single knockout (KO) mice exhibited defects in reticulocyte maturation, no other overt developmental defects were observed (Kundu et al., 2008). No major phenotype has been reported for *Ulk2* single KO mice either (Cheong et al., 2011). This suggests at least a partial functional redundancy is shared between ULK1 and ULK2. In line with that, *Ulk1/2* double knockout (dko) mice die shortly after birth, partially due to defective lung development (Cheong et al., 2014). To answer the core question of what is the physiological role(s) of ULK1 and ULK2 in the mouse brain, we generated and characterized the phenotype of CNS-specific *Ulk1* and *Ulk2* conditional double knockout
Figure 1-2. The historical view of ULK/ATG1 kinase.
(Ulk1/2-cdko) mice. The results obtained from the detailed analyses will be described in Chapter 3 and 4.
CHAPTER 2. EXPERIMENTAL PROCEDURES*

Mice

The strategy for targeting the Ulk2 locus was described previously (Cheong et al., 2011). Mice harboring the targeted Ulk2 allele (Ulk2fllox-neo) were crossed with Ella-Cre transgenic mice (Jackson Laboratory, stock: 003724), and progeny with Ulk2ko or Ulk2fllox alleles were crossed with WT mice to eliminate the expression of the Cre-recombinase. Ulk1−/− mice have been described previously (Kundu et al., 2008). The Nestin-Cre line was obtained from the Jackson Laboratory (stock: 003771). Ulk1+/−;Ulk2fllox/fllox females were bred with Ulk1+/−;Ulk2−/−;Nestin-Cre males to generate Ulk1/2-cdko (Ulk1−/−;Ulk2−/−;fllox;Nestin-Cre) mice. Atg7f/f (a generous gift from Dr. Masaaki Komatsu, Tokyo Metropolitan Institute of Medical Science), and Fip200f/f females were crossed with Nestin-Cre males to generate Atg7Nestin cko and Fip200Nestin cko mice. All animals were maintained in a mixed background between C57BL/6 and 129. All animal experiments were approved by and performed in accordance with guidelines provided by the Institutional Animal Care and Use Committee at St. Jude Children’s Research Hospital.

Immunostainings and Histologic Analyses

Embryonic and newborn mouse brains were dissected in phosphate-buffered saline (PBS) and fixed overnight in 4% paraformaldehyde (PFA) in PBS at 4°C. Mice older than postnatal day (P) 1 were transcardially perfused with 4% PFA and fixed overnight. The brains were then cryoprotected using 30% sucrose in phosphate-buffered saline (PBS) overnight at 4°C and embedded in OCT for cryosectioning. Frozen sections were washed with 0.2% Triton X-100 in tris-buffered saline (TBST) and incubated in the blocking solution (5% normal goat serum or normal donkey serum in TBST) for 1h at room temperature. Sections were incubated with primary antibodies diluted in the blocking solution overnight at 4°C, washed with TBST, and incubated with Alexa Fluor–conjugated secondary antibodies (Invitrogen), diluted at 1:1000 in the blocking solution for 2 h at room temperature. Sections were mounted in ProLong Gold antifade reagent with DAPI (Invitrogen). Primary antibodies against the following targets were used: NeuN (Synaptic System, 1:500), NeuN (Millipore, 1:1000), Cleaved caspase-3 (Cell Signaling, 1:200), GFAP (Dako, 1:500), Iba1 (Wako, 1:500), P62 (Abnova, 1:500), Ubiquitin (Dako, 1:500), p-S6 (Cell Signaling, 1:400), p-elF2α (Cell Signaling, 1:100), Chop (Santa Cruz, 1:100), BiP (Abcam, 1:500), and ATF6 (Abcam, 1:500), Calbindin

(Millipore, 1:500), L1 (Millipore, 1:500), NF165 (Developmental Studies Hybridoma Bank, 1:100), TAG-1 (Developmental Studies Hybridoma Bank, 1:5), NCAM (Developmental Studies Hybridoma Bank, 1:100), Satb2 (Abcam, 1:500), Ctip2 (Abcam, 1:200), anti-Tbr1 (Abcam, 1:500), rabbit anti-GFAP (DAKO, 1:500), rabbit anti-Calretinin (Millipore, 1:500), rabbit anti–5-SERT (Millipore, 1:1000), guinea pig anti-VGlut2 (Millipore, 1:2000).

For histologic analysis, formalin-fixed brains were processed and embedded in paraffin by using standard methods. Sagittal sections (4µm) were cut and stained with hematoxylin and eosin (H&E), Luxol Fast Blue, and counterstained with Nissl by staff in the St. Jude Veterinary Pathology Core, and the sections were examined by a pathologist blinded to the experimental groups. Fluoro-Jade C (Millipore, Cat# AG325) staining was carried out on 20-µm frozen sections per the manufacturer’s instructions. Quantification of CA1 pyramidal neurons was done with ImageJ software (National Institute of Health, USA). The counts of DAPI in a 500µm region of CA1 were determined manually. For each animal, the average number of neurons was calculated from at least 5 sections. Cell numbers in the mutants were normalized to their corresponding controls.

To quantify the dorsoventral corpus callosum (CC) width, low-magnification images were taken with a Marianas 2 microscope. Using SlideBook 6.0 software, the CC width was measured in 5 rostrocaudal sections per animal, and the values were presented as the mean ± s.e.m. To quantify the number of axonal bundles crossing the pallium-subpallium boundary (PSPB) at E14.5, a line spanning the PSPB was drawn, and the number of axonal bundles crossing the line was quantified from 3 comparable sections from each Ulk1/2-deficient mouse and the control. Imaris was used to count the numbers of different cell types in the brains of control and Ulk1/2-deficient mice. At least three comparable sections from individual control and Ulk1/2-deficient mice were quantified, and values were shown as the mean ± s.e.m.

**Behavioral Analyses**

For the limb-clasping test, mice were suspended by pulling their tails. For the rotarod assay, mice were put on a rotating rod, the speed of which gradually increased from 0 to 40 rpm at a rate of 10 rpm/min. The time recorded was when the mouse fell from the rod, and 5 min was used as the cutoff for this analysis. For the gait analysis, the paws of mice were dipped in nontoxic colored ink: fore paws in blue ink and hind paws in red ink. The mice were then allowed to walk through a tunnel placed on top of white paper. The paw prints were then air dried and scanned.

**Morphometric Analyses of Electron Micrographs**

Mice were deeply anesthetized with CO₂, perfused transcardially with 10 mL phosphate buffer followed by 10 mL 2.5% glutaraldehyde/2% PFA in 0.1 M CaCO₃. The brains were removed and fixed in the same fixative and postfixed in 2% osmium.
tetroxide in 0.1 M sodium cacodylate buffer with 0.3% potassium ferrocyanide for 2 h. Vibratome sections (100-mm) of the brain were cut sagittally and collected in cold PBS, rinsed in phosphate buffer, dehydrated through a series of graded ethanol-to-propylene oxide solutions, infiltrated and embedded in epoxy resin, and polymerized at 70°C overnight. Semithin (0.5-µm) sections were stained with toluidine blue for light microscope examination. Ultrathin (80-nm) sections were cut and imaged using an FEI Tecnai F 20 TEM FEG electron microscope with an AT XR41 camera.

For quantitative analysis of endoplasmic reticulum (ER), all images of neurons in the hippocampal CA1 regions of 2 controls and 2 mutants (8-wk-old) were taken at the same parameters. For each sample, images were randomly obtained from at least 20 neurons. For the quantification of ER number and diameter, the TIFF images were opened with NIS-Elements software (Nikon) and calibrated individually. A 2-µm line was drawn to span as many ER tubules as possible. The number of ER tubules the line crossed was recorded as the ER number. The diameters of the ER tubules covered by the line were also measured individually. To quantify the percentage of ER occupation in the cytosol, the same sets of images were opened in the ImageJ software and calibrated accordingly. Then a 25-µm² area was drawn in the cytosol where the ER tubules were the most abundant. The ER tubules in each image were selected and masked manually. The total area of ER in the defined cytosolic region was then measured and recorded. Ten cells from each sample were analyzed.

Immunoprecipitation

Endogenous ULK1 was extracted from the brain of 4-wk-old WT mice or from MEFs by using a Triton-based cell lysis buffer (40 mM HEPES, 120 mM NaCl, 1 mM EDTA, 1.5 mM Na3VO4, 50 mM NaF, 10 mM β-glycerophosphate, 20 mM MoO4, 0.5% Triton X-100, protease inhibitor, phosphatase inhibitor). The lysates were incubated with anti-ULK1 antibody (Santa Cruz Biotechnology, sc10900) overnight at 4°C and precipitated with Protein G agarose beads (Thermo Scientific). For immunoprecipitation of GFP–SEC16A, whole-cell extracts were prepared from 293T cells by using the described Triton-based buffer and precipitated with anti-GFP antibody–conjugated sepharose beads (Abcam, ab69314) after overnight incubation at 4°C. The beads were washed 5 times with cold Triton-based buffer and incubated at 95°C for 5 min in SDS sample buffer (Sigma Aldrich). Phosphatase treatment was performed on GFP IPs by using calf alkaline intestinal phosphatase (Sigma Aldrich) per the manufacturer’s protocol, before elution in SDS sample buffer. Anti–FLAG M2-agarose beads (Sigma Aldrich, A2220) were used for immunoprecipitation of FLAG-tagged proteins.

COPII Vesicle–Formation Assay

The preparation of cytosol (source of soluble COPII proteins) and medium-speed membrane-pellet (source of membrane and cargo) fractions has been described before (Ge et al., 2013; Ge et al., 2014). Each vesicle-budding reaction consists of 3 µL

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membrane from the medium-speed pellet (OD600 = 10 for total membrane), 25 µL cytosol (5 mg/mL), 0.75 µL GTP, and 5 µL 10× ATP regeneration (Ge et al., 2013; Ge et al., 2014). B88 buffer (20 mM HEPES-KOH, pH 7.2, 250 mM sorbitol, 150 mM potassium acetate, and 5 mM magnesium acetate) was added last to adjust the reaction mixture to a final volume of 50 µL. The reaction was performed at 30°C for 1 h followed by centrifugation at 20,000 xg for 20 min. Supernatant aliquots (35 µL) were transferred to an ultracentrifuge tube for sedimentation at 100,000 xg in a Beckman TLA100.3 rotor for 30 min. The supernatant fractions were removed, and the small membranes were analyzed by immunoblotting with the antibodies indicated in the figures.

Plasmid Constructs

The pmGFP-SEC16A (Addgene, Cat# 15775) and pmGFP-SEC16B vectors (Addgene, Cat# 15776) have been previously described (Bhattacharyya and Glick, 2007). SERT–GFP (Origene Technologies, Rockville, MD, Cat# RG210187) and VSVG (ts045)–GFP vectors (Addgene, Cat# 11912) were used in the ER-to-Golgi trafficking assay (Presley et al., 1997). The pCMV6-MYC-DDK–human SEC16A plasmid (Cat# RC223625), pCMV6-MYC-DDK–human ULK1 (Cat# RC215643), and pCMV6-MYC-DDK-human ULK2 (Cat# RC206010) were purchased from OriGene Technologies. FLAG-tagged ULK1 and deletion constructs in the pME18S vector (a generous gift from Dr. Toshifumi Tomoda, Beckman Research Institute of City of Hope, Duarte, CA) were previously described (Joo et al., 2011; Yan et al., 1998). The FLAG-tagged ULK2 construct in the PCSII vector was a generous gift from Dr. Do-Hyung Kim (University of Minnesota, Twin Cities, MN). Plasmid vectors, pCS2+, containing SEC24A, SEC24B, SEC24C, and SEC24D open-reading frames, with a 3× HA tag at the N terminus, were described previously (Kim et al., 2007). The Ulk1 and Ulk1 kinase–inactive (KI) (K46A) mutant, as described previously (Joo et al., 2011), were subcloned into the 5′– or 3′–end of EcoR1–restriction enzyme sites of the MSCV-IGFP-MII (generously provided by Dr. Terrence Geiger, St. Jude) and pcDNA3.1 vectors for retrovirus generation and transient transfection, respectively. Human SEC16A phosphorylation-site mutants were introduced into the pmGFP–SEC16A and pCMV6–entry-(MYC-DDK–tagged) human SEC16A vectors by using mutagenic oligonucleotides.

Generation of Murine Embryonic Fibroblasts, Cell Culture, Transfection, Gene Silencing, and Drug Treatment

Ulk1-ko, Ulk2-ko, and Ulk1/2-dko murine embryonic fibroblasts (MEFs) were prepared from the E12.5 embryos of Ulk1+/−; Ulk2+/− × Ulk1+/−; Ulk2+/− crosses. Embryos were collected, washed twice with PBS, incubated in trypsin, and then separated into single-cell suspensions. Cells were expanded in DMEM (Invitrogen) supplemented with 10% fetal bovine serum (FBS) (Hyclone), MEM nonessential amino acids (NEAA; Life Technologies), L-glutamine (Life Technologies), β-mercaptoethanol (Life Technologies), and gentamycin (Life Technologies). These primary MEFs were then immortalized using
FuGENETM 6 (Roche) to transfect the cells with 1 µg of an expression vector expressing SV40, as described previously (Joo et al., 2011).

After transfection, cells were maintained in normal growth media. Atg7−/− MEFs and stably transduced MEFs were generated as described previously (Joo et al., 2011). Briefly, MSCV-IGFP-MII containing no cDNA (empty vector), WT ULK1, or mutant ULK1, and helper vector (generously provided by Dr. Terrence Geiger) were cotransfected into 293T cells by using FuGENE 6. Transfected cells were then incubated in DMEM with 10% FBS for 48 h. Supernatant was collected twice daily and used to infect GP+E86 retroviral–producer cells in the presence of Polybrene (8 µg/mL). Transduced GP+E86 cells were FACS-sorted by the presence of GFP, and cell-free supernatant was used to transduce Ulk1−/− MEFs. MEFs and 293T cells were grown in DMEM supplemented with 10% FBS, penicillin/streptomycin (Invitrogen), and Glutamax (Invitrogen) at 37°C (5% CO2). The cells were transfected with FuGENE 6 according to the manufacturer’s protocol for transient overexpression of cDNA constructs.

Knockdown experiments in MEFs were performed using Lipofectamine RNAi Max (Life Technologies) per the manufacturer’s protocol and with the following siRNA constructs obtained from Dharmacon: pooled nontargeting siRNA (D-001810-10-05), SMARTpool: ON-TARGETplus Ulk1 siRNA (L-040155-00-0005); SMARTpool: ON-TARGETplus Sec16a siRNA (L058170-02-0005); ON-TARGETplus Sec16a siRNA Targeted Region: 3’UTR (J-058170-11-0005); SMARTpool: ON-TARGETplus Sec24a siRNA (L-056263-01-0005); SMARTpool: ON-TARGETplus Sec24b siRNA (L-048898-01-0005); SMARTpool: ON-TARGETplus Sec24c siRNA (L-059052-01-0005); SMARTpool: ON-TARGETplus Sec24d siRNA (L-065430-01-0005); SMARTpool: ON-TARGETplus Atg7 siRNA (L-049953-00-0005); SMARTpool: ON-TARGETplus Atg13 siRNA (L-053540-01-0005); and SMARTpool: ON-TARGETplus Atg14 siRNA (L-172696-00-0005). Knockdown was confirmed by RT-qPCR (TaqMan) analyses, immunoblot analyses, or both. To chemically induce ER stress, 293T cells were treated with Brefeldin A (Sigma Aldrich) and dissolved in DMSO (final concentration, 10 µg/mL) for 6 h. Cells were then lysed and analyzed by immunoblotting.

**Immunoblot Analyses and Antibodies**

Whole hippocampi or MEF cells were lysed in Triton-based cell lysis buffer. Proteins in cleared lysates were electrophoretically separated on 4% to 12% bis-Tris gels (Life Technologies). Proteins were then transferred to either a nitrocellulose or a PVDF membrane.

After incubation with a 5% skim milk block, blots were probed with antibodies directed against the following targets: P62 (Sigma Aldrich, Cat# P0067), LC3 (MBL, Cat# PM036) ULK1 (Sigma Aldrich, Cat# A7481), p-ULK1 (S555) (Cell Signaling, Cat# 5869), p-(Ser/Thr) Phe (Cell Signaling, Cat# 9631), SEC16A (Novus, Cat# NB100-1799), SEC16A pS846 (custom polyclonal antibody from Rockland), ATG7 (Cell
Signaling, Cat# 2631), ATG13 (Sigma Aldrich, Cat# SAB4200100), p-ATG13 (S318) (Rockland, Cat# 600-401-C49), ATG14 (MBL, Cat# PD026), CDC37 (Santa Cruz Biotechnology, Cat# sc-13129), SERT (Millipore, Cat# PC177L), ATF3 (NOVUS, Cat# NBP1-02935), p-elf2α (Cell Signaling, Cat# 3597), CHOP and BIP (a generous gift from Dr. Linda Hendershot, St. Jude), GFP (Abcam, Cat# ab6556), FLAG (Sigma Aldrich, Cat# F1804), HA (Cell Signaling, Cat# 3724), Myc (Cell Signaling, Cat# 2276), DDK (OriGene, Cat# TA50011-100), ACTB (Santa Cruz Biotechnology, Cat# I-19), and GAPDH (Sigma Aldrich, Cat# G9545). Membranes were then incubated with horseradish peroxidase–conjugated secondary antibodies, and bands were detected using chemiluminescence-detection kits (Amersham).

**Proteomics**

The mass spectrometric (MS) analysis was performed per an optimized platform, as previously reported (Xu et al., 2009). ULK1-interacting proteins were visualized by performing silver staining (Life Technologies) or SYPRO Ruby protein gel staining (Sigma Aldrich) according to the manufacturer’s protocol. Proteins in gel bands were reduced by adding dithiothreitol (DTT) and then alkylated by adding iodoacetamide. The gel bands were washed, dried in a speed vacuum, and rehydrated with a trypsin-containing buffer for overnight proteolysis. The digested peptides were extracted, dried, reconstituted, and loaded onto a capillary reverse-phase C18 column by an HPLC system (Waters ACQUITY UPLC). Peptides were eluted in a gradient, ionized by electrospray ionization, and detected by an in-line mass spectrometer (Thermo scientific LTQ Orbitrap Elite). MS spectra were collected, and the top 20 abundant ions were sequentially isolated for MS/MS analysis. This process was cycled over the entire liquid chromatography gradient. The acquired MS/MS spectra were used to search protein databases to obtain possible peptide matches. All matched MS/MS spectra were filtered by mass accuracy and matching scores to reduce the protein false-discovery rate to less than 1%.

**Quantitative Real-Time PCR**

Total RNA of adult hippocampi was extracted with RNeasy Mini Kit (Qiagen) according to the manufacturer’s instructions. Total RNA was isolated from cells by using TRIzol Reagent (Life Technologies). The reverse-transcription reaction was carried out using the SuperScript III first-strand synthesis kit (Life Technologies) per the manufacturer’s instructions. TaqMan gene-expression assays containing FAM-labeled primer/probe sets specific for Ulk1 (Mm-00437238_m1), Ulk2 (Mm-00497023_m1), Sec16a (Mm00462283_m1), Sec24a (Mm00613960_m1), Sec24b (Mm01313235_m1), Sec24c (Mm00655499_m1), Sec24d (Mm00660744_m1), and 18S were obtained from Applied Biosystems. The real-time PCR reactions were performed in a total reaction volume of 25 µL by using FastStart TaqMan Probe Master (Roche) reagent, and results were analyzed using the ABI 7900 real-time PCR-detection system (Applied Biosystems). Relative expression was normalized to 18S RNA and calibrated to the respective controls.
Immunofluorescence Microscopy

To assess SEC24C puncta quantity and ULK1 distribution, MEFs were incubated in complete growth media, fixed in 4% PFA, permeabilized with digitonin (100 µg/mL in PBS), blocked in 1% BSA, and then labeled with one of the following antibodies: anti-ULK1, anti-SEC24C (Abcam, ab122635), anti-HA or anti-FLAG. Cells were then treated with secondary antibodies conjugated to Alexa-555 or Alexa-647 (Invitrogen). To assess endogenous SEC16A distribution, MEFs were fixed in cold methanol for 15 min; slowly brought to room temperature in PBS; and blocked in PBS, 4% BSA, and 0.1% Triton X-100. MEFs were then labeled with anti-SEC16A antibody (Bethyl Laboratories, Cat# KIAA0310), followed by Cy3-conjugated secondary antibody (Jackson Immunoresearch, Cat# 711-165-152). To evaluate VSVG–GFP and SERT–GFP trafficking, MEFs were fixed and prestained with wheat germ agglutinin and Alexa Fluor 647 conjugate (Life Technologies, Cat# W32466) and then permeabilized with Triton X-100 (100 µg/mL in PBS) if additional staining with anti-FLAG antibody was needed.

To quantify SEC24C and GFP-SEC16A puncta, MEFs were imaged using a spinning-disc confocal Zeiss AxioObserver operated by Marianas system (Intelligent Imaging Innovations, Denver CO) with a 63× oil objective, and the number of puncta was analyzed using SlideBook 5.5 software (Intelligent Imaging Innovations, Denver CO). To quantify endogenous SEC16A puncta, MEFs were imaged using a swept-field confocal microscope (Nikon Ti-E) equipped with a Roper CoolSNAP HQ2 CCD camera using a Nikon 60×, 1.4 numerical aperture plan apochromat oil objective lens. Acquisition parameters were controlled by NIS-Elements software, and image analysis was conducted using Metamorph software. Confocal microscopy of ULK1 and GFP–SEC16A colocalization was performed on a Nikon TE2000E2 microscope equipped with a Nikon C2 confocal microscope using 488-nm, 561-nm, and 638-nm DPSS lasers for excitation. Images were acquired using a Nikon 60× 1.45 NA objective and analyzed using the NIS-Elements software. The following filters were used: 515/60, 605/75, and 675/50. VSVG-GFP– and SERT-GFP–transfected MEFs were imaged using a spinning-disc confocal Zeiss AxioObserver operated by Marianas system (Intelligent Imaging Innovations, Denver CO) with a 63× oil objective.

To quantify nuclear localization of CHOP, MEFs were grown in normal growth medium, transiently transfected with VSVG (ts045)–GFP or SERT–GFP, and immunostained with anti-CHOP antibody (Santa Cruz, Cat# sc-575). At least 30 images were acquired, as described above for each condition. The percentage of cells with nuclear CHOP was obtained from 3 independent experiments. As a positive control, the MEFs were treated with Brefeldin A (10 µg/mL) for 6 h, after which 100% nuclear localization of CHOP was observed.

Immunofluorescence Microscopy in *C. elegans*

*C. elegans* gonads were dissected on poly-L-lysine-coated slides, frozen in liquid nitrogen, and methanol-fixed at −20°C (Audhya et al., 2005). Tissues were
immunostained using a polyclonal rabbit antibody directed against C. elegans SEC16 (Witte et al., 2011) and subsequently labeled with a Cy3-conjugated anti-rabbit secondary antibody. Proximal oocytes were examined on a Nikon Ti-E swept-field confocal microscope, with a Roper CoolSNAP HQ2 CCD camera and using a Nikon 60× 1.4 numerical aperture plan apochromat oil objective lens. Acquisition was controlled using NIS-Elements software, and image analysis was performed using Imaris Bitplane software. Thirty z-sections at 0.2-µm steps were acquired. The number and fluorescence intensities of SEC16 puncta were quantified using the Spots module of Imaris Bitplane in at least 10 animals for each condition. Puncta fluorescence was classified as low-, medium-, or high-intensity thresholds, and the percentage of puncta in each category was compared between N2 control and unc-51–mutant animals. Immunostaining of rabbit anti–MOD-5 antibody (Jafari et al., 2011) and rabbit anti–5-HT antibody (purchased from Dr. H.W.M. Steinbusch, Maastricht University, Masstricht, The Netherlands) was performed with whole-mount WT worms and mod-5(n314) and unc-51(e369) mutants, as described previously (Jafari et al., 2011; Sze et al., 2000).

**Endoglycosidase H Assay**

For monitoring ER-to-Golgi trafficking of SERT, extracts prepared from SERT-GFP–transfected MEFs were used for GFP immunoprecipitation. Immunopurified proteins were incubated in the presence or absence of Endo H enzyme or peptide-N-glycosidase F (PNGase) enzyme (New England Biolabs, Cat# P0704) for 1 h, eluted from the beads in SDS sample buffer with β-mercaptoethanol, and separated by SDS-PAGE. SERT was detected by immunoblot analysis. Glycosylated proteins such as SERT and VSVG are Endo H–sensitive (Endo H–S) while in the ER, and become Endo H–resistant (Endo H–R) after ER-to-Golgi trafficking. PNGase treatment results in deglycosylated proteins.

**Cell-Surface Biotinylation**

The levels of SERT expression in MEFs or platelet PMs were compared using the membrane-impermeant biotinylation reagent, NHS-SS-biotin (Pierce, Inc., Rockford, IL), as described previously (Brenner et al., 2007; Ziu et al., 2012). Briefly, platelet or MEF pellets were washed twice with ice-cold PBSCM solution (PBS containing 0.1 mmol/L CaCl₂ and 1 mmol/L MgCl₂), incubated with NHS-SS-biotin (1.5 mg/mL) on ice with very gentle shaking for 20 min, rinsed briefly, incubated with PBSCM containing 100 mmol/L glycine on ice for 20 min, and lysed with 1% SDS–1% Triton X-100. Biotinylated PM proteins were recovered from the cell lysates by using streptavidin-agarose beads (Pierce, Inc.). After washing the beads with high-salt, low-salt, and 50 mmol/L Tris-HCl (pH 7.5), the biotinylated proteins were treated with either Endo H enzyme or PNGase F enzyme (New England Biolabs, Cat# P0704) for 1 h, eluted from the beads in SDS sample buffer with β-mercaptoethanol, and separated by SDS-PAGE. SERT was detected by immunoblot analysis.
Axonal Tracing

Embryonic day (E) 18.5 brains were fixed in 4% PFA overnight at 4°C. Small crystals of 1,1’-dioctadecyl-3,3,38,38, tetramethylindocarbocyanine perchlorate (DiI) (Molecular Probes) were inserted into the parietal cortex, somatosensory cortex, or dorsal thalamus (dTh) to trace different axonal tracks. The tissues were then stored in 4% PFA at 37°C for 3 to 4 weeks and cut on a vibratome into 100-µm sections. Sections were counterstained with Hoechst 33258 (Molecular Probes) and mounted with 50% glycerol.

Cytochrome Oxidase Staining

Brains were removed and fixed overnight in 4% PFA at 4°C. Barrel cortices were flattened on superfrost plus slides and cut tangentially (parallel to Layer IV) into 100-µm sections on a vibratome. Free-floating sections were incubated with cytochrome oxidase (CO) reaction solution, which included 0.5 mg/mL DAB, 0.5 mg/mL cytochrome C (Sigma), and 40 mg/mL sucrose in PBS at 37°C until staining appeared. Sections were washed 3 times in PBS, mounted with Aqua Polymount (Polysciences), and imaged using bright field light microscopy.

Statistical Analyses

Statistical analyses were performed using SigmaPlot; significance was assessed by 2-tailed Student’s t-test or by 1- or 2-factor ANOVA analysis followed by Holm-Sidak post-hoc analysis.
CHAPTER 3. THE NONCANONICAL ROLE OF ULK/ATG1 IN ER-TO-GOLGI TRAFFICKING IS ESSENTIAL FOR CELLULAR HOMEOSTASIS*

The third chapter of my dissertation is the publication from analyzing the phenotype of postnatal brain of Ulk1/2-cdko mice and the possible underlying mechanisms (Joo et al., 2016). I shared first authorship with Dr. Joung Hyuck Joo, a staff scientist in the lab who completed majority of the in vitro experiments. I performed all the animal experiments, and partially contributed to the biochemical studies. The following texts are quoted from the manuscript.

Mice Lacking Ulk1/Ulk2 Expression in the CNS Have a Distinct Pattern of Neuronal Loss

“Ulk1 and Ulk2 mRNA are expressed throughout the CNS, with the highest levels in the cerebellum (Ulk1>Ulk2) and hippocampus (Ulk2>Ulk1) (Figure 3-1A). To investigate the role(s) of these kinases in the CNS, we generated mice lacking both Ulk1 and Ulk2 in the CNS [Ulk1–/–;Ulk2–/fl;Nestin-Cre; hereafter referred to as Ulk1/2-cdko (conditional–double-knockout) mice] and those expressing Ulk1 but not Ulk2 in the CNS (Ulk1+/+;Ulk2–/fl;Nestin-Cre and Ulk1+/–;Ulk2–/fl;Nestin-Cre; hereafter referred to as “controls”) (Figure 3-1B, C). We confirmed the loss of Ulk1 and Ulk2 mRNA expression in the brains (i.e., hippocampal region) of Ulk1/2-cdko mice by RT-qPCR analyses (Figure 3-1D). Although the Ulk1/2-cdko mice were born at the expected Mendelian ratio, approximately 40% died within 24 h (Figure 3-2A). Survival of the remaining Ulk1/2-cdko mice diminished over the ensuing 12 wk, and all were dead by 28 wk (Figure 3-2A). Newborn Ulk1/2-cdko mice were visibly indistinguishable from their littermates, but at 3 wk showed substantial, sex-independent weight loss (Figure 3-2B). The weight of the brains of 8-wk-old Ulk1/2-cdko mice did not differ from that of littermate controls (Figure 3-2C).

Ulk1/2-cdko mice showed abnormal limb-clasping reflexes (Figure 3-1E) similar to that reported in Atg5–conditional knockout (cko) and Atg7-cko animals (Hara et al., 2006; Komatsu et al., 2006). However, unlike ATG5- or ATG7-deficient mice, which develop cerebellar ataxia (Hara et al., 2006; Komatsu et al., 2006), Ulk1/2-cdko mice had a normal gait (Figure 3-1F) and only a mild delay in motor skill learning compared to littermate controls (Figure 3-1G).

To identify the neurologic deficits in Ulk1/2-cdko animals, we analyzed their

Figure 3-1. Generation and behavioral characterization of *Ulk1*/*2* cdko mice.

(A) In situ hybridization (expression density map) from the Allen Brain Atlas showing the expression of *Ulk1* and *Ulk2* mRNA in the brains of 8-wk-old mice, with magnified views of the hippocampus and cerebellum. (B) Genomic organization of wild-type and targeted *Ulk2* loci. The genomic organizations of the wild-type (*Ulk2*^wt^) and targeted (*Ulk2*^floxed-neo^) *Ulk2* loci are shown (top 2 diagrams). Mice harboring the targeted *Ulk2* allele (*Ulk2*^floxed-neo^) were crossed with *EIIa-Cre* transgenic (tg) mice, and progeny harboring *Ulk2*^ko^ or *Ulk2*^floxed^ alleles (bottom 2 diagrams) were back-crossed with WT mice to eliminate Cre-recombinase expression. (C) Diagram of the generation of *Ulk1*/*2*-conditional double-knockout (*Ulk1*/*2*-cdko) mice. *Ulk1*^+/−;*Ulk2*^floxed/floxed^ mice were bred with *Ulk1*^+/−;Ulk2*^+/−* Nestin-Cre (tg) mice to generate *Ulk1*/*2*-cdko [*Ulk1*^+/−;*Ulk2*^+/−* Nestin-Cre (tg)] mice. Controls used in the experiments included *Ulk1*^+/−;*Ulk2*^floxed/floxed^ Nestin-Cre (tg) and *Ulk1*^+/−;*Ulk2*^−/−* Nestin-Cre (tg) mice. (D) The deletion of *Ulk1* and *Ulk2* in the cdko mice was verified by RT-qPCR. The levels of *Ulk1* and *Ulk2* mRNA in the hippocampus of 8-wk-old *Ulk1*/*2*-cdko mice (*n* = 3) were normalized to wild-type, age-matched controls (*n* = 2). *P* < 0.001 (Student's *t*-test). (E) Abnormal limb-clasping reflexes in an 8-wk-old *Ulk1*/*2*-cdko mouse (right photograph). When lifted by their tails and slowly lowered toward a horizontal surface, control mice (left photograph) extend their hind limbs and bodies in anticipation of contact. In contrast, *Ulk1*/*2*-cdko mice show a pathologic reflex, clasping their fore and hind limbs. (F) The ink paw-print test, in which the forepaws were marked in blue ink and the hind paws in red ink, revealed normal gait patterns in control and *Ulk1*/*2*-cdko mice. (G) Motor coordination was tested using a rotarod assay. Control (*n* = 5) and *Ulk1*/*2*-cdko (*n* = 7) mice were placed on a rotating rod that was accelerated from 0 to 40 rpm at a rate of 10 rpm/min, and the time spent on the rod was measured. Data shown are the means (± SEM). *P* < 0.001 (Student's *t*-test). (H) Hematoxylin and eosin (H&E)-stained sections of the cerebellum in 16-wk-old control, *Ulk1*/*2*-cdko, and *Atg7*-cko mice demonstrate the loss of Purkinje cell neurons in the *Atg7*-cko mice. Intact Purkinje cell neurons are indicated by arrows. Scale bar: 50 µm. (I) The mean number (± SEM) of Purkinje cell neurons per 200 µM in 4-month-old control mice (*n* = 5), 4-month-old *Ulk1*/*2*-cdko mice (*n* = 3), and 2- to 4-month-old *Atg7*-cko mice (*n* = 2). *P* < 0.001 (ANOVA).
Figure 3-2. Degeneration of pyramidal neurons in the CA1 region of ULK1/2-deficient mice.

(A) Survival of control (n=20) and Ulk1/2-cdko (n=15) mice. (B) Average body weight ±SEM of male (M) and female (F) Ulk1/2-cdko (n=11) mice compared to control (n=26) mice. (C) The average brain weights ±SEM of control (n=5) and Ulk1/2-cdko (n=5) mice did not significantly differ (Student’s t-test). (D) –Representative images of serial brain sections stained with hematoxylin and eosin (H&E) or DAPI and antibodies against the neuronal marker NeuN. Scale bars: 200 µm (E) Average number of pyramidal neurons (normalized to that in littermate controls) ±SEM in a 500-µm² area of CA1 (n=3 mice/genotype for each age group). *P <0.001 (Student’s t-test) when compared with control. (F) Representative images of brain sections from 16-wk-old mice stained with antibodies against NeuN and the cerebellar Purkinje cell marker, calbindin. Scale bar: 500 µm. (G) Representative images of Fluoro-Jade C stained and cleaved Caspase 3 (Casp-3) immunostained brain sections from 16-wk-old mice. The CA1 region is indicated by brackets. Scale bars: 200 µm (Fluoro-Jade C); 50 µm (cleaved Casp-3). (H) Representative images of serial brain sections from hippocampal region of 16-wk-old mice stained with antibodies against GFAP or IBA1 and counterstained with anti-NeuN and DAPI. Scale bars: 50 µm.
Brain histology at various time points. Hematoxylin and eosin staining showed progressive degeneration of pyramidal neurons in the hippocampal CA1 region after 3 wk (Figure 3-2D, E). The dystrophic hippocampal neurons also showed loss of expression of the neuronal marker NeuN (Figure 3-2D). The hippocampal degeneration in these mice was more severe than that in age-matched Atg7-cko mice (Figure 3-2D). In contrast, neuronal loss from other brain regions (e.g., cerebellum) was more severe in Atg7-cko mice than in Ulk1/2-cko mice (Figures 3-2F and Figure 3-1H, I). The hippocampal degeneration in Ulk1/2-cko mice was accompanied by increased fluoro-Jade C+ staining, which labels degenerating neurons, and activated caspase-3 staining in the CA1 region (Figure 3-2G). Immunostaining for the glial marker GFAP and microglial marker IBA1 was increased in the hippocampal CA1 region (Figure 3-2H), which was consistent with glial activation in response to neuronal damage. These histopathologic changes were not seen in littermate controls. These results highlight the functional redundancy of ULK1 and ULK2 in the CNS and the differential sensitivity of various neuronal populations to their loss.

ULK1/2 Deficiency Activates the Unfolded Protein Response in Hippocampal Neurons

To determine whether the hippocampal degeneration in Ulk1/2-cko mice is associated with defective autophagy, we examined steady-state levels of the ubiquitin-binding protein P62/SQSTM1 (hereafter referred to as P62), an autophagy substrate whose levels correlate inversely with autophagy flux (Ichimura and Komatsu, 2010). we did not detect any P62+ or ubiquitin+ inclusions in the hippocampal neurons of Ulk1/2-cko mice (Figure 3-3A). These results indicate that in neurons, the autophagy-mediated turnover of ubiquitinated proteins that occurs under basal physiologic conditions proceeds in the absence of ULK1 and ULK2, and that defective autophagy is probably not the primary cause of neuronal degeneration in Ulk1/2-cko mice.

To learn the cause of the neuronal degeneration in Ulk1/2-cko mice, we examined the hippocampal region by transmission electron microscopy. Although we did not detect any accumulation of the atypical membranous structures or abnormal mitochondria that are seen with deficiency of other autophagy-related genes (Hara et al., 2006; Komatsu et al., 2006; Liang et al., 2010), the ER compartment was expanded (Figure 3-3B, C). Because ER expansion (Schuck et al., 2009) and mTOR inhibition in neurons (Di Nardo et al., 2009) can result from ER stress and activation of the unfolded protein response (UPR) pathway, which may ultimately lead to cell death, we examined the expression of markers associated with UPR activation in hippocampal pyramidal neurons. Three major sensors of ER stress initiate the UPR: PERK, ATF6, and IRE1. These sensors are activated upon release from the ER chaperone BiP, which is competed away by accumulation of misfolded proteins in the ER (Walter and Ron, 2011). We did not detect spliced XBP1, an alternative product that is produced by activating IRE1 in mRNA samples prepared from the CA1 region. However, the levels of phosphorylated eIF2α, a substrate of PERK, and nuclear localization of ATF6 were both increased in the CA1 region of Ulk1/2-cko mice with degenerating neurons (Figure 3-3D, E).
Figure 3-3.  *Ulk1/2* deficiency in hippocampal neurons is associated with activation of the UPR pathway.

(A) Representative images of brain sections from 4-wk-old *Atg7*-cko, 8-wk-old *Ulk1/2*- cdko and 8-wk-old control mice stained with antibodies against ubiquitin and P62, and counterstained with DAPI. Scale bars: 200 µm. (B) Representative electron micrographs of neurons in CA1 region from 8-wk-old mice. Scale bars: 1 µm. (C) Morphometric analyses of electron micrographs of hippocampal CA1 neurons from 8-wk-old control (*n=2*) and *Ulk1/2*-cdko (*n=2*) mice. *P* < 0.001 (Student's *t*-test). (D-F) Representative images of brain sections from the hippocampal region of16-wk-old mice stained with antibodies against p-eIF2α, ATF6 or CHOP. Sections were counterstained with anti-NeuN and DAPI. Scale bars: 50 µm. (G) Diagram of the UPR pathway, highlighting the activation of the PERK and ATF6 arms in the *Ulk1/2*-cdko mice.
Consistent with activation of the PERK–eIF2α and ATF6 pathways in the CA1 region of \textit{Ulk1/2}-cdko mice, the levels and nuclear localization of CHOP, a transcription factor that promotes ER stress–mediated apoptosis, were increased (\textbf{Figure 3-3F, G}). These results indicate that the degeneration of \textit{Ulk1/2}-deficient pyramidal neurons in the CA1 region disrupts a process that leads to ER stress and UPR activation but not the accumulation of p62+/ubiquitin+ inclusions or membranous structures seen in mice lacking other autophagy genes (e.g., \textit{Fip200}, \textit{Atg5}, or \textit{Atg7}).

\textbf{ULK1 and ULK2 Mediate the Phosphorylation of SEC16A}

We used an unbiased proteomics-based approach to identify ULK/ATG1-interacting proteins and gain insight into ULK/ATG1 function. We analyzed immunoprecipitates (IPs) of endogenous ULK1 from WT MEFs by liquid chromatography/mass spectrometry. SEC16A was among proteins identified only in ULK1-containing samples. The ULK1–SEC16 interaction was confirmed by immunoblot analyses of ULK1 IPs in WT MEFs (\textbf{Figure 3-4A}) and hippocampal extracts from WT mice (\textbf{Figure 3-4B}). SEC16A is a large protein that localizes to ER-exit sites (ERES) and facilitates the recruitment of soluble subunits of the coatomer protein complex II (COPII), including SEC23, SEC24, SEC13, and SEC31 (Miller and Barlowe, 2010). The COPII machinery distinguishes transmembrane and luminal secretory cargo from resident ER proteins and packages them into transport vesicles destined for Golgi. Defects in ER-to-Golgi trafficking can lead to the accumulation of protein in the ER and activation of the UPR (Fang et al., 2015; Preston et al., 2009); therefore, we further characterized the interaction between the ULKs and SEC16A.

ULK1 exists in a complex with ATG13, FIP200, and ATG101; the stability of the complex and ULK1 levels are reduced in ATG13-deficient cells (Hosokawa et al., 2009). RNAi-mediated silencing of \textit{Atg13} expression reduced ULK1 steady-state levels but did not diminish the ULK1–SEC16A interaction (\textbf{Figure 3-4C}). Given the high degree of homology between ULK1 and ULK2 in their N-terminal domains (Mizushima et al., 2011), it was not surprising that we detected MYC-DDK–tagged ULK2 in GFP–SEC16A+ IPs (\textbf{Figure 3-4D}). These results suggest that the interaction between the ULKs and SEC16A does not require ATG13.

Many ULK-interacting proteins are phosphorylated by ULK1 and ULK2. Therefore, we wondered if SEC16A is an ULK substrate. To test this hypothesis, we used a phospho-serine/threonine–specific antibody that recognizes phosphorylated serine/threonine residues with tyrosine, tryptophan, or phenylalanine at the −1 position or phenylalanine at the +1 position [hereafter referred to as p(S/T)Phe] (Kalabis et al., 2006). This antibody is predicted to recognize the S15 (MQVSFVCQ) and S30 (LDTSFKILD) residues of Beclin-1 and S249 (ILKSFELVK) of VPS34 that are phosphorylated by ULK1 (Egan et al., 2015; Russell et al., 2013). GFP–SEC16A coexpressed with ULK1 or ULK2 but not with the kinase-inactive (KI) ULK1 mutant (K46A) showed robust serine phosphorylation (\textbf{Figure 3-4D, E}). These results indicate that SEC16A is phosphorylated in an ULK1/2-dependent manner.
Figure 3-4. ULKs mediate the phosphorylation of SEC16A and facilitate in vitro budding of COPII vesicles.

(A) Representative immunoblot analyses of IPs from WT MEFs transfected with either control nontargeting siRNA (Ctrl) or Sec16a siRNA. IPs were performed using anti-ULK1 antibodies in the presence or absence of epitope-specific blocking peptides (BP).

(B) Representative immunoblot analyses of ULK1 IPs from the hippocampus of 4-wk-old WT mice. (C) Representative immunoblot analyses of ULK1 IPs from WT MEFs transfected with either control nontargeting siRNA (Ctrl) or Atg13 siRNA. (D) Representative immunoblot analyses of 293T cells transfected with GFP–SEC16A and either C-terminal–MYC-DDK–tagged ULK1 or ULK2 expression constructs.

Phosphorylated SEC16A was detected using the anti-p(S/T)Phe antibody. (E) Representative immunoblot analyses of GFP IPs from 293T cells transfected with GFP or GFP–SEC16A and untagged WT or KI ULK1 expression constructs. (F) Representative low- and high- magnification pseudocolored images of Ulk1-ko MEFs cotransfected with ULK1 and GFP–SEC16A and stained with antibodies against ULK1. Sections were counterstained with DAPI. (G) Line scans showing the degree of colocalization between ULK1 (red) and SEC16A (green) in lines (not shown) drawn within the boxes labeled 1-4 in the high magnification images from panel F. Scale bars: 10 μm (low magnification) and 5 μm (high magnification).

(H) Diagram of the in vitro COPII-budding reaction. (I) Representative immunoblot analyses of budded vesicles from in vitro COPII-budding reactions using cytosolic fractions from WT or Ulk1-ko MEFs and membrane fractions from Ulk1-ko MEFs. RPN1 is an ER-resident protein; LMAN1/ERGIC53 and SEC22B are COPII cargo whose incorporation into budded vesicles is inhibited by SAR1H79G, a dominant-negative form of the COPII-specific GTPase. (J) Representative immunoblot analyses of budded vesicles from in vitro COPII-budding reactions using cytosolic fractions from 293T cells transfected with empty vector (—), WT, or KI ULK1 combined with membrane fractions from untransfected 293T cells.
ULK1 Activity Regulates the Budding of COPII Vesicles in Vitro

The assembly and budding of COPII vesicles is initiated upon activation of the small COPII GTPase SAR1. Sar1 promotes the assembly of soluble SEC23–SEC24 heterodimers, which form the inner coat of the transport vesicles, and soluble SEC13–SEC31 heterodimers, which form an outer lattice (Zanetti et al., 2012). SEC16A has a punctate-distribution pattern in cells in culture, consistent with its localization to ERES, where it facilitates the assembly of the soluble COPII components (Bhattacharyya and Glick, 2007; Watson et al., 2006). ULK1 localized to a subset of GFP–SEC16A⁺ puncta (Figure 3-4F, G), suggesting that it is recruited to ERES, where it may regulate COPII transport.

We examined the contribution of ULK1 activity in the budding of COPII vesicles in vitro. The assay combines the cargo-containing membrane fraction from 1 cell source with cytosolic extracts, which supply soluble COPII proteins and putative regulatory proteins, from another cell source (Figure 3-4H). Cytosolic extracts derived from autophagy-deficient Atg5-ko MEFs support COPII budding in vitro; and the incorporation of 2 endogenous cargo, SEC22B and LMAN1, into budded COPII is not inhibited by adding the type-III PI3 kinase inhibitor 3-methyladenine (Ge et al., 2014), indicating that essential components of the autophagy machinery are not involved in the budding reaction. In contrast, budded COPII (i.e., SAR1-dependent) vesicles containing SEC22B and LMAN1 were reduced in reactions containing cytosolic extracts derived from Ulk1-ko MEFs (Figures 3-4I). These results suggest a role for ULK/ATG1 activity in the budding of COPII vesicles and/or incorporation of cargo into COPII vesicles.

ULK1 and Related Kinases Have a Conserved Role in Assembling COPII Components at ERES

Because SEC16A promotes the assembly of soluble COPII components at ERES, and ULK1 localized to a subset of SEC16A⁺ puncta in MEFs and was required for efficient formation of LMAN1 and SEC22B-containing vesicles in vitro, we wondered if ULK1 and homologs regulate the assembly of COPII components at ERES. First, we examined the staining pattern of endogenous SEC16 in oocytes from loss-of-function mutant unc-51(e369) Caenorhabditis elegans. The proportion of high-intensity SEC16⁺ ERES in unc-51 mutant oocytes was significantly lower (P <0.05) than that in WT controls (Figure 3-5A, B). The amount of SEC13 in high molecular–weight gel-filtration fractions, which contain fully assembled COPII complexes comprising SEC13-containing heterotetramers and SEC23-containing heterodimers, was also lower (P <0.05) (Figure 3-5C). These results indicate that UNC-51/ATG1 is involved in the proper assembly of ERES in C. elegans.

The proportion of high-intensity SEC16A⁺ ERES in Ulk1/2-dko MEFs (Figure 3-5D, E) was significantly reduced (P <0.05), similar to that in unc-51–mutant oocytes (Figure 3-5A, B). Silencing Sec16a expression in mammalian cells decreases the accumulation of soluble COPII components at ERES (Watson et al., 2006). Given the
Figure 3-5. ULKs regulate the assembly of COPII complexes at ERES. 

(A) Representative low- and high-magnification images of distal gonads from WT and unc-51–mutant C. elegans stained using a fluorescently labelled anti-SEC16 antibody. 

(B) Mean percentages ±SEM of ERES that fall within specified intensity thresholds of SEC16 staining. For each condition, at least 1000 unique ERES from C. elegans gonads were examined. Scale bars: 10 µm. *P <0.05 (Student’s t-test) when compared with WT. 

(C) Mean percentages (±SEM) of total SEC13 in peaks 1-3 from Superose 6 gel filtration experiments similar to the one shown in Figure S4A. *P <0.05 (Student’s t-test) when compared with WT. 

(D) Representative images of endogenous SEC16A immunostaining in WT, Ulk1-ko, Ulk2-ko, and Ulk1/2-dko MEFs. Scale bars: 10 µm. 

(E) Mean percentages (±SEM) of all ERES that fall within specified intensity thresholds of SEC16A staining. *P <0.001 and †P <0.05 (ANOVA) when compared with WT. 

(F) Mean number (±SEM) of SEC24+ puncta per cell in WT MEFs transfected with the indicated HA-tagged SEC24 isoform (A, B, C, or D) and control nontargeting (Ctrl), Ulk1, or Sec16a siRNA. Ten HA+ cells per population were scored. Red arrow highlights the decrease in SEC24C puncta number in Ulk1-depleted cells. *P <0.001 and †P <0.05 (ANOVA) when compared with siCtrl. 

(G) Mean number (±SEM) of SEC24C+ puncta per cell in WT, Ulk1-ko, Ulk2-ko, and Ulk1/2-dko MEFs. Ten cells per population were scored. *P <0.001 (ANOVA) when compared with WT. 

(H) Representative images of endogenous SEC24C staining in WT and Ulk1-ko MEFs. Scale bars: 10 µm. 

(I) Mean number (±SEM) of SEC24C+ puncta per cell in Ulk1/2-deficient (i.e., Ulk1 ko and Ulk2 shRNA) MEFs stably transduced with the indicated viral vector [i.e., (—) empty vector; WT ULK; or KI ULK1 mutant]. Ten cells per population were scored. *P <0.001 (ANOVA) when compared with empty vector-transduced cells. 

(J) Mean number (±SEM) of SEC24C+ puncta per cell in WT MEFs transfected with the indicated siRNA. Ten cells per population were scored. *P <0.001 (ANOVA) when compared with Ctrl siRNA-transfected cells.
reduced number of high-intensity SEC16A+ puncta in Ulk1/2-deficient MEFs, we compared the effects of silencing Sec16a and Ulk1 in WT and Ulk2-ko MEFs on the accumulation of cargo adapters, SEC24 A-D, at ERES (Figures 3-5F). RNAi-mediated knockdown of Sec16a significantly reduced (P <0.001) the number of puncta formed by each of the overexpressed SEC24 proteins in MEFs (Figures 3-5F). To our surprise, RNAi-mediated knockdown of Ulk1 in WT or Ulk2-ko MEFs selectively reduced the number of SEC24C+ puncta (Figures 3-5F). The number of endogenous SEC24C+ puncta was also significantly reduced (P <0.001) in Ulk1-ko and Ulk1/2-dko MEFs (Figure 3-5G, H), which is consistent with impaired assembly of SEC24C-containing COPII complexes at ERES in cells lacking ULK1. The defect in ERES assembly in either Ulk1- or Ulk1/2-deficient MEFs was rescued by ectopic expression of WT ULK1 but not the KI mutant (Figure 3-5I), indicating that ULK1 catalytic activity is required for recruiting SEC24C to ERES. Whereas transient RNAi-mediated knockdown of Ulk1 or Sec16a in WT MEFs caused significant decreases (P<0.001) in the number of SEC24C+ puncta, knockdown or knockout of Atg7, Atg13, or Atg14 did not (Figure 3-5J).

These results indicate that ULK/ATG1 has an evolutionarily conserved role in regulating ERES assembly, which is not shared by other autophagy-related proteins. ULK1 and ULK2 activities appear to converge on multiple aspects of SEC16A function, including the normal accumulation of SEC16A at ERES, and the assembly of SEC24C-containing ERES. These functions, however, appear to have different thresholds of sensitivity to loss of ULK/ATG1 function. SEC24C expression was relatively increased in hippocampal neurons, compared to the other SEC24 isoforms and neurons in other brain regions (data not shown). Therefore, these neurons may be particularly sensitive to the disrupted assembly of SEC24C-containing ERES, which occurs in the absence of Ulk1/2 expression.

**ULK1 and Related Kinases Regulate ER-to-Golgi Trafficking**

Because Ulk1/2-deficiency impaired the accumulation of SEC24C at ERES and only minimally affected VSVG trafficking (data not shown), which relies to a greater extent on the A and B isoforms of SEC24 than on the C and D isoforms for ER export (Bonnon et al., 2010), we examined the role of the ULKs in trafficking SEC24C-specific cargo. The serotonin transporter SERT interacts specifically with SEC24C, and its PM localization depends on SEC24C-dependent export from the ER (Montgomery et al., 2014; Sucic et al., 2011). SERT is expressed on the surface of platelets and specific types of neurons. It regulates the concentration of serotonin (5-HT) at synapses and in blood plasma via uptake of 5-HT into neurons and platelets, respectively (Mercado and Kilic, 2010). Ulk1-ko MEFs showed reduced levels of the Endo H–resistant forms of SERT-GFP in whole-cell extracts (Figure 3-6A, lower panels). Similarly, transient RNAi-mediated knockdown of Sec16a or Ulk1 in WT MEFs significantly reduced (P <0.05 and P <0.01, respectively) the ratio of Endo H–resistant to Endo H–sensitive SERT-GFP (Figures 3-6B). However, knockdown of other autophagy-related genes (e.g., Atg7, Atg13, or Atg14) had no effect on ER-to-Golgi trafficking of SERT-GFP (Figures 3-6B).
**Figure 3-6. ULKs regulate ER-to-Golgi trafficking.**

(A) Representative immunoblots of Biotin IPs (top panels) and GFP IPs (bottom panels) from SERT-GFP—transfected WT and *Ulk1*-ko MEFs. WT MEFs were incubated in the presence or absence of Endo H or peptide-N-glycosidase F (PNGase) to establish the migration pattern of the different glycosylated forms of SERT. (B) Mean ratios (±SD) of Endo H–R SERT to total SERT in RNAi-treated samples from 2 independent experiments. *P < 0.01 and #P < 0.05 and (ANOVA) when compared with Ctrl. (C) Mean percentages (±SEM) of cells showing colocalization of AlexaFluor 647–conjugated wheat germ agglutinin (WGA) and SERT-GFP. Data were acquired from 3 independent experiments, and more than 100 cells per population were scored. *P < 0.001 (ANOVA) when compared with WT. (D) Representative merged pseudocolored images of SERT-GFP—transfected WT and *Ulk1*-ko MEFs stained with AlexaFluor 647–conjugated wheat germ agglutinin (WGA) and DAPI. Scale bar: 10 µm. (E) Mean percentages (±SEM) of cells with colocalized WGA and SERT-GFP. Data were acquired from 3 independent experiments, and more than 100 cells per population were scored in each experiment. *P < 0.002 (ANOVA) when compared with empty vector–transduced cells. (F) Mean percentages (±SEM) of siRNA transfected cells with colocalized WGA and SERT-GFP. Data were acquired from 3 independent experiments, and more than 100 cells per population were scored. *P < 0.001 (ANOVA) when compared with *siCtrl*-transfected cells. (G) Representative images of WT, *unc-51* mutant and mod-5 mutant *C. elegans* stained with antibodies against MOD-5/SERT and 5-HT. The arrows highlight the 5-HT or MOD-5/SERT staining of NSM processes.
Treating cells with NHS-biotin prior to lysis allows selective biotinylation of surface proteins and provides a sensitive biochemical method of detecting defects in trafficking of proteins to the PM. Consistent with defective ER-to-Golgi trafficking of SERT, the levels of biotinylated surface SERT were reduced in anti-biotin IPs prepared from \(\text{Ulk1-ko}\) MEFs (Figure 3-6A, upper panels). \(\text{Ulk1-ko}\) and \(\text{Ulk1/2-dko}\) MEFs also showed significant reduction \((P < 0.001)\) in PM localization of SERT–GFP (Figure 3-6C, D). The defect in cell surface localization of SERT in \(\text{Ulk1/2-deficient}\) MEFs was rescued by expressing WT but not KI ULK1 (Figure 3-6E), indicating that ULK1 catalytic activity is required for SERT trafficking. Transient RNAi-mediated knockdown of \(\text{Sec16a}\) or \(\text{Ulk1}\) significantly reduced \((P < 0.001)\) SERT localization to the PM of WT MEFs, but knockdown of other autophagy-related genes (e.g., \(\text{Atg7}, \text{Atg13}, \text{or Atg14}\)) did not (Figure 3-6F).

In \(\text{C. elegans}\), SERT (MOD-5) is expressed on the PM, along the length of axons of 5-HT–producing pharyngeal neurosecretory motor (NSM) neurons, with minimal localization of SERT/MOD-5 in cell bodies. Thus, \(\text{C. elegans}\) is a sensitive system for detecting defects in SERT trafficking in neurons in vivo. In contrast to its pattern of expression in WT NSM neurons, SERT/MOD-5 accumulated in the cell bodies of \(\text{unc-51}\)–mutant NSM neurons and failed to localize to 5-HT–stained NSM axons (Figure 3-6G). These findings are consistent with a defect in ER-to-Golgi trafficking associated with disruption of UNC-51/ATG1 function. In addition to NSM neurons, AIM and RIH interneurons express SERT/MOD-5. These interneurons do not synthesize 5-HT; instead, they regulate 5-HT levels at extrasynaptic 5-HT targets via uptake mediated by SERT/MOD-5 (Jafari et al., 2011). As reported previously, SERT/MOD-5 is essential for accumulating 5-HT in AIM and RIH interneurons (Jafari et al., 2011), and these neurons in loss of function \(\text{mod-5}\)–mutant \(\text{C. elegans}\) showed no 5-HT accumulation (Figure 3-6G). However, the \(\text{unc-51}\)–mutant \(\text{C. elegans}\) showed normal levels of 5-HT accumulation (Figure 3-6G), suggesting that SERT trafficking and SERT-mediated 5-HT uptake is normal in AIM and RIH neurons. The basis for the cell-type specificity of ULK/ATG1-mediated SERT trafficking in \(\text{C. elegans}\) remains unknown. These findings indicate that ULK1 and homologs have an evolutionarily conserved and cell-type specific regulatory role in the trafficking of cargo such as SERT, which is not shared by other autophagy-related proteins, including ATG13.

**ULK-Mediated Phosphorylation of SEC16A at S846 Is Required for ER-to-Golgi Trafficking and ERES Assembly**

ULKs mediate SEC16A phosphorylation, and both proteins function in ER-to-Golgi trafficking; thus, we investigated the role of ULK-mediated phosphorylation of SEC16A in this process. As mentioned above, we detected ULK1/2-dependent phosphorylation of SEC16A by using a p(S/T)Phe antibody. Site-directed mutagenesis was used to substitute serine and threonine residues in SEC16A that matched the antibody-recognition motif with alanines. The resulting GFP–SEC16A mutants were screened for their ability to disrupt ULK1-dependent phosphorylation (data not shown). We observed a reproducible decrease in the ULK1-dependent phosphorylation signal on
the GFP–SEC16A S846A–mutant protein compared to WT (data not shown). To confirm
that S846 was phosphorylated in an ULK1-dependent manner, we reintroduced the
S846A mutation into a MYC-DDK–tagged SEC16A-expression construct. Use of this
construct demonstrated that the S846A substitution was sufficient to abolish the ULK1-
dependent phosphorylation signal on SEC16A that was recognized by the p(S/T)Phe
antibody (Figure 3-7A).

To determine the consequence of ULK-mediated phosphorylation of SEC16A at
S846, we expressed WT SEC16A, the phospho-defective S846A mutant, and the
phosphomimetic S846D mutant in Sec16a-depleted cells and assessed their ability to
restore the defects in ERES assembly and ER-to-Golgi trafficking common to Ulk1/2-
deficient and Sec16a-depleted MEFs. First, we detected fewer SEC16A+ puncta in
Sec16a-depleted cells upon overexpression of the S846A mutant than in cells expressing
WT SEC16A or the S846D mutant, despite comparable levels of protein (Figure
3-7B, C). The number of SEC24C+ puncta was also fewer in Sec16a-depleted cells
expressing the S846A mutant than in those expressing WT SEC16A or the S846D mutant
(Figure 3-7B, D). These results are consistent with the notion that ULK-mediated
phosphorylation of SEC16A is required for efficient recruitment and/or retention of
SEC16A and SEC24C at ERES. WT SEC16A did not increase the number of SEC24C+
puncta in Ulk1-deficient MEFs (Figure 3-7E, F), suggesting that the residual Ulk2
expression in these MEFs was insufficient to yield high enough levels of phosphorylated
SEC16A to promote accumulation of SEC24C at ERES. In contrast, expressing the
S846D mutant in Ulk1-deficient MEFs significantly increased (P <0.001) the number of
SEC16A+ puncta and SEC24C+ puncta and the proportion of puncta showing
colocalization of SEC16A and SEC24C (Figure 3-7E, H). Similarly, the defect in SERT
trafficking in Ulk1-ko MEFs was rescued by overexpressing the S846D SEC16A mutant
but not the WT SEC16A or the S846A mutant (Figure 3-7I, J). These data indicate that
ULK1/2-mediated phosphorylation of SEC16A at S846 drives the assembly of COPII
components at ERES and trafficking of associated secretory cargo.

The Defect in ER-to-Golgi Trafficking in ULK-Deficient Cells Leads to an ER–
Stress Response That Is Reversed by the S846D SEC16A Mutant

Defects in ER-to-Golgi trafficking can cause proteins to accumulate in the ER and
activate the UPR system, but we did not detect any evidence of UPR activation in Ulk1-
and/or Ulk2-ko MEFs under basal growth conditions (data not shown) or in Ulk1-ko
MEFs expressing VSVG–GFP (Figure 3-8A). By contrast, Ulk1-ko MEFs expressing
SERT–GFP showed a significant increase (P <0.001) in nuclear localization of CHOP
(Figure 3-8A, B). Moreover, expression of the S846D SEC16A mutant in Ulk1-ko
MEFs, which rescued the defect in ER-to-Golgi trafficking of SERT, prevented nuclear
localization of CHOP (Figure 3-8C, D). These findings indicate that the defect in ULK-
and SEC16A-dependent ER-to-Golgi trafficking of certain cargo activates the UPR
pathway, similar to that observed in Ulk1/2-deficient hippocampal neurons.”
Figure 3-7. ULK-mediated phosphorylation of SEC16A at S846 is required for assembly of SEC24C+ ERES and ER-to-Golgi trafficking.

(A) Representative immunoblots of DDK IPs prepared from 293T cells transfected with either MYC-DDK–tagged WT SEC16A or a mutant form of MYC-DDK–tagged SEC16A harboring the S846A and the specified ULK1-expression construct. (B) Representative pseudocolored images of MEFs stained with DAPI and antibodies against MYC-DDK-tagged SEC16A and SEC24C. SA: SEC16A S846A mutant; SD: SEC16A S846D mutant. (C) Mean numbers (±SEM) of SEC16A+ puncta per cell. (D) Mean numbers (±SEM) of SEC24C+ puncta per cell. Ten cells (DDK+, if appropriate) per population were scored in C and D. *P <0.001 (ANOVA). (E) Representative pseudocolored images of MEFs stained with DAPI and antibodies against SEC24C and MYC-DDK-tagged SEC16A. (F) Mean numbers (±SEM) of SEC24C+ puncta per cell. (G) Mean numbers (±SEM) of SEC16A+ puncta per cell. (H) Mean percentages (±SEM) of SEC16A+ puncta that were also SEC24C+. Ten cells (DDK+, if appropriate) per population were scored in F-H. Scale bar = 10 µm. *P <0.001 and #P <0.05 when compared with WT (ANOVA). (I) Representative merged pseudocolored images of MEFs stained with AlexaFluor 647–conjugated WGA and antibodies against MYC-DDK-tagged SEC16A. White arrowheads indicate PM. (J) Mean percentages (±SEM) of cells from each population with colocalized WGA and GFP. Data were acquired from 3 independent experiments, and more than 100 cells per population were scored in each experiment. Scale bar: 10 µm. *P <0.001 (ANOVA) when compared with empty vector.
Figure 3-8.  SEC16A mutant with the phosphomimetic S846D overcomes the defect in ER-to-Golgi trafficking and prevents activation of the UPR in ULK-deficient cells.

(A) Mean percentages (±SEM) of cells with nuclear staining of CHOP. Data were acquired from 3 independent experiments, and more than 100 GFP+ cells per population were scored in each experiment. *P < 0.001 (ANOVA) when compared with empty vector-transduced cells. (B) Representative pseudocolored images of WT and Ulk1-ko MEFs transfected with indicated GFP-tagged cargo [i.e, (−), empty vector; VSVG-GFP or SERT-GFP] stained with an antibody against CHOP and DAPI. Scale bar: 10 µm. (C) Mean percentages (±SEM) of cells with nuclear staining of CHOP. Data were acquired from 3 independent experiments, and more than 100 GFP+DDK+ cells per population were scored in each experiment. *P < 0.001 (ANOVA) when compared with empty vector-transduced cells. SA: S846A SEC16A mutant; SD, S846D SEC16A mutant. (D) Representative pseudocolored images of MEFs stained with antibodies against MYC-DDK-tagged SEC16A (channel not shown) and CHOP and DAPI. Scale bar: 10 µm.
CHAPTER 4. AUTOPHAGY-INDEPENDENT ROLE OF ULK1 AND ULK2 IS ESSENTIAL FOR AXON GUIDANCE IN THE DEVELOPING MOUSE FOREBRAIN

_Ulk1/2_-Deficiency in the CNS Results in Abnormal Axon Guidance

Because of the observation that approximately 40% of CNS specific _Ulk1/2_-cdko mice die shortly after birth and the previously described role of ULK/ATG1 homologues in axon guidance and growth, we decided to examine the integrity of the major axon tracts formed by projection neurons in _Ulk1/2_-DKO mice. Towards this end, we interbred _Ulk1^+/−;Ulk2^−/−_ mice. Twice the number of _Ulk1^+/--;Ulk2^−/−_ (63/206; 30.58%) compared to _Ulk1^−/--;Ulk2^−/−_ (29/206; 14.08%) embryos were identified upon genotyping at E18.5, suggesting that loss of both _Ulk1_ and _Ulk2_ increased embryonic mortality. Nevertheless, we observed no significant difference in body weight (Figure 4-1A) or gross brain morphology (Figure 4-1B) in E18.5 _Ulk1/2_-DKO embryos compared to littermate controls (_Ulk1^+/+;Ulk2^−/−_ or _Ulk1^+/−;Ulk2^−/−_). Visual inspection of newborn pups and genotyping of pups on or after post-natal day 1 (P1) confirmed the previously reported perinatal mortality of live born _Ulk1/2_-DKO pups (Cheong et al., 2011; Cheong et al., 2014; Kundu et al., 2008).

Commissural projection neurons project their axons to the contralateral cortex, thus connecting the two cortical hemispheres. The cell bodies of commissural projection neurons in rodents reside primarily in cortical layers II/III (80%), layer V (20%) and to a lesser extent layer VI; their axons traverse the telencephalic midline, giving rise to three commissural tracts: the corpus callosum (CC), the hippocampal commissure (HC) and the anterior commissure (AC). We used an antibody against the axonal marker L1 cell adhesion molecule (L1) to examine the three commissural tracts in E18.5 embryos. The CC was thinner in _Ulk1/2_-DKO embryos compared to that in littermate controls (Figure 4-1C) and many of the callosal axons were overfasciculated (Figure 4-1D1, D1’, D4, D4’). Although the AC in _Ulk1/2_-DKO embryos was severely hypoplastic, the HC showed no overt abnormalities (Figure 4-1D2, D2’, D5, D5’).

We also examined the integrity of the corticothalamic axons (CTAs), which originate from neurons residing in layer VI of the cortex and project to the thalamus, and thalamocortical axons (TCAs), which originate from neurons in various thalamic nuclei and project to corresponding cortical areas. The CTAs and TCAs of _Ulk1/2_dko mice showed several abnormalities as they crossed the pallial-subpallial boundary (PSPB). Instead of forming a parallel fan-like structure, the _Ulk1/2_-deficient axons were disorganized, overfasciculated and traversed the PSPB more anteroventrally than axons in littermate controls (Figure 4-1D3, D3’, D6, D6’, E). The abnormal path of the CTAs and TCAs in _Ulk1/2_-deficient animals was already evident at E14.5, around the time when the CTAs and TCAs first begin crossing the PSPB (Figure 4-1F-G), suggesting that ULK1 and ULK2 regulate an intermediate step in axon pathfinding (i.e. crossing midline or specific boundaries).
Figure 4-1. **ULK1 and ULK2 are required for axonal guidance in the forebrain.**

(A) No significant difference in body weight of control (Ulkl<sup>+/+</sup>;Ulkl<sup>−/−</sup> and Ulkl<sup>+/−</sup>;Ulkl<sup>−/−</sup>) and DKO mice at E18.5. (B) No overt differences in gross brain morphology between control and DKO mice at E18.5. (C) Quantification of the dorsoventral width of the CC of control (n = 5) and Ulkl/2-DKO mice (n = 5) at E18.5. The average CC width of each mutant was normalized to that of its corresponding control. **P<0.01. (D) Brain sections of E18.5 control (D1, D2, D3) and Ulkl/2-DKO (D4, D5, D6) were stained with antibodies against the general axonal marker L1 (pseudocolored green). The callosal axons are heavily overfasciculated, and the corpus callosum is thinner in Ulkl/2-DKO mice (D1′, D4′). The anterior commissure fails to develop (D2′, D5′). CTAs and TCAs progressed abnormally as they cross the PSPB (D3′, D6′). The Ulkl/2-DKO axons are highly disorganized and extend aberrantly towards the external capsule (EC) (arrows in D5-D6). The CTAs and TCAs are also overfasciculated. (E) Sagittal view of the the abnormal CTA and TCA tracts highlighted by arrowheads. (F) The axon pathfinding defects in Ulkl/2-DKO mice are detected as early as E14.5 (dashed lines), and (G) number of axons crossing PSPB is quantified from control (n = 3) and Ulkl/2-DKO mice (n = 3). Abbreviations: AC, anterior commissure; CC, corpus callosum; dTh, dorsal thalamus; St, striatum. Scale bars: 500 µm in D1-D6, and E, G; 250 µm in D1′-D6′.
ULK1 and ULK2 Regulate Axon Guidance via an Autophagy-Independent Pathway

FIP200 and ATG13 are components of the canonical 3-MDa ULK/ATG1 kinase complex that regulates both selective and non-selective autophagy in response to cellular stress (Alers et al., 2012; Mizushima, 2010). ULK/ATG1 also interacts with the COPII scaffold SEC16A to promote ER-to-Golgi trafficking of specific cargo in the absence of stress (Joo et al., 2016). Therefore, we immunoprecipitated endogenous ULK1 from extracts prepared from the cerebral cortex of newborn (P0) mice to determine which of these interactions may be relevant in the developing cortex. While we were able to detect the interaction between ULK1, ATG13 and FIP200 in both P0 and adult cortex, the interaction between ULK1 and SEC16A was restricted to P0 cortex (Figure 4-2A), raising the possibility that either the autophagy-inducing function of ULK/ATG1 or its role in ER-to-Golgi trafficking may play an important role in the regulation of axon guidance.

In order to assess the contribution of the autophagy-inducing function of ULK/ATG1 in axon guidance, we decided to investigate the extent of overlap in the neurodevelopmental phenotypes of mice lacking CNS expression of Fip200, Atg7 and Ulk1/2. Mice lacking CNS expression of Ulk1 and Ulk2 (Ulk1+/−;Ulk2−/−;Nestin-Cre; hereafter referred to as Ulk1/2Nestin cDKO mice) show progressive hippocampal degeneration beginning around 8 weeks of age (Joo et al., 2016). Although some of the Ulk1/2Nestin cDKO mice survived up to 4 months, 40% of them died within the first 24 hours after birth (Joo et al., 2016). The surviving mice showed no accumulation of P62+ or Ubiquitin+ deposits in all the cortical layer either (Figure 4-2B) or any appreciable increase in steady state levels of P62 (Joo et al., 2016). After confirming the loss of Ulk1 and Ulk2 expression in the brains of Ulk1/2Nestin cDKO mice by RT-qPCR analyses (Figure 4-3A), we examined L1 staining of brain sections from Ulk1/2Nestin cDKO mice and littermate controls on post-natal day P1. These stains revealed dysgenesis of the corpus callosum (including the presence of Probst bundles), hypoplastic AC, and disruption of CTAs and TCAs in Ulk1/2Nestin cDKO mice (Figure 4-2B), similar to that observed in the Ulk1/2-DKO mice. Mice lacking CNS expression of Fip200 (Fip200Nestin cKO) or Atg7 (Atg7Nestin cKO) showed the expected accumulation of P62+ and ubiquitin+ deposits in all the cortical layers (layer I-VI) examined (Figure 4-2B), but no defects in commissural axon tracts or disruption of CTAs and TCAs (Figure 4-3B).

We also examined the trajectory of unmyelinated axon projections (i.e. mossy fibers) from granule cells in the dentate gyrus to the hippocampal CA3 regions in the brains of Ulk1/2Nestin cDKO, Fip200Nestin cKO, Atg7Nestin cKO and littermate controls at P21. Immunostaining with antibodies against the neuronal marker NeuN and Neurofilament 165 allowed visualization of the mossy fibers located above (i.e., suprapyramidal bundle) and below (i.e., infrapyramidal bundle) the pyramidal cell layer of the CA3 (Figure 4-4A1). In Atg7Nestin cDKO, Fip200Nestin cDKO mice and control animals, the fibers in the infrapyramidal bundle extended along the pyramidal cell layer in the proximal portion of CA3 (i.e closer to the dentate gyrus), and then crossed the pyramidal cell layer at mid and distal portions of CA3 to join the mossy fibers in the suprapyramidal bundle (Figure 4-4A4, A5). Mossy fibers in the Ulk1/2Nestin cDKO mice
Figure 4-2. ULK1 and ULK2 are not required for autophagy-mediated turnover of ubiquitinated protein in the cerebral cortex.

(A) Immunoprecipitation (IP) of ULK1 using extracts prepared from the cortex of P0 and adult WT mice and performed in the presence or absence of epitope-specific blocking peptides (PB). The interaction between ULK1, ATG13 and FIP200 was detected in extracts prepared from P0 and adult cortex; whereas the interaction between ULK1 and SEC16A was only detected in extracts prepared from P0 (but not adult) cortex. (B) Immunostains performed on sections of cortex from control, Ulk1/2Nestin cDKO, Atg7Nestin cKO and FIP200Nestin cKO mice using antibodies against P62 (pseudocolored red) and Ubiquitin (pseudocolored green). Sections were counterstained with DAPI (pseudocolored blue). Scale bars: 50 µm.
Figure 4-3. FIP200 and ATG7 are not required for axon guidance in the forebrain.
(A) Real-time PCR confirms the loss of both Ulk1 and Ulk2 in the brain of Ulk1/2\textsuperscript{Nestin} cDKO mice. (B) L1 staining of serial sections of Ulk1/2\textsuperscript{Nestin} cDKO brains at P1 reveals abnormally fasciculated axons, including in the CTAs and TCAs (inset), corpus callosum dysgenesis and anterior commissure hypoplasia. These axon guidance abnormalities are similar to those identified in the Ulk1/2-DKO mice and are not observed in autophagy-deficient Atg7\textsuperscript{Nestin} cKO or Fip200\textsuperscript{Nestin} cKO mice. Abbreviations: AC, anterior commissure; CC, corpus callosum; IC, internal capsule; EC, external capsule; HC, hippocampal commissure; Pb, Probst bundle. Scale bars: 500 µm
Figure 4-4. ULK1 and ULK2 regulate mossy fiber guidance via an autophagy independent-mechanism.

(A) Sections of hippocampus from adult control, Ulk1/2\textsuperscript{Nestin} cDKO mice, Atg7\textsuperscript{Nestin} cKO, and FIP200\textsuperscript{Nestin} cKO mice were immunostained with antibodies against NF165 and NeuN. Representative images show the abnormal projections from the infrapyramidal bundle in two Ulk1/2\textsuperscript{Nestin} cDKO mice (A2', A3') but not in the Atg7\textsuperscript{Nestin} cKO (A4') or FIP200\textsuperscript{Nestin} cKO mice (A5'). (B) Immunostaining of markers for mitotic cells (Ki67), neuronal stem cells (Sox2), and mature neurons (NeuN) in the dentate gyrus of adult control and Ulk1/2\textsuperscript{Nestin} cDKO mice. (C) The mean number ± SEM of Ki67\textsuperscript{+}, Sox2\textsuperscript{+}, and NeuN\textsuperscript{+} cells in the dentate gyrus showed no statistical significance between controls (n = 3) and Ulk1/2\textsuperscript{Nestin} cDKO (n = 3) mice. Abbreviations: DH: dendate hilus; ipb: infrapyramidal bundle of mossy fibers; SO: stratum oriens; spb: suprapyramidal bundle of mossy fibers; SR: stratum radiatum; ns: not significant. Scale bars: 200 µm in A1-A5, B; 50 µm in A1'-A5'.

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**Figure 4-4.** ULK1 and ULK2 regulate mossy fiber guidance via an autophagy independent-mechanism.

(A) Sections of hippocampus from adult control, Ulk1/2\textsuperscript{Nestin} cDKO mice, Atg7\textsuperscript{Nestin} cKO, and FIP200\textsuperscript{Nestin} cKO mice were immunostained with antibodies against NF165 and NeuN. Representative images show the abnormal projections from the infrapyramidal bundle in two Ulk1/2\textsuperscript{Nestin} cDKO mice (A2', A3') but not in the Atg7\textsuperscript{Nestin} cKO (A4') or FIP200\textsuperscript{Nestin} cKO mice (A5'). (B) Immunostaining of markers for mitotic cells (Ki67), neuronal stem cells (Sox2), and mature neurons (NeuN) in the dentate gyrus of adult control and Ulk1/2\textsuperscript{Nestin} cDKO mice. (C) The mean number ± SEM of Ki67\textsuperscript{+}, Sox2\textsuperscript{+}, and NeuN\textsuperscript{+} cells in the dentate gyrus showed no statistical significance between controls (n = 3) and Ulk1/2\textsuperscript{Nestin} cDKO (n = 3) mice. Abbreviations: DH: dendate hilus; ipb: infrapyramidal bundle of mossy fibers; SO: stratum oriens; spb: suprapyramidal bundle of mossy fibers; SR: stratum radiatum; ns: not significant. Scale bars: 200 µm in A1-A5, B; 50 µm in A1'-A5'.
entered the proximal part of the infrapyramidal region within CA3 and either terminated prematurely, turned prematurely into or aberrantly away from the pyramidal cell layer (Figure 4-4A2, A3). Together, these results suggest that the role of ULK1 and ULK2 in axonal guidance occurs via an autophagy-independent pathway.

FIP200 is essential for maintenance of the neuronal stem cell pools in post-natal brains (Wang et al., 2013), and the dentate gyrus is one of only two brain regions with continuous neurogenesis in adult animals. Therefore, we examined the dentate gyrus of Ulk1/2Nestin cDKO mice in more detail, but observed no obvious difference in the number of neuronal stem cells, mitotic neurons, or mature neurons in the dentate gyrus in 2-4 month old animals (Figure 4-4B, C). Thus, consistent with the notion that the ULKs are dispensable for autophagy in the central nervous system in unstressed animals, the ULKs are not required for the FIP200- and autophagy-mediated post-natal regulation of neural stem cell function.

**ULK1 and ULK2 Are Not Essential for Cortical Neurogenesis**

We next sought to further characterize the neurodevelopmental defects in Ulk1/2-deficient mice in order to gain some insight into non-canonical functions of the mammalian ATG1 homologues. In order to exclude the possibility that the abnormalities in the callosal axons, CTAs and TCAs in Ulk1/2-deficient mice were caused by impaired production of cortical projection neurons, we examined cortical layer structure of E18.5 DKO brains. Nissl staining and the expression of the callosal neuron marker Satb2, Layer V marker Ctip2, and Layer VI marker Tbr1 were all normal in the DKO cortex (Figure 4-5A, B). Thus, ULK1 and ULK2 appear to be dispensable for specification of projection neurons in the embryonic cortex. The Ulk1/2-DKO embryos did not show an increase in apoptosis (Figure 4-5C, D) or neural tube closure defects (data not shown), both of which are seen in Ambra1-deficient animals (Fimia et al., 2007).

**ULK1 and ULK2 Regulate the Midline Crossing of Callosal Axons**

In order to trace the projection of callosal axons across the midline, we embedded Dil crystals into the parietal cortex of control and Ulk1/2-DKO embryos at E18.5. Compared to that of controls, the fluorescence intensity of callosal axons reaching the contralateral cortex was dramatically reduced in the Ulk1/2-DKO brains (Figure 4-6A). Luxol Fast Blue staining for myelinated axons also showed significantly reduced CC thickness in Ulk1/2Nestin cDKO brains at 2 months, indicating persistent deformity of the CC in Ulk1/2-deficient mice (Figure 4-6B, C).

Abnormal CC development can result from cell autonomous defects, or various cell non-autonomous abnormalities, such as environmental/axon guidance cues from midline glia and guidepost neurons. Having ruled out obvious defects in cortical neurogenesis (Figure 4-5), we examined the midline glial structures and guidepost neuron population. Glial cells stained with GFAP were properly formed in the midline
Figure 4-5. ULK1 and ULK2 are not essential for neurogenesis.

(A) Nissl staining (A1-A2) and expression of the callosal neuron marker Satb2, layer V marker Ctip2, and layer VI marker Tbr1 in control and Ulk1/2-DKO cortices at E18.5 (A3-A4), and the quantifications of Satb2+, Ctip2+, and Tbr1+ cells in layer II-IV, layer V, and layer VI, respectively, are statistically not significant (ns), indicating that Ulk1/2 are not required for neurogenesis. (C-D) Staining of cleaved Caspase 3 on cortices from control and Ulk1/2-DKO at E18.5, and number of Caspase 3+ cells per section is quantified (n = 3 for both control and Ulk1/2-DKO). Scale bars: 50 µm in A1-A2, A3’-A4’, and C; 500 µm in A3-A4.
Figure 4-6. ULK1 and ULK2 regulate development of the corpus callosum.

(A) Dil was placed in the parietal cortex of E18.5 Ulk1/2-DKO (n = 2) and control (n = 2) mice to trace the midline crossing of callosal axons. Representative sections on the left and bar graph on the left highlight the dramatic reduction in fluorescence intensity in the contralateral cortex (rostral and caudal levels) of Ulk1/2-DKO mice (asterisks). (B) Luxol Fast Blue and Nissl staining reveals persistent anomalies of the corpus callosum (CC) in the adult Ulk1/2Nestin cDKO mice. (C) The dorsoventral width of the CC of adult control (n = 3) and Ulk1/2Nestin cDKO mice (n = 3) was quantified, and the average width of each mutant was normalized to that of its corresponding control. There is a significant (**) $P<0.01$ reduction in the width of the corpus callosum in Ulk1/2Nestin cDKO mice relative to that in controls. (D) Representative brain sections from E18.5 Ulk1/2-DKO and P0 Ulk1/2Nestin cDKO mice and age-matched controls. The midline glial structure developed properly. (E) Calretinin immunostaining of the glutamatergic guidepost neurons in control and Ulk1/2-DKO brains at E15.5 and E18.5 showed that the production of those neurons was not altered in the Ulk1/2-deficient mice. Scale bars: 500 µm in A1-A4, and B1-B2; 100 µm in A1'-A4', B1'-B2', and D-E.
glial structures, including the glial wedge, indusium griseum glia, and midline zipper glia (Figure 4-6D). The guidepost neurons marked by calretinin appeared normal in the mutants at E15.5; thus, their abnormal distribution along the midline at E18.5 was more likely a consequence of improper positioning of the axons crossing the midline rather than the cause (Figure 4-6E).

Ulk1/2-Deficiency Impairs Organization of the Somatosensory Cortex

We next sought to characterize the axon guidance defects in Ulk1/2-deficient CTAs and TCAs in more detail. First, we embedded crystals of the fluorescent carbocyanine dye Dil into the presumptive somatosensory cortex of E18.5 embryos to label CTAs and trace their projection. Whereas the Dil crystals labeled the highly organized CTAs crossing the PSPB in control brains, the axons tended to traverse the striatum aberrantly in Ulk1/2-DKO brains (Figure 4-7A1, A1', A3, A3'). Despite the disorganization and overfasciculation of the axons, the CTAs still found and targeted their final destination in the dorsal thalamus (dTh), as evidenced by the Dil signal in the dTh of Ulk1/2-DKO brains (Figure 4-7A2, A4). We also immunostained the brains of Ulk1/2Nestin cDKO mice and controls at P21 for vesicular glutamate transporter 2 (VGluT2) and found no obvious differences in the intensity or distribution of this presynaptic marker in the dTh (Figure 4-7B). Cytochrome oxidase (CO) staining highlighted the intact array of barreloids in the thalamus (Figure 4-7B). Therefore, despite the abnormal fasciculation and path taken by the CTAs, we did not detect gross abnormalities in postnatal circuitry in the dTh.

We then embedded Dil crystals into the dTh of E18.5 to trace TCA projections. Similar to the CTAs, the TCAs were disorganized and overfasciculated as they crossed the PSPB (Figure 4-7C). TCAs project to the cortex through the intermediate zone during embryonic development and migrate radially to Layer IV at birth (Lopez-Bendito and Molnar, 2003). The extensive TCA arbors segregate into clusters surrounded by rings of Layer IV neurons’ somas, whose dendrites are oriented to synapse with TCA afferents. Thalamic innervations are essential to remodel cortical neurons postnatally, and this organization gives rise to the barrels in the somatosensory cortex, which relay information from a single whisker (Lopez-Bendito and Molnar, 2003). The decreased levels of cortical staining in E18.5 Ulk1/2-DKO mice using an antibody against the TCA marker calretinin suggested that fewer TCAs reached cortex after crossing the PSPB than in controls (Figure 4-7D). The organization of the TCA projections postnatally in the brains of P7 mice was assessed by immunolabeling for the cortical barrel markers SERT and VGluT2 (Figure 4-7E, F). The somatosensory map in the Ulk1/2Nestin cDKO mice as revealed by CO stain was highly disorganized compared to that in the controls (Figure 4-7G).
**Figure 4-7. ULK1 and ULK2 are required for proper organization of the somatosensory cortex.**

(A) Representative images of fluorescence signal from Dil crystals embedded in the somatosensory cortex reveals the highly disorganized of CTA projections crossing the PSPB in E18.5 *Ulk1/2*-DKO brains. These *Ulk1/2*-deficient axons, despite their disorganization and abnormal fasciculation, ultimately reach their targeted destination and innervate the dorsal thalamus (A2, A4). (B) Normal VGluT2 staining and cytochrome oxidase (CO) staining in the dorsal thalamus in the *Ulk1/2*-DKO brains at P21 suggests proper cortical innervation of that structure. (C) Dil tracing shows the severely disorganized TCA projections in the *Ulk1/2*-DKO brains. (D) Representative image of Calretinin staining reveal a dramatic reduction in the number of TCA projections reaching the cortex in E18.5 *Ulk1/2*-DKO brains. (E-F) Representative images of sections immunostained using antibodies against the cortical barrel markers SERT (pseudocolored red), VGluT2 (pseudocolored green) (E), and the number of barrel units per section was quantified from control (n = 3) and *Ulk1/2*<sup>Nestin</sup> cDKO (n = 3) brains at P7 (F). (G) CO reveal disorganization of the somatosensory barrel cortex in P7 *Ulk1/2*<sup>Nestin</sup> cDKO brains. ** P<0.01. Abbreviations: dTh: dorsal thalamus; A: anterior; M: medial; VPL: ventral posterolateral nucleus; VPM: ventral posteromedial nucleus. Scale bars: 100 µm in A1'-A3', B, C2'-C4', D1'-D2', E, and G; 500 µm in A1-A4, C1-C4, and D1-D2.
TAG-1 Is Mislocalized in Corticothalamic Axons

The defasciculation defects resulting from Ulk1/2-deficiency were similar to those observed in unc-51– or atg1-mutant invertebrates. In unc-51–mutant C. elegans, the axon guidance receptor UNC-5 is mislocalized to the cell soma and fails to mediate repulsion of axons away from the netrin-rich ventral midline.(Ogura and Goshima, 2006) In Drosophila, atg1 regulates axonal localization of Fasciclin II, an IgG-family cell adhesion molecule (CAM), which is important for axonal guidance and fasciculation in the mushroom body. Therefore, we hypothesized that ULK1 and ULK2 may also regulate cellular localization of specific axon guidance receptors in projection neurons. Although most of the axon guidance receptors that we examined, including DCC, ROBO1, PSA-NCAM, and neural cell adhesion molecule (NCAM), were clearly present in the distal segments of projecting CTAs of both Ulk1/2-DKO and littermate controls (Figure 4-8A and data not shown), transient axonal glycoprotein-1 (TAG-1) was notably absent from the distal terminus of the CTAs in Ulk1/2-DKO embryos (Figure 4-8B). The expression of TAG-1 in the dorsal cortex appeared to be unaltered. We also dissected cortex (proximal CTAs) and striatum (distal CTAs) of E18.5 brains, and western blot analyses clearly confirmed the immunostaining results that TAG-1 level is dramatically reduced in the distal CTAs of the DKO animals (Figure 4-8C). These results indicate that ULK1 and ULK2 may regulate the trafficking of specific axon guidance molecules, such as TAG-1 in CTAs, in developing mouse brains similar to ULK/ATG1 homologues in invertebrates.
Figure 4-8. Abnormal axonal fasciculation in the Ulk1/2-deficient animals is associated with mislocalization of TAG-1 in the projection neurons. (A) NCAM distribution was unaltered in the Ulk1/2-DKO brains compared to the controls. (B) The intensity of TAG-1 immunostaining was dramatically decreased in distal CTAs of the Ulk1/2-DKO brain at E18.5 compared to the controls. All of the sections were counterstained with L1. (C) Western blot analyses on the extracts prepared from cortex (proximal CTAs) and striatum (distal CTAs) confirmed normal expression of NCAM but decreased TAG-1 level in the striatum of the Ulk1/2-DKO brain. Scale bars: 200 µm.
CHAPTER 5. DISCUSSION*

The Role of ULK/ATG1 in ER-to-Golgi Trafficking Is Essential for Cellular Homeostasis

Through genetic and biochemical approaches, we revealed an unexpected, evolutionarily conserved role for ULK/ATG1 in ER-to-Golgi trafficking under basal conditions, which is essential for maintaining cellular homeostasis. ULK1 and ULK2 play important roles in selective and nonselective autophagy induced in response to mitochondrial damage, infection, proteotoxic stress, or various forms of metabolic stress (Joo et al., 2011; Lim et al., 2015; Noda and Fujioka, 2015; Wong et al., 2013). Given the presumed importance of the ULKs in both forms of autophagy, the lack of ubiquitin+ and P62+ inclusions or accumulation of abnormal mitochondria in the neurons of Ulk1/2-cdko animals was surprising. These results suggest that unlike ATG5, ATG7, and FIP200, all of which mediate the degradation of ubiquitinated proteins in neurons under basal physiologic conditions (Hara et al., 2006; Komatsu et al., 2006; Liang et al., 2010), ULK1 and ULK2 are not required for “basal” autophagy. Although we cannot exclude the possibility that another ULK/ATG1 homolog (e.g., ULK3) compensates for the lack of ULK1 and ULK2 in CNS neurons, our findings raise the possibility that an ATG13/FIP200 complex regulates the autophagy-mediated turnover of ubiquitinated proteins in an ULK/ATG1-independent manner. Recent studies in isolated cells have shown that ATG13 and FIP200 support starvation-induced autophagy in the absence of ULK1/2, albeit to a lesser extent than in their presence (Alers et al., 2011; Hieke et al., 2015). Such a pathway may support sufficient levels of autophagy in neurons under basal physiologic conditions to prevent the accumulation of ubiquitinated protein deposits.

Although ULK1 and ULK2 are not required for basal autophagy in neurons, our data indicate that a primary function of the ULKs in the absence of metabolic stress is regulating ER-to-Golgi trafficking. The CNS is spared in response to most metabolic stresses, perhaps making it one of a few environments in which the ULKs’ role in ER-to-Golgi trafficking can be readily separated from that in autophagy. ULK/ATG1-mediated COPII transport does not require the presence of other autophagy-related proteins (e.g., ATG13, ATG14, and ATG7), confirming that ULK/ATG1-regulated ER-to-Golgi trafficking is not an indirect consequence of ULKs affecting autophagy. Moreover, the absence of ATG13 does not disrupt the ULK1–SEC16A complex, as it does the canonical ULK/ATG1 complex composed of ULK1, ATG13, FIP200, and ATG101. These results indicate that regulating ER-to-Golgi trafficking under basal physiologic conditions is a noncanonical function (i.e., one that does not require ATG13) of the ULKs.

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The uncoordinated phenotype of *C. elegans unc-51* mutants is associated with defective axon guidance and is not recapitulated in autophagy-deficient mutants, such as *epg-1 (Atg13)* and *epg-9 (Atg101)* mutants (Liang et al., 2012; Tian et al., 2009). RNAi-mediated silencing of other autophagy-related genes, e.g., *bec-1 (Atg6)*, *M7.5 (Atg7)*, *lgg-1 (Atg8)*, or *F41E6.5 (Atg18)*, also does not replicate the neuronal defects observed in *unc-51* mutants (Ogura and Goshima, 2006). Our data suggest that the N-terminal domain of ULK/ATG1 and the central domain of SEC16A, which are the most highly evolutionarily conserved regions of both proteins, mediate the interaction between ULK/ATG1 and SEC16A. Indeed, the role of ULK/ATG1 in ER-to-Golgi trafficking appears to be conserved in *C. elegans*, as *unc-51* mutants show evidence of abnormal COPII assembly and trafficking of MOD-5 (SERT). Therefore, it is tempting to speculate that the defect in ER-to-Golgi trafficking in *unc-51* mutants contributes to their uncoordinated phenotype.

Although the noncanonical function of ULKs in ER-to-Golgi trafficking does not require other autophagy-related proteins, recent studies have implicated the COPII pathway in autophagosome biogenesis (Ge et al., 2013; Ge et al., 2014; Graef et al., 2013; Ishihara et al., 2001; Suzuki et al., 2013). Given that ULK/ATG1 activity is regulated by energy/nutrient availability, we speculate that the ULK/ATG1–SEC16A interaction represents a specific point of convergence among the metabolism, autophagy, and COPII-transport pathways.

**ULK1/2 Are Essential for Axon Guidance in the Developing Mouse Forebrain**

Based on the observation that around 40% *Ulk1/2 cdko* mice die shortly after birth, and *C. elegans UNC-51* and *D. melanogaster Atg1* promote axonal development, we examined the potential embryonic neurodevelopmental defects in the *Ulk1/2 cdko* mice. We uncovered an evolutionarily conserved role of ULK1/2 in mammalian neurodevelopment. Using both germ-line knockout and conditional knockout mouse models, we demonstrated that loss of both *Ulk1* and *Ulk2* expression in the CNS led to pathfinding defects affecting callosal axons, AC, CTAs, TCAs, and mossy fibers via a non-canonical (i.e. autophagy-independent) pathway. The autophagy-inducing activity of ULK/ATG1 depends on its stable interaction with ATG13 and FIP200 (Ganley et al., 2009; Hosokawa et al., 2009; Jung et al., 2009), and can be regulated by AMBRA-mediated ubiquitinylation of ULK1 (Nazio et al., 2013). AMBRA1 is required for neuronal differentiation and survival during early embryogenesis, and FIP200 is required for maintenance of cortical neurons and neuronal stem cells postnatally (Wang et al., 2013). Although a kinase complex composed of ULK1, ATG13 and FIP200 was detected in both P0 and adult cortex, we failed to observe excessive neuronal cell death in *Ulk1/2*-deficient embryos, accumulation of p62+ and ubiquitin+ deposits, or loss of cortical neurons or neuronal stem cells in *Ulk1/2*-deficient adults. Moreover, the neurodevelopmental abnormalities found in *Ulk1/2*-deficient mice were not recapitulated in mice lacking expression of *Fip200* (or *Atg7*) in the CNS. Together, these findings are consistent with the notion that autophagy proceeds in the absence of ULK1/2 in neurons under basal physiologic condition (Joo et al., 2016) and that ULK1/2 regulate axon
guidance via autophagy-independent pathway(s). Similar results were observed in *C. elegans*, where the axon guidance and defasciculation defects observed in unc-51 mutants were not recapitulated in epg-1 (Atg13) and epg-9 (Atg101) mutants (Liang et al., 2012; Tian et al., 2009).

Our studies demonstrated that ULK1 and ULK2, similar to their counterparts in *C. elegans* and *D. melanogaster*, are essential for proper axon guidance and defasciculation in mammals in vivo. *C. elegans* unc-51 mutant neurons display multiple axonal defects, including premature termination, abnormal trajectories, and extra axonal branches (Ogura et al., 1994). UNC-51 binds to UNC-14, a RUN-domain protein that acts as an effector of Rap and Rab GTPases and regulates membrane trafficking (Lai and Garriga, 2004; Ogura et al., 1997). Disruption of either unc-14 or unc-51 expression leads to aberrant localization of UNC-5 to the cell soma rather than axons, and a failure to repulse axons away from the netrin-rich ventral midline (Ogura and Goshima, 2006). UNC-51 also interacts with VAB-8, which controls the cell-surface expression of SAX3/Robo in touch neuron growth cones (Lai and Garriga, 2004; Watari-Goshima et al., 2007). Thus, the defect in axon guidance in *C. elegans* unc-51 mutants is at least in part due to disruption of the UNC-51/ RUN-14/ VAB-8 complex, which regulates the trafficking of various axon guidance molecules to axons (Ogura and Goshima, 2006). Similarly, the axon guidance molecule Fasciclin II, which regulates fasciculation of mushroom body axons in *D. melanogaster*, is mislocalized in atg1 mutants (Mochizuki et al., 2011). In mammals, ULK1 interacts with the synaptic GTPase-activation protein SynGAP, and Syntenin, both of which regulate Rab5-mediated neuronal endocytotic pathway (Tomoda et al., 2004). ULK1 and ULK2 have been implicated in TrkA-receptor trafficking and the endocytosis of nerve growth factor (Zhou et al., 2007). ULK/ATG1 also has a conserved role in regulating assembly of specific ER exit sites (i.e. SEC24C+) and trafficking of associated cargo (Joo et al., 2016). Together, ULK/ATG1-mediated trafficking of axon guidance molecules contributes to the axonal development from *C. elegans* to mammals.

TAG-1 is a neural cell adhesion molecule that is expressed during brain development and has been implicated axon outgrowth, neurite extension and fasciculation, and neuronal migration during development (Karagogeos, 2003). Nevertheless, the precise consequence of the mislocalized TAG-1 in Ulk1/2-deficient animals remains unclear. Among the cargoes whose ER-to-Golgi trafficking is regulated by ULK1/2 is the serotonin transporter, SERT (Joo et al., 2016). SERT is transiently expressed in TCAs, where it helps to regulate serotonin levels (Chen et al., 2015). The appropriate levels of serotonin at synaptic junctions is critical for the normal organization and development of the thalamocortical afferent circuits, and impaired uptake of the serotonin due to genetic or pharmacologic disruption of SERT function in TCAs impairs barrel cortex formation (Chen et al., 2015; Miceli et al., 2013). The disorganization of the somatosensory cortex in Ulk1/2 cdko is similar to (albeit more severe than) that resulting from genetic or pharmacologic disruption of SERT activity in developing mouse brains (van Kleef et al., 2012). Therefore, we speculate that ULK1 and ULK2, similar to their invertebrate homologues, regulate one or more steps in the trafficking of multiple secretory cargoes, TAG-1 and SERT among them, from the ER to the PM, and that
defects in trafficking contribute to the defects in axon guidance in the brains of developing *Ulk1/2*-deficient animals.

**Emerging Noncanonical Functions of ULK/ATG1**

Our studies have uncovered two important noncanonical functions of ULK1/2 in the mouse brain under basal physiological condition. Complementary to our findings, evidence has also accumulated over the years that ULK/ATG1 regulates cellular trafficking under normal physiological conditions, which appears to be autophagy-independent. Moreover, recent data have established that ULK1 and ULK2 are essential for innate immunity, cell death, and glycolysis following starvation (Figure 5-1). As studies on ULK/ATG1 continue, more noncanonical functions of ULK/ATG1 are likely to be revealed in the future.

**ULK1/2 Regulate Metabolism**

Although ULK1 and ULK2 are important integrators relaying metabolic stress signals to downstream autophagy machinery, whether ULK1 and ULK2 modulate other metabolic pathways, such as glucose metabolism, remains unclear. A recent study demonstrated that during AA and serum starvation, ULK1/2 phosphorylate key glycolytic enzymes, including hexokinase (HK), phosphofructokinase 1 (PFK1), enolase 1 (ENO1), and the gluconeogenic enzyme fructose-1,6-bisphosphatase (FBP1) to secure more carbon flux into the pentose phosphate pathway and maintain cellular energy supply (Li et al., 2016). More importantly, knockdown of other key autophagy genes, including *Atg5*, *Atg7*, and *Becn1* had no effect on glucose consumption under the same conditions (Li et al., 2016), indicating that driving glucose metabolism, in addition to initiating autophagy to maintain sufficient intracellular energy level, is a unique feature of the ULK kinases compared to other autophagy proteins.

Intriguingly, another study revealed that *Ulk1* knockdown not only compromised basal and rapamycin-induced autophagy but also reduced fatty acid oxidation and enhanced fatty acid uptake at the steady state in the differentiated adipocytes, and knockdown of *Ulk2* produced the opposite effect, compared to that of *Ulk1* (Ro et al., 2013). Meanwhile, expression levels of insulin receptor, insulin receptor substrate 1, and glucose transporter were all increased in *Ulk1*-depleted adipocytes (Ro et al., 2013). However, the relevant substrate of ULK1 and ULK2 remains to be discovered, the physiological relevance of Ro et al.’s study waits to be confirmed, and the mechanisms underlying the mutually opposite functions of ULK1 and ULK2 need to be revealed. In any case, these preliminary studies opened up a new area where ULK1 and ULK2 emerge as key regulators in various metabolic pathways, such as glucose breakdown and fatty acid oxidation, among others, under different nutrient conditions.
Figure 5-1. The versatile functions of ULK/ATG1 beyond autophagy regulation.
Depending on the cellular physiological status, ULK/ATG1 is capable of targeting different molecules to dictate cell fate. Under normal growth conditions, such as embryonic development, ULK/ATG1 binds to myriad proteins, such as *C. elegans* UNC-14 and VAB-8, *D. melanogaster* UNC-76, and mammalian SynGAP and Syntenin, to mediate vesicle trafficking in neurite outgrowth. Ubiquitilation of UNC-51, triggered by vesicle cargo such as NGF and TrkA, can potentially regulate the function of ULK/ATG1. Upon AA/serum starvation, ULK1/2 phosphorylate key enzymes of glycolysis, including HK, FBP1, PFK1, and ENO1, to ensure sufficient energy supply in addition to autophagy regulation, which is essential for cell survival. Once detected, viral DNA binds to and activates cGAS to synthesize cGAMP. As a second messenger, cGAMP, on the one hand, activates STING to promote IRF3-dependent transcription of IFNs. On the other hand, cGAMP inhibits AMPK by an unknown mechanism and subsequently activates ULK1 to phosphorylate STING. Phosphorylation of STING by ULK1 suppresses its ability to initiate transcription of IFNs, thus providing a negative feedback mechanism to prevent excessive immune response. Sustained stress imposed by ROS leads to translocation of ULK1 from cytosol to nucleus, where ULK1 binds and modulates PARP1 function to promote necrotic cell death. See text for details.
Vesicular Trafficking

Another important aspect of ULK/ATG1’s function lies in vesicular trafficking. *C. elegans* UNC-51 directly binds to UNC-14, a RUN-domain protein that acts as an effector of Rap and Rab GTPases and modulates membrane trafficking (Ogura and Goshima, 2006; Ogura et al., 1994). In *unc-14* and *unc-51* mutants, UNC-5 is aberrantly localized to the cell soma rather than axons (Ogura and Goshima, 2006). The interaction between UNC-51 and the novel protein VAB-8 also implicates UNC-51 in regulating guidance-receptor trafficking, considering that VAB-8 controls the cell-surface expression of SAX3/Robo in touch neuron growth cones (Lai and Garriga, 2004; Watari-Goshima et al., 2007). Although there seems to be UNC-51-dependent phosphorylation of both UNC-14 and VAB-8, the mechanism underlying the regulation of UNC-14 and VAB-8 is not yet clear. The *Drosophila atg1* mutants exhibit abnormalities during embryonic formation of axonal tracts similar to *unc-51* mutant (Mochizuki et al., 2011; Toda et al., 2008). ATG1 physically binds and phosphorylates the kinesin adaptor UNC-76 at Ser143. The phosphorylated UNC-76 has an elevated affinity for Synaptotagmin-1, a major transmembrane protein of synaptic vesicles (Toda et al., 2008). More importantly, the defective axonal transport of synaptic vesicles in *unc-76* mutants is rescued by a phospho-mimetic UNC-76 (S143D), highlighting the essential role of ATG1-mediated phosphorylation of UNC-76 in axonal transport (Toda et al., 2008). In mammals, early studies identified the synaptic GTPase-activation protein (SynGAP) and Syntenin, an endocytic vesicular membrane protein with PDZ domains, as novel binding partners of ULK1; both proteins regulate the Rab5-mediated neuronal endocytotic pathway (Tomoda et al., 2004). Moreover, ULK1 and ULK2 regulate TrkA-receptor trafficking and the endocytosis of nerve growth factor in mouse dorsal root ganglion neurons in culture (Zhou et al., 2007). Mechanistically, NGF can induce the interaction of ULK1 with TrkA receptor complexes through promoting K63-polyubiquitination of ULK1 and binding ULK1 to the scaffolding protein p62 (Zhou et al., 2007). It is unclear whether ULK1 phosphorylates any of the homologous binding partners in mammals, and the physiological relevance of these interactions needs further investigation.

Interferon Response

Pathogenic nucleic acids, such as DNA, RNA, and nucleotides, are detected by host sensing mechanisms (Wu and Chen, 2014). These cell surface sensors regulate activation of several downstream effectors that lead to the transcription of type I IFNs and pro-inflammatory cytokines that regulate innate and adaptive immune responses (Wu and Chen, 2014). IFNs are then secreted and bind cell surface receptors where they initiate transcription of interferon-stimulated genes (ISGs) to mediate appropriate IFN response and anti-viral effect (Ivashkiv and Donlin, 2014). Recent studies have implicated ULK1 and ULK2 in IFN production as well as IFN signaling. Following exposure of cytosolic DNA, the DNA-binding protein cyclic guanosine monophosphate–adenosine monophosphate (cGAMP) synthase (cGAS) triggers synthesis of cGAMP to activate stimulator of interferon genes (STING), which then phosphorylates interferon regulatory factor 3 (IRF3), the transcription factor directly involved in transcription of
IFNs (Wu et al., 2013). Meanwhile, cGAMP is required for dephosphorylation of AMPK at Thr172 and subsequent dephosphorylation of Ser555 of murine ULK1. This also leads to activation of ULK1, which directly phosphorylates STING at Ser366 to block its kinase activity towards IRF3 and subsequently inhibits transcription of related immune genes (Konno et al., 2013). Thus, ULK1 fine tunes the immune response to pathogenic nucleic acids through a negative-feedback mechanism to prevent STING-dependent inflammatory disorders. Interestingly, another study reported that upon IFN stimulation, AKT-dependent phosphorylation of ULK1 at Ser757 is required for phosphorylation of p38 MAPK and eventually proper transcription of ISGs in multiple cell lines (Saleiro et al., 2015). More importantly, abolishing ATG5 had no impact on the transcription of ISGs, suggesting that ULK1 and ULK2 mediate IFN response, probably via a noncanonical, autophagy-independent mechanism (Saleiro et al., 2015). Notably, patients with myeloproliferative neoplasms (MPNs) were found to have elevated expression levels of ULK1 compared to controls, and suppression of ULK1 expression blocked the anti-neoplastic functions of type I IFNs against malignant erythroid precursors derived from the patients (Saleiro et al., 2015). These results suggest that increased ULK1 expression could correlate with favorable responses to type I IFN therapy in MPNs and other types of IFN-responsive cancers, but this remains to be explored further.

**Necrotic Cell Death**

Generally perceived as a pro-survival mechanism, autophagy can also be destructive and lead to cell death under certain conditions (Maiuri et al., 2007). How individual autophagy machinery, including ULK1 and ULK2, contributes to cell death remains largely unknown. A previous study suggested that treating osteosarcoma cells with sub-lethal dose of camptothecin leads to p53-mediated upregulation of ULK1 at transcription level, which induced sustained autophagy and subsequently cell death (Gao et al., 2011). Our recent study for the first time has demonstrated that ULK1 translocates to the nucleus and sensitizes cells to necrotic cell death in response to hydrogen peroxide treatment (Joshi et al., 2016). Mechanistically, nuclear ULK1 interacts with poly (ADP-ribose) polymerase 1 (PARP1) and enhances PARP1 activity, which accelerates ATP depletion and cell death in an autophagy-independent manner (Joshi et al., 2016). Whether PARP is a novel substrate of ULK1 or there is an unknown substrate of ULK1 bridging ULK1 and PARP, and the signaling events leading to the translocation of ULK1 from cytosol to nucleus need further investigation. Interestingly, ULK2 has also been shown to localize to the nucleus with the aid of an importer receptor-Kapβ2, even under basal conditions in HEK293 cells. The functional significance of the nuclear ULK2 has not yet been elucidated (Shin et al., 2015).

**ULKs and Neurological Diseases**

For neurons, eliminating dysfunctional organelles and cellular waste over a lifetime without harnessing cell division to buffer the waste burdens is particularly challenging (Nixon, 2013). Hence, neurons rely heavily on autophagy for survival, and
dysregulated autophagy is often linked to neurodegenerative diseases (Nixon, 2013). As expected, conditionally knocking out Atg5, Atg7, or Fip200 in the CNS causes widespread neuronal loss, associated with accumulation of p62- and ubiquitin-positive inclusion bodies, a pathological feature shared by numerous human neurological diseases (Hara et al., 2006; Komatsu et al., 2006; Liang et al., 2010; Nixon, 2013). Our study suggests that, unlike other canonical autophagy proteins, ULK1 and ULK2 are not required for “basal” autophagy in the neurons (Joo et al., 2016). However, the existence of stable ULK1-ATG13-FIP200 in the cortex at various ages suggests that ULK1 and/or ULK2 might induce autophagy in response to certain stresses, such as aberrant accumulation of protein aggregates. This hypothesis is supported by a recent study, which demonstrated that in response to proteasome inhibition or expression of polyQ-Htt, ULK1 phosphorylates the autophagy receptor protein p62 thus inducing autophagy-mediated degradation of excessive protein aggregates (Lim et al., 2015). Evidence that dysfunction of ULKs contributes to human neurological diseases is sparse. A recent study revealed that both ULK1 and ULK2, together with other autophagy proteins, accumulate in the mature Lewy bodies in the brains of patients with Parkinson’s disease or dementia with Lewy bodies (Miki et al., 2016). This likely represents a cellular protective mechanism in which ULK1 and ULK2 cooperate to degrade the excessive aggregates of disease proteins through autophagy, rendering ULK1 and ULK2 attractive targets for treating certain neurodegenerative diseases. Alternatively, ULK1- and ULK2-mediated cellular trafficking may affect the developing brain. Brain wiring abnormalities are believed to contribute significantly to a spectrum of psychological diseases, including the ones with a neurodevelopmental origin (autism, for example). No mutations in Ulk1 or Ulk2 have so far been reported in human patients suffering from neurological disorders. However, similar to our observations in ULK1/2 deficient mice, where the survival rate is lowered, Ulk1 or Ulk2 mutations may cause early mortality in humans too.

**Future Directions**

Recent work from several labs, including our own, has provided significant insights on the cellular functions and the regulation of ULKs. The physiological significance of ULKs in vivo, however, still remains largely unclear, thus calling for more detailed analyses of Ulk-deficient mouse models. As it is possible that the in vivo function of ULKs might be tissue-dependent, it is essential that the phenotypic analyses on the Ulk-deficient animals be performed under basal conditions with minimal bias. It will also be crucial to ascertain whether the phenotype is autophagy-dependent or -independent by comparing Ulk-deficient animals with mice depleted of other autophagy genes such as Atg7, and Fip200. Despite the well-established significance of ULKs in metabolic stress-induced autophagy in several cell lines, investigations on in vivo models may yield different results, leading to different conclusions. Thus, it is prudent to use multiple means and modes of stress (such as starvation and ischemia-reperfusion) to induce autophagy in Ulk-deficient mice and their effects should be studied.

Given the plethora of cellular processes that ULKs may be involved in, it is reasonable to predict that there are more substrates of these kinases than those...
summarized in Chapter 1. Thus, using different approaches to identify novel substrates of ULKs will be of continuous interest. Not only do the phosphorylation sites of these potential substrates need to be located with traditional biochemical assays, but also the possible functional significance of these interactions and phosphorylations needs to be explored. In addition, identifying novel upstream regulators of ULKs will be critical to understanding how ULKs integrate the signals and dictate the cellular response under different conditions. Another puzzle to solve is how the function of ULKs is coordinated in different environments. It is possible that under stress conditions, ULKs are hijacked into the autophagy pathway by having their normal function compromised. Alternatively, the noncanonical function of ULK/ATG1 may also contribute directly to autophagy. This notion is validated by recent reports that the ER membrane is a source for autophagosome formation, and that the COPII pathway is involved in autophagosome biogenesis (Ge et al., 2013; Ge et al., 2014; Graef et al., 2013; Ishihara et al., 2001; Suzuki et al., 2013). Thus, it is tempting to speculate that ULK/ATG1-regulated ER-to-Golgi trafficking may act as a membrane source during autophagy induction. Nevertheless, identifying the upstream molecules that dictate the specific function of ULKs under different conditions remains an important but also challenging task.

Finally, disruption of ULK function may contribute to the pathogenesis of multiple human diseases, including cancer and neurodegeneration, which doesn’t necessarily involve the autophagy-dependent mechanism. In light of our novel description of the noncanonical function of ULKs, it becomes necessary to revisit the previously established idea that ULKs function solely in an autophagy-dependent fashion in cancer cells. As the noncanonical function of ULKs could potentially also contribute to tumor cell growth or demise, depending on the specific context, it is therefore critical to separate the different function of ULKs in any specific situation. Now that the knockout mouse models of ULK1 and ULK2 are available, crossing the Ulk1 and Ulk2 KO mice with mouse models of cancer and neurological diseases and access the role of ULK1 and ULK in the disease progression will be an attractive direction to explore. This will not only facilitate a clearer understanding of the basic mechanisms of pathogenesis, but also will help develop new therapeutic strategies to combat many devastating diseases.
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