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The Cul3 Ubiquitin Ligase: an Essential Regulator of Diverse Cellular Processes

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The Cul3 Ubiquitin Ligase: An Essential Regulator of Diverse Cellular Processes

by

Brittney Marie Davidge

A dissertation submitted in partial fulfillment of the requirements for the degree of

Doctor of Philosophy
in
Biology

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ABSTRACT:

Cul3 forms E3 ubiquitin ligase complexes that regulate a variety of cellular processes. This dissertation describes Cul3’s role in several of these pathways and provides new mechanistic details regarding the role of Cul3 in eukaryotic cells. Cyclin E is an example of a protein that is regulated in a Cul3-dependent manner. Cyclin E is a cell cycle regulator that controls the beginning of DNA replication in mammalian cells. Increased levels of cyclin E are found in some cancers, in addition, proteolytic removal of the cyclin E N-terminus occurs in some cancers and is associated with tumorigenesis. Cyclin E levels are tightly regulated and controlled in part through ubiquitin-mediated degradation initiated by one of two E3 ligase complexes, Cul1 and Cul3. Cul1 mediated degradation of cyclin E is triggered by cyclin E phosphorylation, however the mechanism Cul3 uses to ubiquitinate cyclin E is poorly understood. In order to gain a better understanding of how Cul3 mediates cyclin E destruction we identified the degron on cyclin E that is important in Cul3 dependent degradation. In addition, we show this degron is lacking in LMW cyclin E (found in abundance in breast cancer), providing a novel mechanism for how these cyclin E modifications result in increased cyclin E levels by avoiding the Cul3 degradation pathway.
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LIST OF ABBREVIATIONS USED

APC: Anaphase Promoting Complex

AQP2: Aquaporin 2

BACK: BTB and C-terminal Kelch

BCR: BTB, Cul3, Rbx1

BTB: Bric-a-brac, Tramtrack, Broad-complex

Cdk: Cyclin-dependent kinase

CHX: Cycloheximide

CKI: Cyclin-dependent kinase inhibitor

Cul1: Cullin 1

Cul2: Cullin 2

Cul3: Cullin 3

Cul4A: Cullin 4A

Cul4B: Cullin 4B

Cul5: Cullin 5

Cul7: Cullin 7

DCT: Distal convoluted tubule

ECM: Extracellular matrix

FHHt: Familial Hyperkalemic Hypertension: Also called PHAII or Gordon’s syndrome.

HECT: Homologous to E6-AP Carboxyl Terminus: a class of E3 ubiquitin ligases
**HIFα**: Hypoxia-inducible factor alpha: HIFα has been suggested to be a substrate of RhoBTB3.

**KO**: Knockout

**LMW**: Low molecular weight

**LRR**: Leucine-rich repeat

**LRR3 (SALM1)**: LRR-domain protein 3, also called SALM1 (synaptic adhesion like molecule 1)

**LRR5 (FMOD)**: LRR-domain protein 5, also called FMOD (fibromodulin)

**MEFs**: Mouse embryonic fibroblasts

**ORC**: Origin recognition complex

**PHAII**: Pseudohypoaldosteronism type II: Also called FHHt.

**PKA**: Protein kinase A

**Rb**: Retinoblastoma protein

**ROS**: Reactive oxygen species: Oxidative stress

**SCF**: Skp1, Cul1, F-box

**TG2**: Transglutaminase 2 (tissue transglutaminase)

**WNK1**: With no lysine (K) 1: A kinase involved in NCC regulation.

**WNK4**: With no lysine (K) 4: A kinase involved in NCC regulation.

**WT**: Wild-type
GLOSSARY

**Apical membrane:** The apical membrane is found in the nephron where it contacts the urine. AQP2 is localized to the apical membrane.

**AQP2:** Aquaporin 2 is a water channel protein that resides in the apical surface of the collecting duct. AQP2 helps to maintain blood pressure via the re-absorption of water.

**ATP1B1:** A subunit of the Na+/K+ ATPase. This subunit was identified as a possible binding partner of Klhl3 (Chapter 3).

**Basolateral membrane:** The basolateral membrane contacts the interstitial fluid in the nephron.

**BTB:** A domain found in proteins that function as Cul3 substrate adaptors. The BTB domain binds to Cul3.

**Cdk:** Cyclin-dependent kinases are responsible for cell cycle progression. A Cdk is active when it is bound to a cyclin.

**Cdk2:** The Cdk binding partner of cyclin E. Cyclin/Cdk2 complexes regulate the G₁/S transition and release from quiescence.

**CKI:** Cyclin-dependent kinase inhibitor. There are two families of CKIs: INK4, which inhibit Cdk4/6, and the CIP/KIP family, which includes p21 and p27.

**Cre:** Cre recombinase is an enzyme that can be used in a conditional knockout mouse model to recombine the DNA that is surrounded by two LoxP sites (floxed).
Ctb73: A putative substrate adaptor for Cul3, Ctb73 contains an N-terminal BTB domain, a central BACK domain, and a C-terminal PHR domain.

Cul1: One of two E3 ligases responsible for cyclin E ubiquitination. Cul1 uses substrate adaptors with two subunits: The Skp1 linker protein and an F-box protein.

Cul3 Δ403-459: A Cul3 mutation that results in the skipping of exon 9 of the Cul3 protein. This mutation causes FHHt in humans.

Cul3: An E3 ubiquitin ligase that targets many substrate, including cyclin E, for ubiquitination. Cul3 uses BTB-domain containing proteins as substrate adaptors.

Cul3K712R: Cul3K712R is inactive because it cannot be neddylated.

Cul3Δ51-67: This Cul3 mutant lacks the region near its N-terminus that is responsible for binding BTB proteins (substrate adaptors).

Cul5: Cul5 is an E3 ligase that uses B/C box or VHL proteins and substrate adaptors.

Cyclin E: Cyclin E controls the G1/S transition by binding and activating Cdk2. Cyclin E/Cdk2 complexes phosphorylate themselves in addition to many other substrates including the Rb protein. Eukaryotes contain two cyclin E proteins, cyclin E1 and cyclin E2, which are products of two different genes.

Cyclin homology domain: This domain is responsible for cyclin E’s ability to bind Cdk2 and is found in the central part of cyclin E’s structure.

Cyclins: A class of proteins responsible for binding and activating Cdks, resulting in cell cycle progression.
Dynamitin/p50: A subunit of the dynactin complex. This complex is responsible for endosomal transport. P50 links the microtubule binding and cargo binding portions of the complex. P50 was identified as a Klhl3 interacting protein and possible substrate (Chapter 3).

E1 ubiquitin activating enzyme: E1 enzymes are responsible for activating the C-terminal Gly-Gly motif on ubiquitin and preparing it for attachment to a substrate. Activation of ubiquitin by an E1 occurs via an ATP-dependent reaction.

E2 ubiquitin conjugating enzyme: E2 enzymes work with E3 ubiquitin ligases in order to transfer ubiquitin to substrates.

E2F: A transcription factor involved in cell cycle regulation. E2F is sequestered by the Rb protein, which releases E2F upon sufficient phosphorylation by cyclins and Cdks. Once E2F is free of Rb, it is able to activate transcription, resulting in entry into S-phase.

E3 ubiquitin ligase: E3 ligases provide substrate specificity to the ubiquitin system. Cullin E3s work with an E2 to facilitate ubiquitin attachment. HECT E3 ligases and cullin ligases represent two classes of E3s.

EEA1: A marker of early endosomes.

ENaC: A sodium channel located in the nephron.

F-box: Cul1 utilizes substrate adaptors, such as Fbxw7, that contain F-box domains to recognize substrates.

FAZF: A BTB protein that also contains a zinc-finger domain. FAZF binds LRR5 (FMOD) (Chapter 4).
**Fbxw7**: An F-box protein that functions as a substrate adaptor for Cul1. Fbxw7 is involved in Cul1-mediated ubiquitination of cyclin E.

**FHHt**: Familial Hyperkalemic Hypertension: Also called PHAII or Gordon's syndrome. Mutations in Cul3 and Klhl3 cause this disease.

**Floxed**: “Flanked by LoxP”: This is a DNA construct that uses LoxP sites to make conditionally –expressing mouse alleles. Cre recombinase can be expressed in mice that carry a floxed allele, resulting in the deletion of the selected DNA. The Cre LoxP system is useful for studying essential genes, as it allows for tissue specific deletion.

**G₀**: G₀ is another name for the quiescent stage of the cell cycle. A cell arrested in quiescence is not preparing to divide, but can re-enter the proliferative cell cycle if given the appropriate mitogenic signals.

**G₁**: This is the first growth phase that takes place during interphase of the cell cycle. G₁ occurs prior to S-phase.

**G₂**: G₂ is the growth phase following DNA synthesis but prior to mitosis.

**Gli**: A family of mammalian transcription factors that are involved in the hedgehog signaling pathway. The Gli proteins are homologues of Ci in *Drosophila*. Gli proteins, as well as Ci, are ubiquitinated by Cul3.

**GSK3**: A kinase that phosphorylates cyclin E.

**Interphase**: Interphase consists of the G₀, G₁, S, and G₂ phases of the cell cycle.

**KCTD13**: KCTD13 is the BTB substrate adaptor that targets RhoA for Cul3-mediated degradation.
**Keap1**: A prominent member of the BTB-BACK-Kelch family of proteins, Keap1 targets the transcription factor Nrf2 for degradation.

**Kelch domain**: A common substrate-recognition domain that is found in many BTB-domain containing proteins. Some notable members of this family are Keap1, which regulates Nrf2, and Klhl3, which is involved in blood pressure regulation.

**Klhl17 (actinfilin)**: A BTB-BACK-Kelch protein that is thought to be a Cul3 substrate adaptor.

**Klhl2 (Mayven)**: A BTB-BACK-Kelch protein and Cul3 substrate adaptor, Klhl2 is closely related to Klhl3.

**Klhl24**: Klhl24 is a BTB-BACK-Kelch protein and Cul3 substrate adaptor that ubiquitinates keratin 14.

**Klhl3**: A BTB-BACK-Kelch protein that has been shown to cause FHHt in humans when mutated.

**Klhl3R528H**: A dominant mutation in Klhl3 that has been associated with FHHt.

**LAMP-1**: A marker that can be used to identity late endosomes.

**MATH domain**: A secondary domain that is found in the Cul3 substrate adaptor SPOP.

**MEI-1**: A katanin protein involved in microtubule dynamics, MEI-1 is ubiquitinated by Cul3 and MEL-26.

**MEL-26**: A BTB protein that works with Cul3 to ubiquitinate MEI-1.

**MG132**: A proteasome inhibitor that when added to cultured cells, results in increased stability of some ubiquitin-proteasome system substrates.
Mitosis: The final portion of the proliferative cell cycle when a cell completes cell division, resulting in two daughter cells.

MSI2: A substrate of the Cul3RhoBT2 ligase that is associated with breast cancer.

MUF1 (LRRC41): An LRR-domain containing protein that also contains a B/C/ box, making it a possible Cul5 substrate adaptor. MUF1 can bind Cdk2 (Chapter 4).

Na+/K+ ATPase: This transporter is found in a variety of tissues and is responsible for creating an ion gradient in the cells of the nephron. The beta subunit of this transporter was identified as a potential Klhl3-interacting protein (Chapter 3).

NCC transporter: NCC is a sodium chloride transporter found in the DCT of the nephron. Improper regulation of NCC causes FHHt hypertension.

Nedd8: Nedd8 is a ubiquitin-like molecule involved in Cul3 function. Cul3 is modified by Nedd8 on K712.

Nrf2: Nrf2 is a transcription factor involved in the stress-response. Nrf2 is a substrate of the Keap1 BTB-Kelch protein and Cul3.

OSR1: OSR1 is a kinase that activates NCC via phosphorylation.

PEST: A short sequence that was identified in proteins with high turnover. Cyclin E has a PEST sequence located near its C-terminus.

PHR domain: A secondary domain that is similar to a kelch domain. Several BTB-proteins also contain PHR domains. Kelch and PHR domains are involved in Cul3 substrate recognition.

PLZF: A zinc-finger domain containing BTB protein. PLZF is closely related to FAZF.
**Rb:** The retinoblastoma protein plays a crucial role in cell cycle progression, as it is responsible for binding and sequestering the E2F transcription factor, which is necessary for S-phase to begin. Sufficient phosphorylation of Rb by cyclin/Cdk pairs results in the release of E2F, allowing for transcription of S-phase associated genes to begin.

**Rbx1:** The “ring-finger” protein, which interacts with cullin ubiquitin E3 ligases near their C-termini. Rbx1 is believed to play a role in E2 binding.

**RCBTB1:** RCBTB1 is a BTB protein that has been associated with the eye disease retinitis pigmentosa.

**RhoA:** RhoA is a small GTPase that regulates many processes, including the actin cytoskeleton. RhoA is ubiquitinated and degraded in a Cul3\textsuperscript{KCTD13} dependent manner.

**RhoBTB2:** A member of the RhoBTB protein family, RhoBTB2 targets MSI2 for degradation.

**RhoBTB3:** A member of the RhoBTB protein family that binds cyclin E. Others have suggested that RhoBTB3 and other members of the RhoBTB family associate with MUF1.

**S-phase:** The cell cycle stage during which a cell replicates its DNA.

**Skp1:** Skp1 forms part of Cul1-based complexes as it is responsible for linking Cul1 with an F-box protein.

**SPAK:** SPAK is a kinase that regulates NCC. See OSR1.

**SPOP:** Also called Ctb75, SPOP is a Cul3 substrate adaptor (BTB protein).
**Substrate adaptor:** Substrate adaptors are proteins that bind to cullin-based E3 ligases in order to recognize specific substrates for ubiquitination. Each cullin ligase can associate with many different substrate adaptors, allowing each cullin to have many substrates.

**UbE2E1:** UbE2E1 is an E2 ubiquitin conjugating enzyme, which forms degradative (K48) ubiquitin linkages. UbE2E1 binds RhoBTB3 (Chapter 2).

**Ubiquitin:** Ubiquitin is a small protein that is attached to target proteins, often resulting in their degradation.

**WNK1:** With no lysine (K) 1: A kinase involved in NCC regulation. WNK1 can bind and be ubiquitinated by Klhl3.

**WNK4:** With no lysine (K) 4: A kinase involved in NCC regulation. WNK4 interacts with and can be ubiquitinated by Klhl3.

**Zinc Finger domain:** A DNA-binding domain that is present in some BTB proteins, for example PLZF and FAZF.
CHAPTER 1: Introduction and Background
THE CUL3 UBQUITIN LIGASE

Post-translational modifications of proteins such as phosphorylation, methylation, acetylation, and ubiquitination are essential for a wide variety of cell signaling events. The ubiquitin system is one crucial method of cellular signaling in eukaryotic cells in which a small protein called ubiquitin (Figure 1.1) is covalently attached to lysine residues on target proteins, also called substrates (Metzger et al. 2012). Substrates can be either monoubiquitinated (one molecule of ubiquitin is attached) or polyubiquitinated with a ubiquitin chain consisting of multiple ubiquitin molecules (Hicke 2001). Histones are often modified post-translationally by monoubiquitination (Cole et al. 2015). Ubiquitin chains can form different conformations (Grice and Nathan 2016), which are determined by the specific lysine residues on the ubiquitin molecules that comprise the chain (Figure 1.2). Ubiquitinated proteins can be differentiated by the cell depending upon the type and length of ubiquitin chain that is attached. For example, K48-linked ubiquitin chains signal the ubiquitinated protein to be degraded by the proteasome whereas K63-linked chains are involved in different processes (Lim and Lim 2011). The attachment of ubiquitin signals a target protein to undergo a variety of cellular processes including degradation by the 26S proteasome (Grice and Nathan 2016). The ubiquitin system relies upon a complex set of biochemical mechanisms allowing for proper substrate identification and ubiquitin attachment. Ubiquitin signaling is essential for the survival and proper function of eukaryotic cells (Finley et al. 1987).
The transfer of ubiquitin to a substrate requires three enzymes (Figure 1.3). First, an E1, also called an ubiquitin activating enzyme, attaches the C-terminal Gly residue of ubiquitin to its active site in an ATP-dependent reaction (Haas et al. 1982). Next, The E1 transfers the ubiquitin to the active site of an E2 ubiquitin-conjugating enzyme. Lastly, the E2 will work with an E3 ubiquitin ligase to form an isopeptide bond attaching the ubiquitin (via the C-terminal Gly residue) to a lysine residue on the target protein (Haas and Rose 1982; Pickart and Rose 1985). Ubiquitin ligases are a diverse class of molecules, which is important because they are responsible for providing substrate specificity to this system (Gonen et al. 1996). Thus, ubiquitin ligases ensure that the appropriate target protein is ubiquitinated at the proper time.
Figure 1.1: Ubiquitin is a small protein that is attached to substrates. Ubiquitin is a small protein, the C-terminus of which is attached to substrates and other ubiquitin molecules via lysine residues. Different conformations of chains are formed depending upon which ubiquitin lysines are involved; for example K48-linked chains target substrates for proteasomal degradation.
**Figure 1.2:** *Examples of ubiquitin chains.* K48-linked chains have a globular shape and are associated with degradation (Left). K63-linked chains have a linear structure and are associated with non-degradative processes (Center). A substrate can also be monoubiquitinated, which is necessary for regulation of histone proteins (Right).
Figure 1.3: *Ubiquitin attachment is a three-step process.* Attachment of ubiquitin to a substrate requires three events: First, an E1 activating enzyme prepares the ubiquitin molecule for conjugation. Next, the E1 transfers the ubiquitin to the active site Cysteine on the E2 ubiquitin-conjugating enzyme. Lastly, the E2 will work with an E3 ubiquitin ligase, which provides substrate specificity, to attach the ubiquitin to the substrate.

There are two main classes of E3 ubiquitin ligases. The first class is called the HECT (homologous to E6-AP carboxyl terminus) ubiquitin ligases, which can directly catalyze the transfer of an activated ubiquitin molecule to a substrate (Bernassola et al. 2008). The second class of E3 ligases is called the cullin-RING (really interesting new gene) E3 ligases (Petroski and Deshaies 2005). Unlike the HECT E3s, these cullin scaffolds do not themselves catalyze ubiquitin transfer, but
instead function to position the substrate in close proximity to the E2, allowing for the attachment of the ubiquitin molecule (Petroski and Deshaies 2005).

Humans possess eight cullin proteins (Cul1, Cul2, Cul3, Cul4A, Cul4B, Cul5, Cul7, and the APC) each of which forms its own set of E3 ligases complexes (Petroski and Deshaies 2005). The cullin-based ligases are divided into several subtypes including Cul1-based (SCF), Cul2-based, Cul3-based etc. The complexes formed by these different ligases have many structural similarities, but each type of ligase is unique and requires a different set of substrate recognition proteins, for example Cul1 uses Skp1 and an F-box protein to recognize substrates, and Cul3 uses BTB-domain containing proteins to bind substrates (Geyer et al. 2003; Petroski and Deshaies 2005). The requirement of a second protein, or substrate adaptor, to bind a substrate, allows for each cullin to have a diverse set of substrates and affect a wide variety of cellular processes. Cul1 and Cul3 have both been shown to degrade the cell cycle regulator cyclin E and form SCF (Skp1-Cul1-F-box) and BCR (BTB, Cul3, Rbx1) complexes respectively (Clurman et al. 1996; Singer et al. 1999; Petroski and Deshaies 2005). Cul1-based complexes have been extensively studied and the structure of these complexes is well known (Figure 1.4). The structure of Cul3-based complexes is not as well understood, even though they are crucial for cellular functioning and Cul3 is an essential gene in mice (Singer et al. 1999).
**Figure 1.4:** *A comparison between Cul1 and Cul3-based complexes.* Cul1 (SCF, top, purple) uses a two-subunit substrate adaptor consisting of Skp1 and an F-box protein whereas Cul3 (BCR, bottom, red) relies upon one-subunit substrate adaptors called BTB proteins, which contain both a Cul3 binding domain and a substrate interaction domain. Both Cul1 and Cul3 bind the ring-finger protein, Rbx1, and associate with E2 enzymes (green) via the C-terminal domain of the cullin.

Both Cul1 and Cul3 possess an N-terminal domain which interacts with substrate adaptors, a central cullin homology domain, and a C-terminal region that is modified by the ubiquitin-like protein Nedd8, interacts with the ring finger protein Rbx1, and interacts with the E2 enzyme (Hori et al. 1999; Zheng et al. 2002; Wimuttisuk and Singer 2007). Current understanding of cullin architecture posits that the E2 binds to Rbx1, which is located near the C-terminus of the cullin (Kleiger et al. 2009; Metzger et al. 2014). For example, it has been shown that the acidic tail on the E2 enzyme Cdc34 interacts with a basic region near the Rbx1 binding site on Cul1 (Kleiger et al. 2009). SCF modification of Cul1 by Nedd8 increases SCF activity.
and may be involved in recruiting the E2 to the complex, demonstrating the importance of Nedd8 modification for cullin functioning (Kawakami et al. 2001).

In order to recognize substrates, Cul1 relies upon substrate adaptors comprised of Skp1 and an F-box protein. Skp1 helps to associate the F-box protein with the Cul1 complex, whereas the F-box protein is responsible for recognizing and recruiting phosphorylated substrates to the complex (Skowyra et al. 1997; Zheng et al. 2002).

Cul3 relies upon a class of proteins containing BTB (Bric-a-brac, Tram track, Broad Complex) domains to recognize substrates (Xu et al. 2003; Pintard et al. 2004). These proteins contain one or more BTB domains that interact with Cul3, as well as a secondary domain such as a Kelch domain or MATH domain which is responsible for substrate binding (Stogios et al. 2005). BTB proteins work with Cul3 to regulate a variety of cellular processes. The BTB family of proteins has many members including MEL-26 which ubiquitinates the katanin protein MEI-1, RhoBTB3 which is believed to ubiquitinate both cyclin E and HIFα, Klhl3 which is involved in blood pressure regulation by ubiquitinating WNK4, and Klhdc5 (Ctb9) which ubiquitinates the microtubule-severing protein p60/katanin (Pintard et al. 2003; Cummings et al. 2009; Johnson et al. 2009; Lu and Pfeffer 2013; Shibata et al. 2013; Wakabayashi et al. 2013; Zhang et al. 2015). Coordinated ubiquitination of a substrate by more than one ligase is known to occur. For example, the cell-cycle regulator Cyclin E provides us with an example of a substrate that is degraded by two different cullin-based E3 ligases, Cul1 and Cul3 (Clurman et al. 1996; Singer et
al. 1999). The ubiquitination of the Gli proteins in the hedgehog-signaling pathway is also known to occur via both Cul1 and Cul3-dependent mechanisms (Jiang 2006).

**CELL CYCLE REGULATION: Basic mechanisms controlling proliferation**

The cell division cycle is one crucial process in which the ubiquitin system has long been known to play an essential role. Interphase of the eukaryotic cell cycle is divided into three stages. The first stage is a growth phase called G₁ (Ohtsubo and Roberts 1993). G₁ is followed by S phase during which the DNA is replicated, which in turn is followed by G₂. The proliferative cell cycle is completed with “M phase”, the phase during which mitosis occurs (Pardee 1974). Eukaryotes have evolved complex mechanisms to ensure that cells replicate their DNA and divide only when appropriate. Mechanisms to ensure the proper replication of DNA are in place throughout interphase of the cell cycle, which helps to ensure that the genome is copied only once and only during S phase (Nishitani and Lygerou 2002). During S-phase, DNA replication begins from pre-determined locations on the DNA, known as origins, which must be properly labeled for replication to occur (Ford and Chevalier 1995). Origins are labeled either late in M-phase or early in G1 (Ford and Chevalier 1995). The labeling of origins during this brief temporal window is one way to decrease the chance that re-replication will occur, as it prevents the labeling of more origins later on during the cell cycle (Ford and Chevalier 1995). Following the labeling of an origin during G₁, the proteins Cdt1 and Cdc6 will associate with the ORC (origin recognition complex). The next step in the replication process is called licensing, which can only occur during G₁ (Nishitani and Lygerou 2002).
During this step, two proteins, Cdt1 and Cdc6, load the replication machinery onto the origin in preparation for DNA replication (Nishitani and Lygerou 2002).

The cell cycle also has a special state referred to as ‘G₀’ or ‘quiescence’, during which the cell is not preparing to divide. Unlike senescent cells, which can no longer divide, quiescent cells possess the ability to exit the G₀ state and reenter the cell cycle if given the proper molecular signals (Siddiqi and Sussman 2014). The quiescent state of the cell cycle is imperative to the normal functioning of eukaryotic organisms. At any given time, a large portion of the cells in a multicellular organism are arrested in this quiescent state. Quiescence is actively regulated by the cell and is necessary to prevent excessive proliferation and preserve the integrity of the cell’s DNA (Cheung and Rando 2013). Abnormal release from quiescence can result in unwanted proliferation and can be harmful to the organism. Intricate processes regulate the cell cycle during all its stages and are governed by cyclin/Cdk (cyclin dependent kinase) protein complexes. These complexes drive the cell cycle forward via the phosphorylation of a wide variety of substrates.

CELL CYCLE REGULATION: Cyclins and Cdk

Cyclins are essential for proper cell cycle control. Mammalian cells have several different cyclins, and each one is active at a specific time point during the cell cycle (Figure 1.5). Therefore, the level of any given cyclin does not remain constant, but instead rises and falls throughout the different stages of the cell cycle. Proper temporal regulation of cyclin activity throughout the cell cycle is necessary for proper functioning of the cell and the health of the organism as a whole.
Figure 1.5: *Cyclins and Cdk5 regulate cell-cycle transitions*. Cyclin D and its partner Cdk4/6 are responsible for internalizing an external mitogenic signal. An external stimulus causes cyclin D/Cdk 4/6 to phosphorylate target proteins, such as Rb, that result in cell cycle progression and increase cyclin E levels. Cyclin E/Cdk2 complexes phosphorylate substrates during the G1/S transition resulting in the start of S-phase. A cell has passed the restriction point once it no longer requires external signals for cell cycle progression to occur, and cyclins D and E are involved in this process. Cyclin A/Cdk2 controls progression through S-phase and cyclins A and B bind Cdk1, controlling progression through G2 and mitosis.

Cyclins bind to and activate enzymes known as Cyclin Dependent Kinases (Cdns) (Koff et al. 1991). When bound to a Cdk, the Cyclin/Cdk complex phosphorylates target proteins that are responsible for cell cycle progression (Koff et al. 1992). Cyclin E is a mammalian cyclin that binds to Cdk2. The Cyclin E/Cdk2 complex is responsible for the phosphorylation of target proteins during the G1/S
transition of the cell cycle (Figure 1.5). The cyclin E gene is one of many cell cycle related genes that is regulated by the E2F transcription factor (Swiss and Casaccia 2010). E2F binds to the Rb (retinoblastoma) pocket protein, which prevents it from activating transcription (Swiss and Casaccia 2010). Sufficient phosphorylation of Rb by cyclin D and Cdk4/6 upon mitogen stimulation during G1 of the cell cycle results in release of E2F by Rb and increased expression of E2F controlled genes, including cyclin E (Resnitzky et al. 1994; Sherr 1994; Ekholm and Reed 2000). Cyclin E then binds and activates Cdk2, resulting in the phosphorylation of Rb as well as other substrates (Koff et al. 1991; Ohtsubo and Roberts 1993). Cyclin E/Cdk2 substrates include Rb, Cdh1, cyclin E/Cdk2 autophosphorylation, and others (Won and Reed 1996; Abbas et al. 2007; Keck et al. 2007). One essential substrate of cyclin E/Cdk2 is Cdc6, which when stabilized upon phosphorylation by cyclin E/Cdk2, allows for the formation of the pre-replication complex prior to the start of S-phase (Ayad 2005; Mailand and Diffley 2005).

G1 contains a point known as the “Restriction point” at which time the cell has committed to duplicating its DNA and will continue on into S-phase even if the external mitogenic stimuli are removed (Pardee 1974; Blagosklonny and Pardee 2002). Cyclin/Cdk activity contributes to the cell’s ability to reach this point (Blagosklonny and Pardee 2002). This mechanism plays an important role in both the entry into S-phase and release from quiescence.

Eukaryotes contain two cyclin E proteins, cyclin E1 and cyclin E2, which are products of two different genes, CCNE1 and CCNE2 (Sherr and Roberts 1999; Geng
et al. 2003; Perez-Neut et al. 2015). Both cyclin E proteins bind and activate Cdk2 (Perez-Neut et al. 2015). Cyclin E is significant as its overexpression is associated with increased proliferation and release from quiescence as well as aneuploidy, polyploidy, and delayed progression through mitosis (Spruck et al. 1999; Keck et al. 2007). Overexpression of cyclin E in cancer cells is associated with increased tumorigenesis and poor clinical prognosis (Said and Medina 1995). In contrast, mouse fibroblasts that lack both cyclin E genes, cyclin E1 and cyclin E2, are unable to exit from the quiescent state but loss of cyclin E has little effect on cells that are already proliferating (Geng et al. 2003).

As cyclin E plays a significant role in cell cycle progression, it stands to reason that it is a highly modified, highly regulated protein. Cyclin E1 and cyclin E2 are regulated by different mechanisms (Perez-Neut et al. 2015). The remainder of this section discusses regulation of cyclin E1. Cyclin E1 is regulated via phosphorylation, both auto phosphorylation when bound to Cdk2 and phosphorylation by different kinases such as GSK3 (Clurman et al. 1996; Welcker et al. 2003). Cyclin E is also regulated in a ubiquitin-dependent manner, resulting in its degradation by the 26S proteasome (Clurman et al. 1996). Lastly, activity of cyclins and Cdk5 is regulated via a class of proteins called cyclin dependent kinase inhibitors or CKIs (Besson et al. 2008). There are two types of CKIs: The first type includes the INK4 proteins, which exclusively regulate Cdk5 and 6 (Sherr and Roberts 1999). The second class, which regulates cyclin E and Cdk2 in addition to others, includes the proteins p27, p21, and p57 (Sherr and Roberts 1999). Proteins
like p21 and p27 inhibit cyclin/Cdk activity by binding to various cyclins and Cdks (Sherr and Roberts 1999).

Additionally, research has demonstrated that in several cancers, including breast, ovarian, melanoma, and others, cyclin E1 can be proteolytically cleaved resulting in truncated low molecular weight (LMW) cyclin E proteins, which mainly lack their N-termini. These LMW cyclin E are able to activate Cdk2 variants are associated with tumorigenesis and poor prognosis in cancer patients (Said and Medina 1995; Harwell et al. 2000; Porter et al. 2001).

**CELL CYCLE REGULATION: Mechanisms controlling cyclin E abundance**

Cells maintain proper levels of Cyclin E through a combination of production via 1) tight transcriptional regulation, ensuring that it is made only during the correct temporal window and 2) degradation (Clurman et al. 1996; Singer et al. 1999). This degradation is also stringently regulated and is frequently initiated by the addition of ubiquitin. The covalent attachment of ubiquitin to cyclin E results in degradation by the 26S proteasome.

Two ubiquitin ligase pathways are responsible for signaling the degradation of Cyclin E. The first is the Cul1 (Cullin1) ubiquitin ligase, which functions as part of a complex that recognizes a phosphorylated Threonine residue (T395 in humans, T393 in mice) on Cyclin E (Clurman et al. 1996; Koepp et al. 2001; Strohmaier et al. 2001; Loeb et al. 2005). After recognizing Cyclin E, the Cul1 complex can ubiquitinate it for degradation by the proteasome. In addition to Cul1, the E3 ligase Cullin3 (Cul3) also functions in the degradation of cyclin E when it is not bound to
Cdk2 (Singer et al. 1999). The mechanism by which Cul3 degrades cyclin E is not well understood. Previous work from our lab which utilized a condition knockout of Cul3 has shown that Cul3 is necessary for the degradation of cyclin E and maintenance of quiescence in the murine liver (McEvoy et al. 2007).

**CUL3 AND DISEASE: Cancer**

Cul3 is an essential gene, as ablation of Cul3 in mice resulted in embryonic lethality (Singer et al. 1999). As Cul3 is necessary for cellular functioning, it is not surprising to learn that Cul3 has been associated with several human diseases, including cancer (Genschik et al. 2013). Cancer is a complicated disease involving many cellular and physiological pathways including those involved in cell division and responses to stress. Cancer develops when a cell undergoes DNA mutation which results in transformative changes involving a variety of cellular pathways that are meant to control proliferation, resulting in a cancerous cell that can grow and metastasize (Penkert et al. 2016). Determining the exact role(s) Cul3 plays in cancer development poses a difficult task as Cul3 regulates dozens of substrates, many of which remain unknown. Here, some of the possible roles of Cul3 in cancer are summarized.

One well-studied substrate of the Cul3 ligase is Nrf2, a transcription factor involved in the oxidative stress response, which is a substrate of Cul3 and the Keap1 BTB protein (Itoh et al. 1999; Cullinan et al. 2004; Kobayashi et al. 2004; Zhang et al. 2004; Furukawa and Xiong 2005). The involvement of the Nrf2-Keap1-Cul3 pathway in many diseases including cancer has been extensively reviewed
In short, the BTB protein Keap1 and other mechanisms tightly regulate Nrf2 when the cell is not under stress, but upon stimulation by stress induced by ROS (reactive oxygen species), a structural modification of Keap1 occurs which reduces degradation of Nrf2 allowing it to activate transcription of Nrf2-controlled genes (Harder et al. 2015). Nrf2 plays a complicated role in cancer as Nrf2 helps to prevent the development of cancer in healthy cells, but some cancerous cells also have elevated Nrf2 activity which is associated with resistance to chemotherapy (Harder et al. 2015). Nrf2 may be the most studied Cul3 substrate, and it alone provides Cul3 with a definitive role in cancer biology.

Another well-studied Cul3 substrate involved in cancer is the cell cycle regulator cyclin E. Like Nrf2, cyclin E levels are tightly maintained by Cul3 and other mechanisms in most cells (Singer et al. 1999; McEvoy et al. 2007). Overexpression of cyclin E is associated with tumorigenesis and truncated cyclin E variants are also associated with tumorigenesis and cancer (Scuderi et al. 1996; Porter and Keyomarsi 2000; Porter et al. 2001). The regulation of cyclin E by Cul3 will be further analyzed in chapter two. There are many other processes that are regulated by Cul3 that also have demonstrated involvement in cancer. For instance, it has recently been revealed that the substrate adaptor RhoBTB2 is responsible for degrading the oncogenic protein MSI2 in breast cancer (Choi et al. 2016), and mutations in the BTB protein SPOP have been demonstrated in prostate cancer (Mani 2014). The involvement of Cul3 and BTB proteins in cancer has been recently reviewed (Genschik et al. 2013; Chen and Chen 2016).
CUL3 AND DISEASE: FHHt hypertension:

Familial hyperkalemic hypertension (FHHt), also known as Gordon’s syndrome or PHAII, is an inherited form of hypertension also characterized by hyperkalemia that can be treated with thiazide diuretics (O’Shaughnessy 2015). Genetic studies have identified mutations in two components of a ubiquitin ligase complex, the cullin scaffold protein Cul3, and the BTB domain containing substrate adaptor protein Klhl3, that cause this disease (Boyden et al. 2012; Louis-Dit-Picard et al. 2012). Mutations in the kinases WNK1 and WNK4 were previously known to cause FHHt, which is known to result from misregulation of the NCC sodium chloride cotransporter located in the distal convoluted tubule (DCT) of the nephron of the kidney (O’Shaughnessy 2015).

WNK1 and WNK4 are kinases that are involved in regulation of NCC. The WNK kinases regulate the activity of NCC and its expression at the membrane surface by phosphorylating substrates including the SPAK kinase, which is responsible for directly phosphorylating NCC (Yang et al. 2003; Hoorn and Ellison 2012; Chavez-Canales et al. 2014). Recently, WNK4 has been identified as a substrate of the ubiquitin ligase complex formed by Cul3 and Klhl3 (Ohta et al. 2013; Shibata et al. 2013; Wakabayashi et al. 2013). Degradation of WNK4 by the Cul3^Klhl3 (nomenclature: Cul3 scaffold containing the Klhl3 BTB protein) ubiquitin ligase complex can partially explain the role of Cul3 in NCC regulation, as WNK4 is necessary for regulation of SPAK and also NCC degradation by the lysosome (Golbang et al. 2006; Hoorn and Ellison 2012). However, patients with Cul3 mutations have a more severe disease phenotype and develop hypertension at an
earlier age in comparison to patients with mutations in Klhl3, WNK4, or WNK1 (Boyden et al. 2012). This led to the hypothesis that Cul3 regulates ion balance and blood pressure by one or more different mechanisms besides the degradation of WNK4, an idea that will be explored further in chapter three.

The Cul3 mutations that cause this disease result in the skipping of exon 9 (encoding amino acids 403-459) of the Cul3 protein. These mutations are dominant and some, including the Cul3 mutation, are believed to be de novo mutations (Boyden et al. 2012). Previous researchers have hypothesized that the Cul3 mutation is a dominant negative (Boyden et al. 2012). However, we speculated that this idea was unlikely since Cul3 has been previously shown to be essential for both embryonic development and cell survival (Singer et al. 1999; McEvoy et al. 2007). We have since demonstrated that the hypertension mutant, Cul3Δ403-459, is a functional ubiquitin ligase capable of ubiquitinating substrates as well as the BTB protein Klhl3 (McCormick et al. 2014).

CUL3 AND DISEASE: Neuronal, eye, and skin conditions

Besides hypertension and cancer, Cul3 has also been implicated in eye, skin, and neurological conditions. Recent work has shown that Cul3 uses the substrate adaptor Klhl24 to ubiquitinate keratin14 and that dominant mutations in Klhl24 result in an increase in keratin degradation and skin fragility (Lin et al. 2016). Additionally, recent research has shown that regulation of RhoA by Cul3 and the substrate adaptor KCTD13 is necessary for brain development (Lin et al. 2015). Mutations in components of Cul3 complexes, including mutations in Cul3 itself, have
been associated with autism, although a mechanism by which Cul3 might lead to autism has not been proposed (Codina-Sola et al. 2015; Lin et al. 2015; Wang et al. 2016). Another BTB protein, RCBTB1, has been associated with retinopathy and retinitis pigmentosa (Coppieters et al. 2016; Wu et al. 2016). Together, these findings indicate that Cul3 is essential for proper cellular and bodily functions in humans. Further study that seeks to determine the mechanisms by which Cul3 functions as well as the identity of more Cul3 substrates will be necessary in order to understand a vast array of human diseases.

**SIGNIFICANCE**

In this chapter, I have given a brief overview of the Cul3 ubiquitin ligase and its involvement in the degradation of cyclin E as well as its role in a variety of other cellular pathways and human disease. In the upcoming chapters, I will provide new insights into the role of Cul3 in cells including its involvement in cyclin E destruction. The data presented in the next chapter will demonstrate that the lysine residue on cyclin E that Cul3 ubiquitinates is located near the N-terminus of cyclin E. This detail provides insight into the role of Cul3 in cancer and tumorigenesis as cyclin E lacking its N-terminus is associated with cancer. Chapter three will further discuss the role of Cul3 and Klhl3 in hypertension. Lastly, chapter four will describe interactions between LRR (leucine rich repeat)-domain containing proteins and Cul3 complexes as well as some other potential future directions for these projects.
REFERENCES


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CHAPTER 2: The N-terminus of Cyclin E is Required for Cul3-mediated Degradation
INTRODUCTION AND BACKGROUND:

Cyclin E and its binding partner Cdk2 regulate the transition from G1 to S-phase as well as release from quiescence in mammalian cells (Koff et al. 1991; Geng et al. 2003). Thus, it is not surprising that cell cycle errors are associated with alterations to cyclin E function and/or abundance; fibroblasts lacking both cyclin E genes, cyclin E1 and cyclin E2, are unable to release from quiescence (Geng et al. 2003). In contrast, overexpression of cyclin E is associated with cancer and tumorigenesis (Said and Medina 1995). Analysis of the cyclin E protein has revealed several functional domains including a central cyclin homology domain which interacts with Cdk2, a unique N-terminal region, and a C-terminal PEST sequence, which is commonly found in proteins that get degraded by the ubiquitin system (Rogers et al. 1986; Rogers and Rechsteiner 1986; Richardson et al. 1993; Honda et al. 2005; Rath and Senapati 2014). In certain cancers, including breast, ovarian and melanoma, cyclin E is known to be cleaved by proteases resulting in N-terminally truncated low molecular weight (LMW) forms ranging in size from 33 to 45 kilodaltons (Scuderi et al. 1996; Harwell et al. 2000; Porter and Keyomarsi 2000; Porter et al. 2001; Wang et al. 2003; Libertini et al. 2005). LMW cyclin E activates Cdk2 and demonstrates increased cyclin E/Cdk2 activity (Porter et al. 2001). These forms of cyclin E are associated with poor clinical prognosis in cancer patients (Harwell et al. 2000; Porter et al. 2001; Duong et al. 2012).

Cyclin E expression is restricted to the G1/S transition by two distinct E3 ubiquitin ligase complexes which are responsible for the degradation of cyclin E: Cul1 and Cul3 (Clurman et al. 1996; Singer et al. 1999; Strehmaier et al. 2001;
Welcker et al. 2003). Both Cul1 and Cul3 are members of the cullin-RING family of ubiquitin ligases. Cul1 or SCF (Skp1, Cul1, Fbw7) based ligases use Fbw7 as a substrate adaptor to recognize cyclin E (Koepp et al. 2001; Strohmaier et al. 2001; Hao et al. 2007). Cul1 mediated-degradation requires phosphorylation of cyclin E at T77 and T395 in order for ubiquitination of cyclin E to occur (Clurman et al. 1996; Welcker et al. 2003; Loeb et al. 2005; Minella et al. 2008). Cul1-mediated degradation of cyclin E occurs about four hours following release from a thymidine block (S-phase) (Bhaskaran et al. 2013).

In contrast to the Cul1 pathway, the mechanistic details of Cul3 mediated destruction of cyclin E remain largely uncharacterized. Similar to Cul1 complexes, Cul3 ubiquitin ligase complexes consist of a substrate adaptor (BTB protein) which binds near the Cul3 N-terminus to recruit substrates, and C-terminal region that binds to the RING finger protein Rbx1 (Tyers and Jorgensen 2000; Jin and Harper 2002; Duda et al. 2008), which in return recruits an E2 ubiquitin conjugating enzyme (Petroski and Deshaies 2005). Previous studies from our lab have shown that Cul3 degrades cyclin E that is not bound to Cdk2 and regulation of cyclin E by Cul3 is necessary for the maintenance of quiescence in liver (Singer et al. 1999; McEvoy et al. 2007). In addition, it has recently been demonstrated that Cul3 can utilize the substrate adaptor RhoBTB3 to ubiquitinate cyclin E (Lu and Pfeffer 2013).

Despite the advances, many details regarding the mechanism utilized by Cul3 for cyclin E ubiquitination remain as yet unknown, including the location of the degron, or region recognized by the ubiquitin ligase, that Cul3 uses to recognize
cyclin E. Here, a Cul3 degron near the N-terminus of cyclin E is identified, and a unique mechanism for Cul3-mediated cyclin E destruction is proposed.

RESULTS:

**Cyclin E binds directly to Cul3 independently of BTB domain-containing proteins**

Cul3 is believed to require a BTB protein to bind cyclin E, but previous work from our lab has shown Cul3 can bind cyclin E in a yeast two-hybrid screen, which suggests that the proteins interact directly with each other (Singer et al. 1999). In order to determine the nature of the interaction between Cul3 and cyclin E, and whether the two proteins interact directly with each other, several Flag-tagged Cul3 mutants were co-transfected with full-length Myc-tagged cyclin E1 and their binding was measured using immunoprecipitation. The mutants represented disruptions of the major functional regions of Cul3; the BTB domain interaction region, the Nedd8 modification site, and the gain of function 403-459 deletion (Zheng et al. 2002; Wimuttisuk and Singer 2007; Boyden et al. 2012; McCormick et al. 2014; Wimuttisuk et al. 2014). Cyclin E bound to all Cul3 mutants tested, including the Cul3Δ51-67 mutant, which cannot bind BTB proteins (Figure 2.1A lane 3). These data imply that cyclin E is able to bind directly to Cul3 without the aid of a BTB protein, unlike what has been shown for other Cul3 substrates.
Figure 2.1: *Cul3 binds cyclin E directly.*  A: Myc-cyclin E was co-transfected in 293 cells with different Flag-tagged Cul3 mutants including Cul3Δ51-67, which cannot bind BTB proteins, Cul3Δ403-459, which causes FHHt, and Cul3K712 R, which is inactive because it cannot be neddylated. Immunoprecipitations (IPs) were performed using a Flag antibody. A western blot using Myc antibody was performed to detect cyclin E. IP results are shown in the top panel. B: Flag-Cul3 or Flag-Cul3
Δ51-67 was co-transfected with Myc-cyclin E, HA-ubiquitin, and S-tagged-SPOP. An IP for Myc-cyclin E was performed followed by western blotting for to detect S-SPOP (B, top panel) and Flag-Cul3 (B, second panel). The lower three panels show the original protein levels of the lysates prior to immunoprecipitation.

In order to further delineate the interaction between Cul3 and cyclin E, Cul3 and cyclin E1 binding was examined in the presence of SPOP, a BTB protein that does not bind or participate in cyclin E degradation (Kwon et al. 2006; Zhang et al. 2014). We found that when cyclin E1 and SPOP are co-transfected with WT Cul3, cyclin E can co-immunoprecipitate SPOP, but cyclin E was not able to co-immunoprecipitate SPOP when the Cul3 mutant Cul3Δ51-67, which cannot bind BTB domain-containing proteins, was used (Figure 2.1B lane 2 compared to lane 3). This shows us that Cul3 was ‘linking’ cyclin E to SPOP in this binding assay. The ability of cyclin E to immunoprecipitate SPOP only when it is bound to Cul3 is consistent with a direct interaction between cyclin E1 and Cul3 and that the binding interaction occurs outside of the region associated with binding BTB domain-containing proteins.
Figure 2.2: Mutants of cyclin E1. A: Cartoon showing full length, wild-type cyclin E (top) and three truncation mutants throughout the protein, STOP100, STOP200, and STOP300. B: Cartoon depicting the locations of point mutants and alanine scanning mutants (designated by amino acid number) on cyclin E. The cleavage site which produces LMW cyclin E is shown in red.
Figure 2.3: The N-terminus of cyclin E interacts with Cul3. A: Upper blot, immunoprecipitation (IP) for Flag-Cul3 and blot for Myc-cyclin E showing binding. Middle and lower blots show relative levels of transfected protein in cell extracts. B: Top blot, IP for Flag-Cul3 and western blot for Myc-cyclin E showing binding of three N-terminally located alanine scanning mutants (lanes 4, 6, and 8) in comparison to wild-type cyclin E (lane 2). The lower two blots show levels of
transfected protein in the cell extracts. All binding assays were repeated a minimum of two times. The result shown in panel B was repeated four times and is reproducible.

**Mutations in the N-terminal region of cyclin E prevent degradation by the Cul3 complex**

In order to determine what region of cyclin E1 binds to Cul3, two C-terminally truncated cyclin E1 mutants were analyzed for binding to Cul3 and compared to full-length cyclin E1. These consisted of the N-terminal 200 amino acids (STOP 200) of cyclin E and the N-terminal 300 amino acids (STOP 300) of cyclin E (Figure 2.2). A STOP100 truncation of cyclin E1 was also transfected, but it was found to be unstable and therefore not used for further experimentation (Figure 2.2). Cul3 bound to both cyclin E1 truncation mutants as well as wild type cyclin E1 (Figure 2.3A compare lanes 2 and 4 to lane 6). To further pinpoint the binding site, we examined the potential of several cyclin E1 alanine-scanning mutants for binding to Cul3 (Kelly et al. 1998). Each mutant in this set contains a charged amino acid sequence that has been mutated to alanines. Amongst the set of mutants, one alanine-scanning mutant DPDEE→AAAA (amino acids 41-45), which is located near the N-terminus, showed decreased binding to Cul3 (Figure 2.3B first panel lane 6 in comparison to lanes 2, 4, 8, and 10).

After determining that Cul3 only requires the N-terminal portion of cyclin E1 for binding, we sought to determine if this region contains all the necessary signals for Cul3-mediated degradation to occur. In order to examine this, the stability of the different mutants was checked in a novel transfection assay utilizing cells that are deficient for Cul3. It was observed that in Cul3 hypomorphic (floxed) MEFs (Mouse
Embryonic Fibroblasts), transfected cyclin E was more abundant than in WT MEFs, similar to the endogenous levels of cyclin E (McEvoy et al. 2007). Controls that were not Cul3 substrates transfected with equal efficiency demonstrating the specificity for Cul3 substrates in this assay. The same regulation of cyclin E was observed when comparing WT (Cul3 containing) and Cul3 KO HEK293 cells (Ibeawuchi et al. 2015) (Figure 2.4A lanes 1 and 2). We reasoned that if a mutant cyclin E, that lacked the Cul3 degron, was transfected into these two genotypes, we would not see a difference in levels of cyclin E. Therefore, transfected cyclin E1 mutants that are Cul3 substrates would be expected to express at higher levels in the Cul3 KO cells in comparison to wild-type 293 cells. On the other hand, a mutant that is not degraded in a Cul3-dependent fashion would be expected to show equal levels of transfected protein in the two cell types.

Before experimentation began, two controls Myc-cyclin E1 and lysineless cyclin E1 were transfected into the two cell types (WT and Cul3 KO 293s). We observed that when transfected with equal amounts of WT cyclin E1, the protein is detected at higher levels in the KO 293 cells than the WT cells (Figure 2.4A compare lanes 1 and 2). In contrast, lysineless cyclin E1 was expressed evenly in both cell types (Figure 2.4A lanes 3 and 4), indicating the utility of this assay as a measure of the substrate being recognized for degradation instead of merely binding. Next, the stability of the cyclin E1 truncations were examined using this assay and we observed that both the STOP 200 and STOP 300 cyclin E truncations are both more stable in the KO cells, implying that the degron recognized by Cul3 is found within the first half of the cyclin E1 protein (Figure 2.4B compare lane 1 to lane 2 and
compare lane 3 to lane 4). Taken together, these results demonstrate that the Cul3 degron resides in the N-terminal half of the cyclin, in contrast to Cul1 the degron which encompasses, in part, T395 located near the C-terminus (Clurman et al. 1996).

*Figure 2.4:* Mutations near the N-terminus of cyclin E result in increased stability. A: Controls for a stability assay using WT and Cul3 KO 293 cells shows that cyclin E, a Cul3 substrate, is more stable when transfected into Cul3 KO cells (A: Left panel). Lysineless cyclin E, which cannot be ubiquitinated, is shown as a negative control.
(A: Right panel). B: Myc-tagged STOP 200 and STOP 300 truncations of cyclin E were transfected into wild-type and Cul3 KO 293 cells and their expression was measured using a Myc (cyclin E) antibody. C: Three Myc-tagged alanine scanning mutants, DPDEE→AAAAA (residues 41-45, lanes 1 and 2), KIDR→AAAAA (amino acids 48-51, lanes 3 and 4) and DKED→AAAAA (amino acids 79-82, lanes 5 and 6), in addition to a triple point mutant, cyclin E K118R, K123R, K125R (Panel C lanes 7 and 8), were transfected into both WT and Cul3 KO cells and their abundance was measured using a Myc (cyclin E) antibody. D: Diagram showing two mutants in which portions of the N-terminal domain have been deleted. E: Expression of transfected Myc-cyclin EΔ31-82 and Myc-cyclin EΔ2-86 are shown in both the WT and KO cells (E, right panel, lanes 1 through 4). Quantification of each western blot (WB) is listed below each lane as a percent relative to the sample in the KO lane for each pair (KO is always 100 percent). Transfections were repeated a minimum of three times and representative results are shown.
**Figure 2.5:** Localization of cyclin E mutants resembles wild-type. Myc-tagged cyclin E mutants were transfected into HeLa cells and localization was determined using immunofluorescence. The localization of several mutants is shown: DPDEE→AAAAA (row 2), KIDR→AAA (row 3), and DKED→AAAA (row 4). The top row shows the localization of wild-type cyclin E.
Lysine 48 on cyclin E serves as a ubiquitination site for Cul3

To pinpoint specific residues on cyclin E that may be involved in degradation, cyclin E alanine scanning mutants were transfected into the Cul3 WT and KO 293 cells. Three alanine-scanning mutants, DPDEE→AAAAA (amino acids 41-45), KIDR→AAAA (amino acids 48-51), and DKED→AAAA (amino acids 79-82), appear more stable in the wild-type cells, which suggests that they are less likely degraded by Cul3 (Figure 2.4C). All three of these mutants are located near the N-terminus of cyclin E and in order to further describe the importance of this region for Cul3-mediated degradation to occur, two constructs containing deletions in this region were created (Figure 2.4D). The first construct, cyclin EΔ31-82, is missing the region containing all three of the stabilized alanine scanning mutants. The second construct, cyclin EΔ2-86, resembles the LMW cyclin E found in some cancer cells, and is missing the entire N-terminal region (Figure 2.4D). Both of these mutants were transfected into the wild-type and Cul3 KO HEK293 cells, and both demonstrated increased stability in the WT cells (Figure 2.4E, compare lane 1 to 2, and lane 3 to lane 4), suggesting that this region may be necessary for Cul3-mediated degradation of cyclin E to occur.

Sometimes, phenotypes exhibited by mutant proteins can be explained by changes in the localization of the protein within the cell. In order to determine if the phenotypes of the three stable alanine-scanning mutants are the result of localization changes, HeLa cells were transfected with each construct and visualized using immunofluorescent microscopy. The cellular localization of the three mutants was found to be predominantly nuclear, similar to wild-type cyclin E, indicating that
their increased stability was not a result of a change in cellular localization (Figure 2.5). The DPDEE→AAAAA (residues 41-45) mutant cannot bind Cul3 as well as the others (Figure 2.3), which provides an explanation for its increased stability in our assay. As the KIDR→AAAA (residues 48-51) and DKED→AAAA (residues 79-82) mutants both contain lysine residues and are located near the N-terminus of cyclin E (Figure 2.4C). The increased stability of the KIDR and DKED alanine scanning mutants as well as the deletion mutants imply that both lysine residues K48 and K80 are potential ubiquitination sites, and the degron that is recognized by Cul3 likely resides within the N-terminal portion of cyclin E.

To establish if ubiquitination on K48 or K80 of cyclin E regulates its stability, we determined if the KIDR→AAAA mutant or DKED→AAAA mutant stabilized by a mutation in ubiquitin that prevents K48 branching, a type of branching that leads to degradation (Grice and Nathan 2016). To test this, all of the alanine scanning mutants within the first 200 amino acids of cyclin E were transfected into both cell types in the presence or absence of a mutant K48R ubiquitin. We observed that, unlike WT cyclin E which was stabilized in WT cells by the addition of the dominant-negative K48R ubiquitin mutant (Figure 2.6 compare lanes 1 and 2 to lanes 3 and 4), the KIDR→AAAA mutant was unaffected indicating it is no longer a substrate for ubiquitination-dependent degradation (Figure 2.6 lanes13-16). The DPDEE→AAAAA (Cul3 binding mutant, lanes 9-12) mutant behaved similarly to the KIDR→AAAA mutant, which suggests that they might comprise a Cul3 degron, which consists of a binding region (DPDEE) and ubiquitination site (K48 on cyclin E, the lysine contained in the KIDR mutant). The DKED→AAAA mutant (Figure 2.6 lanes
17-20) appears similarly to wild-type (Figure 2.6 lanes 1-4) in this assay suggesting the K48 (the KIDR lysine) and not K80 (the DKED lysine) is the ubiquitination site for Cul3. As a second control, cyclin E T395A, which cannot be degraded by Cul1, was examined and we observed that it is a substrate of Cul3 (Figure 2.6 lanes 5 through 8). In order to confirm that K48 is an ubiquitination site on cyclin E, a point mutant, cyclin E K48R, was constructed for use in the ubiquitination assay. Like the KIDR mutant, cyclin E K48R appears relatively stable (Figure 2.6 lanes 21-24). The increased stability of the KIDR mutant and cyclin E K48R suggests that K48 on cyclin E is indeed an ubiquitination site utilized by Cul3 in vivo. Taken together, these data suggest that the residues DPDEE (binding site) and KIDR (ubiquitination site) are part of a degron that is recognized by Cul3.

![Figure 2.6](image)

**Figure 2.6:** *K48 is a Cul3 ubiquitination site on cyclin E.* Alanine scanning and point mutants located within the first 200 amino acids of the cyclin E protein were transfected in WT and KO 293 cells with or without the addition of an S-tagged K48R ubiquitin construct. Wild-type cyclin E (lanes 1-4) and cyclin E T395A, which cannot be degraded by Cul1 (lanes 5-8), were included as controls. Three cyclin E alanine scanning mutants are shown here, DPDEE→AAAAA (residues 41-45, lanes 9-12), KIDR→AAAA (residues 48-51, lanes 13-16), and DKED→AAAA (residues 79-82, lanes 17-20). The point mutant cyclin E K48R is also shown (lanes 21-24).
The N-terminal domain of cyclin E is required for degradation

In order to truly measure if cyclin E that lacks its N-terminus is differentially degraded by Cul3 compared to wild-type cyclin E, it is necessary to measure the rate of degradation of these proteins. To ascertain the role of the N-terminal domain of cyclin E degradation, cyclin E half-lives were measured and compared in cells of both genotypes (Cul3 WT and KO). Cells were transfected with full-length Myc-tagged cyclin E or Myc-tagged cyclin E Δ2-86. Following addition of cycloheximide (CHX) the cells were harvested every two hours for ten hours. We observed that WT cyclin E has a half-life of about 2 hours in the WT cell line but a half-life of 5 hours in the KO cells demonstrating that loss of Cul3 reduces the half-life of cyclin E (Figure 2.7 compare WT cyclin E (left) to cyclin E Δ2-86 (right). We also observed that cyclin E Δ2-86 has a 5 hour half-life and shows equal stability in both WT and Cul3 KO cells, demonstrating the inability of Cul3 to target it for degradation (Figure 2.7, right panel, lanes 1-5 vs. lanes 6-10).
Figure 2.7: The N-terminal domain of cyclin E is required for Cul3-mediated degradation. Cul3 WT and KO 293 cells were transfected with Myc-tagged cyclin E (Left) or Myc-tagged cyclin EΔ2-86 (Right). After 24 hours, cycloheximide was added and cells were harvested at the indicated time points. Half-lives were determined via Western blots (top), and quantified (bottom). Repetition of this experiment yields similar results.

Cul3 cannot degrade endogenous LMW cyclin E

As low molecular weight (LMW) cyclin E can be found endogenously in some cells, we sought to determine if loss of Cul3 affects the presence of these LMW forms. It has been previously shown that the LMW cyclin E truncations can still be degraded in a Cul1-dependent manner (Delk et al. 2009). Overall, the Cul3 KO 293 cells have more endogenous cyclin E than the wild-type (Ibeawuchi et al. 2015). We observed that the 50 kDa endogenous cyclin E band increases upon inhibition of the proteasome in WT cells to equal the amount of the 50 kDa protein that is present in the KO cells prior to proteasome inhibition, demonstrating that the 50 kDa band is a
substrate of Cul3 in 293 cells (Figure 2.8 lane 1 vs lane 3). Two LMW cyclin E bands are also detected, the smallest of which is about 43 kDa, and they appear to be equally expressed in WT and KO cells (Figure 2.8 lanes 1 and 3). The relative abundance of the LMW bands is elevated equally in both cell types upon proteasome inhibition (Figure 2.8 lanes 2 and 4), indicating that LMW cyclin E still targeted for ubiquitin-mediated proteolysis equally in both cell types.

![Figure 2.8: LMW cyclin E is not degraded by Cul3.](image)

The upper blot shows levels of endogenous cyclin E in WT and Cul3 knockout (KO) 293 cells. The proteasome inhibitor MG132 has been added to the cells shown in lanes 2 and 4. The lower blot shows levels of actin in the same cells.

**Cyclin E2 is not a substrate of Cul3**

Mammals contain two cyclin E genes, CCNE1 (cyclin E1) and CCNE2 (cyclin E2), which produce different proteins (Sherr and Roberts 1999; Geng et al. 2003). The two cyclin E proteins share high homology and many structural similarities, including the Cdk2 interacting domain (Perez-Neut et al. 2015). In order to determine if the observed degradation of cyclin E was specific to cyclin E1, cyclin E2
was transfected into the WT and KO 293 cells. Cyclin E2 is evenly expressed in both cell types, indicating that it does not get ubiquitinated by Cul3 (Figure 2.9, lanes 1 and 3 compared to lanes 2 and 4).

### Table

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<tr>
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<th>+</th>
<th>+</th>
<th>+</th>
<th>+</th>
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<tr>
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<td>KO</td>
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<td>KO</td>
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**Figure 2.9:** *Cyclin E2 is not a Cul3 substrate.* Transfected, Myc-tagged cyclin E2, shown here in duplicate is shown here in WT (lanes 1 and 3) and Cul3 KO (lanes 2 and 4) HEK 293 cells.

### Loss of Cul3 results in early accumulation of cyclin E after release from serum starvation

In order to determine when loss of Cul3 resulted in cyclin E accumulation we turned to the Cul3 knockout 293 cells. Previous work from our lab has shown that loss of Cul3 in mouse embryonic fibroblasts (MEFs) results in increased amounts of cyclin E and a greater percentage of cells in S-phase (McEvoy et al. 2007). Analysis of Cul3 KO 293 cells by flow cytometry shows an increased percentage of cells in S phase when compared to WT 293s (Figure 2.10). This result shows that the excess cyclin E in the Cul3 KO 293 cells is functional as it is capable of initiating DNA replication. To determine at which point during the cell cycle the Cul3 KO cells begin to accumulate excess cyclin E, both WT and KO cells were arrested in G1 via serum starvation and released. The KO cells enter S-phase earlier than their wild-
type counterparts, as four hours after release into G1, only 47.1 percent of wild-type cells are in S-phase or G2 but 61.7 percent of Cul3 KO cells have reached this point (Figure 2.11, WT cells are shown on the left and KO are shown on the right).

In order to determine if the increased percentage of cells in S-phase is a result of cyclin E accumulation during G1, we utilized a similar serum-starve and release experiment. We examined cyclin E accumulation in Cul3 hypomorphic MEFs (mouse embryonic fibroblasts) after serum starvation and release. We observed cyclin E accumulation 2 hours earlier in Cul3 hypomorphic (floxed) MEFs than in WT MEFs (Figure 2.12 WT top row, Cul3 hypomorphic flx/flx bottom row). This is consistent with a role for Cul3 in regulation of cyclin E levels in quiescent cells, as we had observed in an animal model (McEvoy et al. 2007), and indicates Cul3 regulation of cyclin E occurs in a different temporal window than the Cul1 regulation of cyclin E which takes place 4 hours after the onset of DNA replication (Bhaskaran et al. 2013).
Figure 2.10: *The excess cyclin E in Cul3 KO 293 cells increases proliferation.* Flow cytometry analysis of proliferating 293 cells stained with propidium iodide, which labels the DNA allowing for the quantification of DNA content. WT cells are shown on the left and Cul3 KO cells are shown on the right. Cells in G₁ are shown in purple, S-phase cells are shown in yellow, and G₂/M cells are shown in green. At least 20,000 cells were counted for each experiment. The quantification show represents and average of three experiments.
**Figure 2.11:** *Cells lacking Cul3 enter S-phase early.* Cells were serum-starved and released into G1. Cells were harvested at 4-hour intervals, stained with propidium iodide, and analyzed by flow cytometry. Quantification of the resulting cell cycle profiles is shown (Right).

<table>
<thead>
<tr>
<th>Time post-release</th>
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<th>Cul3 KO cells</th>
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<tr>
<td></td>
<td>G1</td>
<td>S/G2</td>
</tr>
<tr>
<td>Arrested</td>
<td>61.5%</td>
<td>38.5%</td>
</tr>
<tr>
<td>4 Hours</td>
<td>53.3%</td>
<td>47.1%</td>
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<tr>
<td>8 Hours</td>
<td>41.1%</td>
<td>58.9%</td>
</tr>
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<td>12 Hours</td>
<td>35.1%</td>
<td>65.0%</td>
</tr>
<tr>
<td>16 Hours</td>
<td>35.2%</td>
<td>64.8%</td>
</tr>
<tr>
<td>25 Hours</td>
<td>66.7%</td>
<td>33.3%</td>
</tr>
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**Figure 2.12:** *Cyclin E levels increase early in cells that are hypomorphic for Cul3.* Western blot showing levels of endogenous cyclin E in mouse embryonic fibroblasts (MEFs) that are wild-type (WT, top row) or deficient for Cul3 (flx/flx, bottom row). Cells that had been serum-starved and released were collected at two-hour intervals as indicated.
**RhoBTB3 interacts with UbE2E1 to facilitate complex formation**

The previous data imply that cyclin E can bind Cul3 without the assistance of a BTB domain-containing substrate adaptor such as RhoBTB3. We therefore became interested in what potential role RhoBTB3 had in Cul3 mediated cyclin E degradation, as it has been suggested that RhoBTB3 facilitates Cul3-mediated ubiquitination of cyclin E (Lu and Pfeffer 2013). Like Cul3, RhoBTB3 binds to the cyclin E STOP 200 and STOP 300 truncations (Figure 2.13 panel A lanes 2 and 4). RhoBTB3 shows a decreased ability to bind the DPDEE→AAAAA (amino acids 41-45) alanine scanning mutant in comparison to other mutants tested, which is also similar to Cul3 (Figure 2.14 panel B, compare lane 6 to lanes 2 and 4). Binding of RhoBTB3 to the DKED→AAAA (amino acids 79-82) alanine scanning mutant also appears reduced (Figure 2.13 lane 8).

Since RhoBTB3 is not essential for the Cul3/cyclin E interaction, we speculated that it might be playing another role in the active complex. We hypothesized that since specific E2 recruitment should be an essential role for the E3 ligase it may be that RhoBTB3 is responsible for recruiting the E2 enzyme that ubiquitinates cyclin E. If BTB proteins are involved in E2 selection, it would ensure that the proper E2 is recruited to form the correct ubiquitin linkage for a given substrate. To test this hypothesis, we examined the binding of Cul3 and RhoBTB3 to the E2 enzyme UbE2E1 which forms degradative K48-linked chains and has been shown to associate with Cul3 (Plafker et al. 2009). We observed that both Cul3 and RhoBTB3 bind to UbE2E1 (Figures 2.14 lane 2 and 2.15 lane 2). We then examined if Cul3 binding to UbE2E1 requires the BTB domain interacting region on Cul3. We
observed that UbE2E1 cannot bind the Cul3 mutant that is incapable of binding BTB proteins, Cul3Δ51-67 (Figure 2.15 lane 3). This indicates that the interaction with the E2 enzyme is likely not entirely mediated by RhoBTB3. To confirm that Cul3 does not mediate the interaction between RhoBTB3 and UbE2E1, Cul3 KO 293 cells were used to analyze binding (Ibeawuchi et al. 2015). As can be seen in lanes two and five, RhoBTB3 binds to the E2 enzyme in the absence of Cul3 (Figure 2.14 lane 4) to the same degree that the two proteins bind in the presence of Cul3 (Figure 2.14 lane 2) indicating that Cul3 is not required for the BTB to interact with the E2 and BTB proteins may be necessary to help associate the E2 enzyme with Cul3. Together these results demonstrate that a BTB protein, in this instance RhoBTB3, can interact with an E2 enzyme independently of Cul3 and may enhance E2 binding in vivo.
Figure 2.13: *RhoBTB3 interacts with the N terminus of cyclin E*. A: Upper blot, immunoprecipitation results showing binding between S-tagged RhoBTB3 and Myc-tagged cyclin E truncations. The lower blots show levels of the transfected proteins in the cell extracts. B: The upper blot shows immunoprecipitations results to check for binding between S-RhoBTB3 and Myc-cyclin E alanine scanning mutants. The Four mutants shown here are 27-29 (RSR, lanes 1 and 2), 30-32 (KRK, lanes 3 and 4), 41-45 (DPDEE, lanes 5 and 6), and 79-82 (DKED, lanes 7 and 8). The lower blots show the expression of the transfected proteins in cell lysates.
Figure 2.14: *RhoBTB3 binds an E2 enzyme.* RhoBTB3 was co-transfected with the E2 ubiquitin conjugating enzyme HA-tagged UbE2E1 in the presence of endogenous Cul3 (WT cells, lanes one and two) and absence of Cul3 (KO cells, lanes three and four). Immunoprecipitations (IPs) were used to check binding.

Figure 2.15: *Cul3 requires a BTB protein to interact with UbE2E1.* Flag-WT Cul3 (lane 2) and Flag-Cul3Δ51-67, which cannot bind BTB proteins, (lane 3) were co-transfected with the HA-tagged E2 enzyme UbE2E1. Immunoprecipitations using Flag antibody were conducted to check for binding. The top row shows the IP results while the bottom two rows show the protein expression in the cell extracts.
DISCUSSION:

Cyclin E protein accumulates in late G1, peaks in early S-phase and rapidly disappears. Increased levels of cyclin E are associated with cell cycle errors and tumorigenesis and loss of cyclin E in MEFs results in the inability of the cells to exit from quiescence (Said and Medina 1995; Geng et al. 2003). Cyclin E undergoes several post-translational modifications in cells including phosphorylation and ubiquitination resulting in its degradation by the proteasome. Research has shown that cyclin E can be proteolytically cleaved resulting in truncated forms of the cyclin, many of which lack portions of the protein near the N-terminus (Porter et al. 2001; Wang et al. 2003). These LMW forms of cyclin E are associated with poor prognosis in cancer patients and overexpression of the LMW cyclin E has also been shown to cause increased cell cycle errors and chromosome instability in cell culture models compared to full length cyclin E (Bagheri-Yarmand et al. 2010). Others have shown that the LMW forms of cyclin E have an increased ability to activate Cdk2, resulting in an increase of Cdk2 kinase activity (Porter et al. 2001; Wingate et al. 2005). It has also been demonstrated that these cyclin E truncations can be degraded in a ubiquitin-dependent manner (Delk et al. 2009).

Two ubiquitin ligase pathways, Cul1 and Cul3, are known to degrade cyclin E. Cul1-mediated degradation has been extensively investigated and it is known to degrade cyclin E that is bound to Cdk2 and require phosphorylation of residues T77 and T395 on cyclin E in order to recognize it for degradation (Clurman et al. 1996; Loeb et al. 2005; Minella et al. 2008). In contrast, the Cul3-based pathway is not as well understood and is known to degrade cyclin E that is not bound to Cdk2 (Singer
et al. 1999). LMW cyclin E containing complexes have been shown to be more poorly inhibited by the cyclin-dependent kinase inhibitors (CKIs) p21 and p27 (Akli et al. 2004; Wingate et al. 2005). It has been proposed that the LMW cyclin E may be capable of sequestering the CKIs and therefore preventing them from interacting with full length cyclin E, which is believed to be more susceptible to inhibition, ultimately resulting in the increased Cdk2 activity that is associated with the LMW cyclin E (Wingate et al. 2005). Our data implies that decreased inhibition of LMW cyclin E by p27 and p21 may only provide a partial explanation for the accumulation of LMW cyclin E and a secondary Cul3-based mechanism may also contribute to the increased activity. This mechanism results from bypassing Cul3 degradation of cyclin E by cleaving off the N-terminal degron of cyclin E making more available to interact with Cdk2 (Figure 2.16).

We show that K48 is a ubiquitination site on cyclin E utilized by Cul3 (Figure 2.6). This finding is notable given what is known about the structure of the cyclin E protein as the N-terminus, which contains K48, is believed to be a disordered, mostly hydrophilic region (Rath and Senapati 2014). The structure of cyclin E bound to Cdk2 has been determined (Honda et al. 2005), but the published structure only includes amino acids 81-363 of cyclin E and therefore did not include the N-terminus of the cyclin. Recently, advanced modeling techniques have predicted a structure for the N-terminal region of cyclin E which shows that this portion of the protein is mostly disordered with the notable exception of a predicted alpha helix encompassing the sequence “DEEMAKID” (amino acids 43-50, Figure 2.18A) (Rath and Senapati 2014). Coincidentally, the alanine scanning mutants in this region
(DPDEE→AAAAA and KIDR→AAAA) exhibited the most notable phenotypes in our study as DPDEE exhibited diminished binding to the Cul3 complex and KIDR→AAAA cannot be degraded in a Cul3-dependent manner (Figures 2.3, 2.4 and 2.6). Our data suggests that this structural motif may form part of the interface on cyclin E, which is responsible for Cul3 interaction and also functions as the Cul3 degron on cyclin E (Figure 2.17A). The putative helix comprising the Cul3 degron, including the K48 ubiquitination site, is conserved in other mammalian species (Figure 2.17B), which is notable as Cul3 also regulates cyclin E in mice (Singer et al. 1999; McEvoy et al. 2007). Additionally, cyclin E2, which cannot be degraded by Cul3 (Figure 2.9), does not resemble cyclin E1 in the region of the newly proposed degron (figure 2.17C). The net charge of the region was calculated by finding the difference between the number of basic residues and the number of acidic residues. Using this method, it can be seen that cyclin E1 is very acidic in this region with a net charge of negative two. Cyclin E2, in contrast, is extremely basic with a net charge of four (Figure 2.17C). Additionally, a Chou-Fasman algorithm was used to predict possible structures of both cyclin E proteins in the degron region (Chou and Fasman 1975). This method revealed what is possibly a helical region in cyclin E1 but an unstructured region in cyclin E2 (Figure 2.17C). Taken together, this information suggests that the Cul3 degron comprises a structural feature that is unique to cyclin E1. These data demonstrate the critical role of the Cul3 complex in degradation of cyclin E by providing evidence that suggests that Cul3’s ability to degrade LMW cyclin E which lack their N-termini may be impaired as the Cul3 degron is located near the cyclin E N-terminus. Cul3 uses a mechanism distinct from the Cul1-based
pathway to recognize cyclin E for degradation, as Cul3 does not require cyclin E to be phosphorylated on T395 for degradation to occur (Figure 2.6 lanes 5-8).

The data presented here support the idea that LMW cyclin E is still a substrate of the Cul1 ubiquitin ligase pathway since LMW cyclin E increases in the Cul3 KO 293 cells upon the addition of MG132, indicating that it is still degraded in a ubiquitin-dependent manner (Figure 2.8). In addition, other labs have also shown that LMW cyclin E can still be degraded by the ubiquitin proteasome system (Delk et al. 2009).

It has been suggested that Cul3 utilizes the substrate adaptor RhoBTB3 to target cyclin E for degradation (Lu and Pfeffer 2013). Our work sheds light on the mechanistic purpose of RhoBTB3 in cyclin E degradation as we demonstrate that RhoBTB3 is involved in the binding of the E2 ubiquitin conjugating enzyme and can bind the E2 UbE2E1 both in the presence and absence of Cul3 (Figure 2.1). This finding is significant as it suggests that Cul3 substrate adaptors may play a role in E2 selection, which would help to explain how Cul3 is capable of forming a variety of ubiquitin chain linkages as different E2s are associated with this process. The revelation that Cul3Δ51-67, which cannot bind BTB proteins, is incapable of binding UbE2E1 (Figure 2.15 lane 3 compared to lane 2) presents a second piece of information in support of this idea that BTB proteins are necessary for E2 selection to occur. These findings may shed light on interactions between Cul3, BTB proteins, and E2s, a finding which may be applicable to other BTB proteins and substrates. In addition to clarifying the mechanistic details regarding Cul3-mediated degradation of cyclin E, the work presented here also identifies the temporal window during
which such degradation occurs. Previous work has shown that loss of Cul3 results in cyclin E accumulation and exit from quiescence in mice (McEvoy et al. 2007), leading us to hypothesize that loss of Cul3 results in earlier increases in cyclin E and entry into S-phase. Here, this hypothesis is supported as cells that are hypomorphic for Cul3 both enter the cell cycle earlier than their WT counterparts and show increased levels of cyclin E earlier than wild-type cells following release from quiescence (Figure 2.11 and 2.12). These data suggest that Cul3 is responsible for maintaining levels of cyclin E earlier in the cell cycle and preceding the start of S-phase, which is again in contrast to the Cul1 degradation pathway that degrades cyclin E later during S-phase (Bhaskaran et al. 2013). Taken together, our data suggests that during G1, Cul3 functions to suppress levels of cyclin E via ubiquitination of its N-terminal domain (Figure 2.18). Lack of cyclin E regulation by Cul3 during G1 might contribute to the increased cyclin E/Cdk2 activity that is observed in LMW cyclin E containing cancer cells (Figures 2.16 and 2.18).
**Figure 2.16:** *LMW cyclin E cannot be degraded by Cul3.* This model demonstrates Cul3’s role in degradation of full-length cyclin E (Left) and contrasts it with improper regulation of LMW cyclin E (Right). Full length cyclin E is ubiquitinated and degraded via Cul3 during G1, leaving some cyclin E available to activate Cdk2. However, in cells containing LMW cyclin E which lacks its N-terminal domain, proper degradation of LMW cyclin E by Cul3 might not occur, which might result in increased activation of Cdk2.
Figure 2.17: Cyclin E1 contains a putative Cul3 degron that is conserved in other mammals but lacking in cyclin E2. A: Others have proposed a helix located in the N-terminal region of cyclin E1 (Rath and Senapati, 2014). The proposed helix (A, shown in blue) overlaps the DPDEE (Cul3 binding) and KIDR (ubiquitination site) alanine scanning mutants (shown in red), suggesting that the Cul3 degron is part of a structural feature on cyclin E. B: The Cul3 degron is highly conserved in mouse (Mus musculus) and rat (Rattus norvegicus) cyclin E1 (outlined in red). C: Sequence comparison of the degron region in cyclin E1 (bottom) to the same region in cyclin E2 (top). Chou-Fasman analysis of the amino acid sequences suggests that the structures differ between cyclin E1 and cyclin E2. Additionally, the Cul3 degron in cyclin E1 is acidic, with a net charge of minus two (difference between the number of acidic residues and the number of basic residues). The same region in cyclin E2 is extremely basic with a net charge of plus four.

Figure 2.18: Cul3 degrades cyclin E during G1. Our data is consistent with a model where Cul3 is responsible for maintaining cyclin E levels during G1 in order to prevent early entrance into S-phase. Cul1 is known to degrade cyclin E after S-phase has begun, so therefore the two ligases work to degrade the cyclin at two different points during the cell cycle.
MATERIALS AND METHODS

**Cell culture and transfections:** Cells (HEK 293, HeLa, and MEFs) were maintained in DMEM supplemented with ten percent fetal bovine serum and penicillin/streptomycin. HEK 293 and HeLa cells were split 1:20 for transfection in 6cm dishes the night before transfection. Transfections were completed using calcium phosphate to precipitate the DNA onto the cells. For immunoprecipitations and expression level assays, cells were harvested 48 hours post transfection. For HeLa and 293 cells, between 1 and 10 µg of plasmid DNA was transfected into each plate. For experiments using cycloheximide, drug was added 24 hours post-transfection at a final concentration of 50 µg per milliliter. Cells were then harvested at the time points indicated. For experiments utilizing MG132, drug was added approximately 18 hours before harvest at a final concentration of 20µM. All transfections were harvested using RIPA buffer and then sonicated before being used for immunoprecipitations or western blots. CRISPR knockout Cul3 293 cells were a gift from Curt Sigmund.

**Plasmids:** 3x-Flag-Cul3 was used for all Cul3-containing transfections. All Cul3 mutants (Cul3 K712R, Cul3Δ51-67, and Cul3Δ403-459) were expressed using the same p3x-Flag vector as wild-type Cul3. Cyclin E mutants were expressed using the CS2+ Myc-tagged expression vector, and CS2+ S-tagged and CS2+HA tagged vectors were utilized for expression of BTB proteins and the E2 enzyme UbE2E1.

**Western blotting and immunoprecipitations:** Western blots and immunoprecipitations were conducted as previously described (Wimuttisuk et al.
2014). In short, a sonicated transfection lysate was added to the desired antibody in an Eppendorf tube and brought to a final volume of 500μL. 40μL of IPA sepharose beads were then added to the mixture, and the IPs were placed on a rotator for two hours at room temperature before being rinsed with RIPA buffer, heated in SDS-loading buffer, and run on an SDS-PAGE gel. The following antibodies were used for immunoprecipitations and/or western blotting: Monoclonal anti-FLAG (Sigma, F1804-50UG), monoclonal anti-Myc (9E10, Santa Cruz), polyclonal anti-c-Myc (A14) (Santa Cruz, discontinued), S-peptide monoclonal antibody (6.2) (Fisher, Cat# MA1-981), monoclonal anti-HA.11 (16B12) (ThermoFisher), polyclonal anti-HA (ThermoFisher, Product # PA1-985), polyclonal anti-β actin (ThermoFisher, Product # PA1-183), polyclonal anti-cyclin E (Singer et al. 1999), polyclonal anti-Cul3 (Singer et al. 1999; McEvoy et al. 2007), and monoclonal anti-cyclin E (HE12, Santa Cruz biotechnology). Quantification of the western blots for all cycloheximide experiments was done using the FluorChem SP software (Alpha Innotech) and graphs were generated using Microsoft Excel.

**Immunofluorescence:** Hela cells were grown on coverslips, transfected, followed by incubation in four percent paraformaldehyde/PBS at room temperature for 10 minutes. Cells were then permeablized using a solution of one percent Triton X-100 with 2mM EGTA and 5mM PIPES, followed by incubation in methanol at -20 degrees for 10 minutes. Cells were then rinsed with PBS and stained overnight with a polyclonal Myc antibody (A14, Santa Cruz). The next day, cells were rinsed and stained with an AlexaFluor 488 conjugated secondary antibody (Abcam, ab150077)
followed by DAPI, rinsed in methanol, and mounted on coverslips for viewing. Microscopy was conducted using a Zeiss M2 microscope and AxioVision software.

**Flow cytometry:** Proliferating cells were harvested when they were 70 percent confluent. Cells were resuspended in 70 percent ethanol in PBS and stored at 4 degrees until analysis. Prior to analysis, fixed cells were stained in a solution of propidium iodide in PBS with RNase A at 37 degrees for a minimum of 30 minutes. Cells were then strained and analyzed on a BD Accuri C6 benchtop flow cytometer. Three proliferating samples of each genotype were analyzed and 20,000-50,000 cells were counted for each sample. Analysis shown in figure 2.10 was completed using FlowJo software (TreeStar). Cell cycle analysis of the samples in Figure 2.11 was completed by hand.

**Construction of cyclin E mutants:** The majority of the alanine scanning mutants were a kind gift from Jim Roberts at the Fred Hutchinson Cancer Center (Kelly et al. 1998). The point mutants were made using site-directed mutagenesis and all point mutants were confirmed by sequencing. Alanine scanning mutants that demonstrated a phenotype were also re-confirmed by sequencing. Truncations were cloned by using mutagenesis to generate a stop codon at the designated location in the cyclin E protein. Both deletions were made using site-directed mutagenesis and were also confirmed by sequencing. Forward primer sequences for mutagenesis are as follows: Cyclin E K80R (DKED point mutant) 5'--TCCCCACACCTGACAGAGAAGATGATGACCG, Cyclin E 3 lysine (K118, 123, 125R) 5'--AGAGGAAGTCTGGAGAATCATGTTAAACAGGGAAAGGACATACTTAAGGG, Cyclin E Δ 2-86 5'--GGACTTGAAATTCCATGGTTTACCACCTCAA, Cyclin E KIDR point mutant
(K48R) 5’-CGCGGTCTCTGAGTTCGAGGCCATTCTTCAT, Cyclin EΔ31-82 5’-
GCTCGCTCCAGGAAGGATGACCGGGTTTAC. Primer sequences for the cyclin E
truncations are as follows: Cyclin E STOP 200 5’-
TCATCTTTATTTATGGAGCCAAACTTGAGGAA, and STOP 300 5’-
TTTCTTTATGGTATATGAGCTGCTTCGGCCTGG. All reverse primers were reverse
complements of the forward primers.

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insights into NEDD8 activation of cullin-RING ligases: conformational control


CHAPTER 3: Identifying Possible Roles of Cul3 and Klhl3 in the Kidney
**BACKGROUND: The role of Cul3 and Klhl3 in FHHt**

Mutations in the BTB-Kelch protein Klhl3 as well as Cul3 have recently been shown to cause an inherited form of hypertension called Familial Hyperkalemic Hypertension (FHHt) (Boyden et al. 2012; Louis-Dit-Picard et al. 2012). This form of hypertension results primarily from the misregulation of NCC sodium chloride cotransporter located on the apical membrane of the distal convoluted tubule of the nephron (O'Shaughnessy 2015). Klhl3 can affect NCC, resulting in disease, via degradation of the WNK4 kinase, which is responsible for controlling NCC through a phosphorylation pathway involving the SPAK and OSR1 kinases (summarized in Figure 3.1) (McCormick and Ellison 2011; Shibata et al. 2013). Mutations in Klhl3 that alter its ability to bind and ubiquitinate WNK4 result in changes in NCC phosphorylation status, which leads to the disease phenotype (Ohta et al. 2013; Shibata et al. 2013; Wakabayashi et al. 2013). Interestingly, the FHHt-causing mutations in Cul3 and Klhl3 are associated with a more severe disease phenotype than the previously known FHHt-causing mutations in WNK4 and the related kinase WNK1 (Boyden et al. 2012). This information suggests that Klhl3 and Cul3 may have additional unknown roles in this pathway. In order to elucidate Cul3’s role in FHHt, two approaches, one focused on Cul3 and one focused on Klhl3, were taken: First, the structure and function of the hypertension-associated Cul3 mutation, Cul3Δ403-459, was studied, and secondly, a screen was performed in order to identify potential binding partners of Klhl3 in vivo.
Figure 3.1: **NCC regulation.** NCC is regulated via phosphorylation, which requires multiple kinases such as SPAK, OSR1, and the WNK kinases. There are several possible points at which Cul3 can regulate this pathway: The Cul3\textsuperscript{Khl2} E3 ligase has been shown to ubiquitinate WNK4, but other possible roles of Cul3 in this process remain unknown.

As previous work from our lab has demonstrated that Cul3 is an essential gene as its loss results in embryonic lethality in mice, we hypothesized that the newly discovered human Cul3 mutation is a functional ubiquitin ligase capable of ubiquitinating substrates and therefore the FHHt-associated Cul3 mutant is not a loss of function mutant (Singer et al. 1999). We sought to determine the biochemical properties of the human Cul3 mutations as well as the role of the Cul3 E3 ligase in the regulation of blood pressure (McCormick et al. 2014). This work, which was published in 2014, demonstrated that the hypertension-causing Cul3 mutant, Cul3\textDelta403-459, is a gain of function mutant that has an increased ability to

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interact with the substrate adaptor Klhl3 as well as an increased ability to ubiquitinate substrates, including its substrate adaptor Klhl3 (McCormick et al. 2014). A kidney-specific deletion of the Cul3 gene in the DCT of mice revealed that in contrast to the human mutation, loss of Cul3 decreases blood pressure (McCormick et al. 2014). The kidneys also showed a marked increase in levels of WNK4, a SPAK regulating kinase that ultimately controls NCC function, which is consistent with the accepted idea that WNK4 is a substrate of Cul3 and Klhl3 (McCormick et al. 2014).

One interesting revelation from this study was the change in subcellular localization of the hypertension-causing Cul3 mutant (Figure 3.2). The localization of wild-type Cul3 is predominantly nuclear, but the mutant Cul3Δ403-459 is mostly cytoplasmic. Normally, Cul3 is recruited to different locations in the cell by BTB proteins (Mathew et al. 2012), which is demonstrated here by the shift of wild-type Cul3 from the nucleus to the cytoplasm in the presence of Klhl3 (Figure 3.2 bottom row) (McCormick et al. 2014). This change in localization of the human Cul3 mutation indicates that the mutant Cul3 may have an altered ability to degrade substrates as a result of its improper localization, resulting in changes to ubiquitination that are substrate-dependent. Our lab has shown that Cul3Δ403-459 can ubiquitinate cyclin E and Klhl3 (McCormick et al. 2014), but others have suggested that Cul3Δ403-459 actually has an impaired ability to degrade the Cul3 substrate RhoA (Ibeawuchi et al. 2015). Future work investigating how Cul3Δ403-459 impacts different substrates will be necessary in order to better determine the
differential regulation of Cul3 substrates that might occur in individuals who carry this mutation.

**Figure 3.2:** *The human Cul3 mutant localizes to the cytoplasm.* HeLa cells were transfected with wild-type (WT) Flag-Cul3, Flag-Cul3Δ403-459, Myc-Klh3, or both Myc-Klh3 and Flag-Cul3 and stained for fluorescent microscopy. Flag or Myc antibodies were used to detect the transfected protein. WT Cul3 is predominantly nuclear (top row) whereas the mutant is mainly localized to the cytoplasm (second row). Myc-Klh3 is a cytoplasmic protein (third row) and is able to recruit WT Flag-Cul3 to the cytoplasm (bottom row). The same pattern can be observed in mDCT kidney cells (McCormick et al. 2014). The experiment shown here has been published (McCormick et al. 2014).
RESULTS:

Identifying proteins that interact with Klhl3

As mutations in the BTB-Kelch protein and putative Cul3 substrate adaptor, Klhl3, have also been shown to cause FHHt (Boyden et al. 2012; Louis-Dit-Picard et al. 2012), it is essential to gain a better understanding of Klhl3 and its role in the cell. One way to do this is to identify potential Klhl3 interacting partners and substrates in the cell. Several yeast two-hybrid screens were performed using full length Klhl3 (Figure 3.3). Klhl3 was cloned into a yeast expression vector and transformed into yeast. The Klhl3-containing colonies were then screened against a cDNA library to identify interacting proteins. Yeast two-hybrid screens are valuable for identifying protein-protein interactions, as they provide an inexpensive \textit{in vivo} system that allows millions of proteins to be screened for interactions with a single ‘bait’ protein of interest. Klhl3 was utilized as ‘bait’ in the appropriate two-hybrid system to identify the possible interacting proteins listed below in Table 3.1.

![Figure 3.3: Klhl3 structure](image)

\textbf{Figure 3.3: Klhl3 structure.} Klhl3 is composed of an N-terminal BTB domain followed by a BACK domain and C-terminal Kelch domain. The BTB domain is responsible for binding Cul3 while the Kelch domain interacts with substrates. The BACK domain connects the BTB and Kelch domains.
Table 3.1: Potential binding partners of KLHL3

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<th>Number of Clones screened:</th>
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<td>KLHL3</td>
<td>Human Testis</td>
<td>1.2x10⁶</td>
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<td>KLHL26</td>
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<tr>
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<td>Human Fetal Liver</td>
<td>3.0x10⁶</td>
<td>Dynamitin (p50)</td>
</tr>
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<td>KLHL3</td>
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<td>Dynamitin (p50)</td>
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Figure 3.4: **P50/dynamitin is a subunit of the dynactin complex.** P50 interacts with the dynein motor. P50 forms the scaffold, which connects the cargo-binding arm (Arp1, green) to the microtubule-binding (p150/Glued, blue) domain. This complex is necessary for transport of endosomes within the cell.
**Dynamitin/p50 binds Klhl3**

One protein of interest that was identified in all three screens was p50/dynamitin. Dynamitin/p50 is a subunit of the dynactin complex, which is required for retrograde transport of endocytic vesicles (Figure 3.4). The p50/dynamitin subunit is of interest as a possible Klhl3 binding partner as p50 overexpression is known to result in trafficking errors and breakdown of the dynein-dynactin complex (Burkhardt et al. 1997), making p50 a good candidate for regulation by the ubiquitin-proteasome system.

P50 was cloned by PCR into a CS+ vector that contained either HA and S epitope tag. These vectors allow the protein of interest to be expressed in mammalian cells and carry an epitope tag for easy detection. Immunofluorescent microscopy reveals that transfected Klhl3 and p50 co-localize in the cytoplasm, which means there is a possibility that a biologically relevant interaction might occur between these two proteins (Figure 3.5). Co-transfection of Klhl3 and p50 followed by immunoprecipitation revealed the ability of the two proteins to interact in mammalian cells (Figure 3.6, lane 1 compared to lane 3). In order to further determine the significance of the interaction between p50 and Klhl3, information was sought in the literature regarding the staining pattern of the endogenous proteins. Endogenous Klhl3 is known to stain in a punctate pattern in 293 cells (Louis-Dit-Picard et al. 2012), which can be characteristic of actin-associated proteins. Endogenous p50 is also known to stain in a similar cytoplasmic pattern (Burkhardt et al. 1997). Taken together, this information suggests that p50 and
Klhl3 interact *in vivo* and it is possible that Klhl3 is normally found associated with the dynactin complex in the cytoplasm.

To further elucidate the interaction that occurs between p50 and Klhl3, a Klhl3 mutant which lacks the BTB (Cul3 binding, refer to figure 3.3) domain was created. This construct was utilized to determine if the kelch domain of Klhl3, which is involved in substrate recognition for the Cul3 ubiquitin ligase complex, is the portion of Klhl3 that is responsible for interacting with p50. This construct also contains the BACK domain, which connects the BTB domain to the Kelch domain. Either wild-type Klhl3 or Klhl3ΔBTB was co-expressed with p50 (Figure 3.6). P50 binds Klhl3ΔBTB and is in fact stabilized in its presence (Figure 3.6, lane 4 compared to lane 2), suggesting that p50 binds to the substrate recognition region (kelch domain) and normally is degraded in a Cul3-dependent fashion. This result indicates that Klhl3ΔBTB is sequestering p50 and protecting it from Cul3-mediated degradation, as the kelch domain of Klhl3 retains its high affinity for p50, but without the BTB domain is unable to associate with the Cul3 complex so p50 is not degraded. This result supports the hypothesis that p50 is a substrate of Klhl3 and Cul3.
Figure 3.5: *P50 and Klhl3 co-localize in the cytoplasm.* Ha-p50 and Myc-Klhl3 were transfected into HeLa cells and stained using antibodies to HA and/or Myc, and viewed using immunofluorescent microscopy. The localization of p50 is shown in green, Klhl3 in red, and the nucleus (DAPI) in blue.
**Figure 3.6:** *P50 interacts with the substrate recognition domain of Klhl3.* Upper blot; immunoprecipitation results showing binding between S-tagged p50 and either wild-type Klhl3 (lanes 2 and 3), or Klhl3 lacking its BTB domain (lanes 4 and 5). The middle and lower blots show protein expression in the cell extracts.

**P50 is stabilized in the presence of an FHHt-associated Klhl3 mutant**

Several mutations in Klhl3 have been shown to cause FHHt in humans (Boyden et al. 2012; Louis-Dit-Picard et al. 2012). A dominant mutation that results in hypertension is Klhl3R528H, a point mutation in the kelch domain of Klhl3 (Boyden et al. 2012). In order to determine if this mutant affects p50, the two proteins were co-transfected in HEK293 cells (Figure 3.7 lane 2). Our preliminary results show that p50 appears to be stabilized by Klhl3R528H but not wild-type...
Klhl3 (Figure 3.7 lane 2 compared to lanes 1 and 4). The increased stability is comparable to the stability that results when p50 is expressed with Klhl3 that cannot bind Cul3 (lacks BTB domain, figure 3.7 lane 3). P50 is still able to bind Klhl3R528H (Figure 3.7 bottom row lane 2). Taken together, these results suggest that Klhl3-mediated degradation might be impaired in the presence of Klhl3R528H.

**Figure 3.7:** *A human Klhl3 mutant results in p50 stability.* S-tagged p50 was cotransfected with a human Klhl3 mutation, Klhl3R528H (lane 2), Klhl3 lacking its BTB domain (cannot bind Cul3, lane 3), or wild-type Klhl3 (Lane 4). P50 expression levels in the presence of each Klhl3 mutant are shown (top panel). The middle blot shows Klhl3 expression and the lower blot shows immunoprecipitation results. This is a preliminary result as it has not yet been repeated.
P50 binds Cul3

As previously demonstrated in chapter two, the Cul3 substrate cyclin E is able to bind Cul3 in addition to a BTB substrate adaptor. In order to determine if p50 also possesses this capability, it was checked for binding with a set of Cul3 mutants. It was concluded that, similar to cyclin E, p50 also has the ability to bind wild-type Cul3 in addition to several Cul3 mutants (Figure 3.8, lanes 2-6). In addition to WT Cul3, p50 also binds Cul3Δ51-67, a mutant that does not bind BTB proteins as well as Cul3K712R, an inactive Cul3 that is not modified by Nedd8 (Figure 3.8, lanes 3 and 5). These data suggest that p50 may be able to interact with Cul3 independently of its interaction with Klhl3 and this interaction is not dependent on neddylation of Cul3. P50 was also able to bind the Cul3 mutant that is associated with hypertension, Cul3Δ403-459, indicating that the region of Cul3 that is deleted in this mutant is not involved in the interaction with p50 (Figure 3.8, lane 4), as well as a double mutant which contains both the hypertension mutation and the BTB binding mutation (Figure 3.8, lane 6).
Figure 3.8: P50 binds Cul3 and Cul3 mutants. Upper blot; S-p50 was transfected with several Cul3 mutants. Immunoprecipitation results are shown. The lower lower blots show expression levels of the transfected proteins in the cell lysates.

Cul3 and Klhl3 ubiquitinate p50/dynamitin in vitro

The data presented above suggest that p50/dynamitin can serve as a substrate of Cul3 and Klhl3 in vitro (Figures 3.5 and 3.9). In order to further investigate this hypothesis and determine if p50 is a substrate of the ubiquitin-proteasome pathway, S-tagged p50 was co-transfected with Cul3 and Klhl3 in the presence or absence of the proteasome inhibitor MG132. P50 levels are destabilized when Cul3 and Klhl3 are present, but its levels are restored by MG132 (Figure 3.9 lane 3), demonstrating that the observed reduction in p50 stability in the presence of Klhl3 or Cul3 may be a result of proteasomal degradation. Additionally, co-transfection of Ubr7, a mutant ubiquitin where all lysines have been mutated to arginine rendering it unable to form degradative ubiquitin chains, also increases
steady-state levels of p50 (Figure 3.9 lanes 4 and 5). Ubiquitin laddering of the p50 protein can also be observed, which suggests that it may be a substrate of ubiquitination (Figure 3.9 lanes 4 and 5). Taken together, these data support the hypothesis that p50 is ubiquitinated by Cul3 and Klhl3 in vitro.

Figure 3.9: P50 is ubiquitinated in the presence of Cul3 and Klhl3. S-tagged Dynamitin/p50 expression levels are shown either alone (Top panel, lane 1) or in the presence of Cul3 and Klhl3 (lanes 2 through 5), with or without the ubiquitin mutant Ubr7 (lanes 4 and 5) or the proteasome inhibitor MG132 (lanes 3 and 5).
P50 levels appear to remain unchanged *in vivo*

In order to determine if p50 is targeted for degradation by Cul3 *in vivo*, levels of p50 were compared in Cul3 WT and Cul3 KO 293 cells (Ibeawuchi et al. 2015). P50 levels appear evenly expressed in the wild-type and Cul3 KO cells (Figure 3.10). This result indicates that if p50 is a substrate of Cul3, ubiquitination may serve a purpose besides degradation, or ubiquitination results in degradation but only of a small pool of p50, which could occur either in a cell cycle or localization-dependent manner. Based on the convincing data demonstrating the stabilization of p50 by transfected ubiquitin mutants or the addition of MG132 (Figure 3.9), the latter possibility seems more likely, as p50 would show no change in stability if ubiquitination by Cul3 were non-degradative.

![Figure 3.10: Levels of endogenous p50 are unaffected by loss of Cul3.](image)

*Figure 3.10: Levels of endogenous p50 are unaffected by loss of Cul3.* Levels of endogenous p50 (middle panel) in wild-type and Cul3 KO 293 cells were determined via Western blotting. Levels of actin in the same cells (bottom panel) are shown for comparison.
Transfected p50 is destabilized by Cul3

To further support the hypothesis that p50 is a Cul3 substrate, it was transfected into the Cul3 WT and Cul3 KO 293 cells to check its stability in the presence and absence of Cul3 (refer to the assay in Chapter 2 Figure 2.4). Like cyclin E, a known Cul3 substrate, p50 appears more stable in the KO cells that it does in the wild-type (Figure 3.11, top panel). This result further supports a role of Cul3 in p50 ubiquitination.

Figure 3.11: Transfected p50 is more stable in Cul3 KO cells. Western blot showing levels of transfected S-p50 in Cul3 KO 293 cells (top panel, right lane) in comparison to WT cells (top panel, left lane). The Cul3 substrate cyclin E is shown as a control (bottom panel).

DISCUSSION:

Mutations in Klhl3, a BTB protein and putative substrate adaptor for Cul3, have been shown to cause the inherited form of hypertension FHHt (Boyden et al. 2012; Louis-Dit-Picard et al. 2012). Others have shown that Klhl3 plays a role in the
ubiquitination of the kinases WNK1 and WNK4, which are responsible for phosphorylating the SPAK kinase, which in turn regulates NCC (Yang et al. 2003; Ohta et al. 2013; Shibata et al. 2013; Wakabayashi et al. 2013). In order to determine other cellular processes that Klhl3 might regulate, a yeast two-hybrid screen was performed to identify potential binding partners of Klhl3 (Table 3.1). One protein that this screen identified is p50/dynamitin, a subunit of the dynactin complex, which is involved in vesicle trafficking along the cytoskeleton (Table 3.1 and Figure 3.4). The p50/dynamitin subunit of the dyanctin complex plays a role in dynein-dependent trafficking within the cell. The work presented here demonstrates that p50/dynamitin binds to the substrate recognition (Kelch) domain of Klhl3 and also to Cul3, and can be ubiquitinated in a Cul3 or Klhl3-dependent manner. Taken together, these results suggest that p50 is a likely substrate of the Cul3Klhl3 E3 ubiquitin ligase in cells (Figures 3.6, 3.7, 3.8, and 3.9). These findings suggest the possibility that Klhl3 might function as a regulator of p50 in cells, thereby suggesting a possible role for Klhl3 in endosomal trafficking.

**FUTURE DIRECTIONS:**

Taken together, these results suggest a role for p50/dynamitin as a substrate of Cul3 and Klhl3 in vitro. However, it remains unclear if p50 is ubiquitinated in a Cul3-dependent manner in living cells and organisms. Future experimentation will be necessary in order to determine the role Klhl3 plays in cells and to confirm the status of p50 as a Cul3 substrate and also show what affects p50 degradation may be having in cells. An experiment that would be illuminating would be to knock
down Klhl3 using siRNA in a cell line where Klhl3 is known to be expressed and measuring if there are any effects on endosomal trafficking and endosome maturation in the Klhl3 knock-down cells. Antibodies to early endosome markers such as EEA1 and late endosome markers such as LAMP-1 would be useful for this experiment (Franken et al. 2013). In order to gather the most convincing results, it is important that Klhl3 is knocked-down for this experiment, as Cul3 has already been shown to affect vesicle trafficking via a different mechanism (Hubner and Peter 2012; Huotari et al. 2012; Gschweitl et al. 2016). An effective siRNA against Klhl3 has been developed for use in this experiment (Figure 3.12)

![Figure 3.12](image)

**Figure 3.12:** An siRNA targeting Klhl3 effectively decreases the expression of transfected Klhl3 protein. Different amounts of an siRNA targeting Klhl3 were transfected into 293 cells with Klhl3. This siRNA can be used in future experiments requiring the knockdown of Klhl3.

**The possibility of aquaporin regulation by Cul3**

One notable anomaly in adult mice that have had Cul3 deleted from their kidneys is depletion of the AQP2 aquaporin protein (McCormick et al. 2014). AQP2, a member of the aquaporin family of water channels, is located on the apical surface of the renal collecting duct, where it functions in the reabsorption of water, helping to maintain blood pressure and ion homeostasis (Takata et al. 2008). Expression of
AQP2 at the apical membrane is highly regulated (Takata et al. 2008). AQP2 is stored in vesicles and is transported to the membrane in a dynein-mediated manner, which is contrary to the usual direction of dynein-mediated transport as a result of microtubules that are able to nucleate near the membrane in this instance (Marples et al. 1998). Translocation of AQP2 to the apical membrane occurs when intracellular cAMP levels are increased upon stimulation by the hormone vasopressin (Klussmann et al. 2001; Olesen and Fenton 2017). In order for translocation to occur, AQP2 must have been previously phosphorylated by PKA (Nedvetsky et al. 2009). There are several possible ways in which Cul3 might be involved in AQP2 regulation: 1) AQP2 could be a substrate of Cul3, 2 and 3) The Cul3 substrates Nrf2 or RhoA, which are believed to regulate AQP2 might be involved, and 4) Cul3 might regulate AQP2 via trafficking. These four possibilities as well as the literature supporting them will be discussed below.

When AQP2 is no longer needed at the membrane, it undergoes clathrin-mediated endocytosis, which is also dynein-dependent (AQP2 recycling is summarized in Figure 3.13) (Takata et al. 2008; Nedvetsky et al. 2009). AQP2 is known to be ubiquitinated on K270, although the E3 ligase responsible is unknown (Lee and Kwon 2009). Research has been conducted in attempt to determine the ubiquitin ligase responsible for AQP2 degradation (Lee et al. 2011). Cul3 was not shown to be associated with AQP2 and it has been suggested that either Cul5 or Nedd4 is responsible for AQP2 ubiquitination (Lee et al. 2011). The ligase involved remains to be determined but ubiquitination of AQP2 is believed to be necessary for its endocytosis and also exocytosis via the formation of a multivesicular body (Lee
and Kwon 2009). It remains a possibility that Cul3 could be involved in this process, but this possibility seems increasingly unlikely as Cul3 was not found associated with AQP2 (Lee et al. 2011).

Figure 3.13: Regulation of AQP2 expression at the membrane. This model illustrates the process by which AQP2 is recycled or degraded within the cells of the nephron. This diagram, has been adapted from a review (Takata et al. 2008).
A possible role for Cul3 in AQP2 regulation in the collecting duct has been recently described via the Keap1-Nrf2 pathway (Suzuki et al. 2017). Nrf2 is a transcription factor involved in the oxidative stress response and it is tightly regulated by the BTB-Kelch protein Keap1 (Wakabayashi et al. 2003). Increased Nrf2 activity during murine development as a result of depleted Keap1 has been shown to result in upregulation of two lectins, Clec4d and Clec4n, that are responsible for the glycosylation of AQP2 (Suzuki et al. 2017). Increased glycosylation of AQP2 results in its increased excretion in exosomes and loss of AQP2 in the collecting duct, resulting in increased urine excretion as a result of the animals’ inability to absorb water and a diabetic phenotype (Suzuki et al. 2017). It has been well demonstrated that Nrf2, a Cul3 substrate, regulates AQP2 via this mechanism, but as Cul3 has many other substrates, it is also possible that other Cul3-based mechanisms are also involved in AQP2 regulation.

A second known regulator of AQP2 expression at the apical membrane is RhoA. RhoA is a member of the Rho family of small GTPases and it is known to be involved in the organization of the actin cytoskeleton (Ridley 2001). RhoA is also a known Cul3 substrate, and depletion of Cul3 has been shown to result in increased formation of actin ‘stress fibers’ in cells (Chen et al 2009). Constitutively active RhoA has been shown to result in the formation of stress fibers and prevent translocation of AQP2 to the apical membrane (Klussmann et al. 2001; Tamma et al. 2001). Therefore, it also remains a possibility that Cul3 regulates AQP2 expression at the membrane at least in part through Cul3-mediated regulation of RhoA. It is possible that increased RhoA activity resulting from loss of Cul3 results in reduced
translocation of AQP2 to the membrane. One way to investigate this hypothesis would be to measure AQP2 abundance in exosomes of either Cul3 KO animals or an exosome-producing cell line where either Cul3 or Klhl3 has been depleted.

Regulation of dynactin-mediated transport by Cul3 provides a fourth possible mechanism that might be utilized by Cul3 to regulate AQP2 abundance. Further investigation into a potential role for Klhl3 in p50/dynamitin regulation may help to shed light on this topic. If p50 is indeed regulated in a Cul3-dependent manner, then impaired trafficking of AQP2-containing endosomes may also be a factor that contributes to the reduction in AQP2 that was observed in Cul3 kidney KO animals. Cul3 is known to be involved in other aspects of vesicle trafficking and endosome maturation (Hubner and Peter 2012; Huotari et al. 2012; Gschweitl et al. 2016), so regulation of p50 may not represent the only point at which Cul3 might regulate AQP2 trafficking. Further experimentation in order to increase understanding of the possible role of Klhl3 in trafficking will be necessary in order to determine if Cul3 is regulating AQP2 via endosomal trafficking. It will be important to utilize Klhl3 knockout cells for this experiment instead of Cul3 knockout cells, as Cul3 is already known to have substrates involved in trafficking processes (Hubner and Peter 2012; Gschweitl et al. 2016), and Klhl3 will have fewer substrates than Cul3, thereby allowing for a clearer determination of which pathways might be involved.

In order to begin to determine which of these four mechanisms are responsible for the loss of AQP2 expression in the kidneys of adult Cul3 Kidney KO animals, there are some simple experiments that should be done. First, as it has
been shown that Nrf2 can result in decreased AQP2 expression, which is likely to occur in the Cul3 Pax8 Cre (Kidney KO) animals, it will be necessary to measure Nrf2 expression both in kidneys of adult animals (WT and kidney KO) and in cells or embryos that are hypomorphic or deleted for Cul3 (Cul3\textsuperscript{flx/flx}) in order to determine if Nrf2 is being regulated by Cul3. As the effects of Nrf2 on AQP2 are a result of Nrf2 levels during embryonic development and not increased Nrf2 levels during adulthood (Suzuki et al. 2017), the results gleaned from experiments with MEFs and floxed (hypomorphic for Cul3 but not Cul3 KO) Cul3 embryos will be the most informative as any increase in Nrf2 will indicate whether or not Cul3 floxed adult animals will suffer from abnormal AQP2 regulation that is Nrf2 dependent. Exosome excretion in Cul3 WT and KO animals, and possibly cell lines, will also be necessary to confirm that the AQP2 is being excreted.

Next, it will be necessary to investigate the contributions that altered levels of RhoA may be having on AQP2 regulation. This can be accomplished by comparing levels of RhoA in WT and Cul3 kidney KO levels by western blot and also immunofluorescence to look for the formation of actin stress fibers, which may be preventing AQP2 from reaching the membrane. Elucidating the possible role of Klhl3 in p50 regulation and endocytic trafficking as was previously discussed will help to determine if Klhl3 plays a role in AQP2 regulation. Lastly, it may be necessary to confirm that Cul3 is not the E3 ligase responsible for AQP2 degradation. As other ligases have already been implicated in this process (Lee and Kwon 2009; Lee et al. 2011), further experimentation may not be necessary.
Other identified proteins that may interact with Klhl3 ATP1β1:

Another protein of interest that was identified in the Klhl3 two-hybrid screens is the beta subunit of the Na+/K+ ATPase, which is expressed in many cell types including the basolateral membrane of the DCT (NCC and ENaC are found on the apical surface). The Na+/K+ ATPase is responsible for maintaining the electrochemical gradient which drives the other renal ion transporters (Hamilton and Devor 2012). ATP1β1 is believed to be ubiquitinated and degraded (Yoshimura et al. 2008).

In addition to its role in renal function, this transporter and its subunits also serve other purposes in different types of cells. ATP1β1 is also associated with cancer, as it plays a role in cell adhesion (Litan and Langhans 2015). Decreased expression of ATP1β1 is associated with poor prognosis in breast cancer patients (Presson et al. 2011). Interestingly, increased expression of TGFβII, which was also identified as a potential Klhl3 binding partner, has also been associated with cancer (Presson et al. 2011). Silencing of ATP1β1 is also associated with clear cell renal carcinoma (Selvakumar et al. 2014).

Interestingly, the Na+/K+ ATPase is also very important in brain and may be associated with Down’s Syndrome (Lubec and Sohn 2003). ATP1β1 has been shown to be downregulated in scrapie-infected mice (Kim et al. 2008) and also has been implicated in brain pathologies in humans (Brignone et al. 2011). ATP1β2 (another subunit of this transporter) has also been demonstrated to be important for interactions between neurons and glial cells (Brignone et al. 2011).
ATP1β1 is a good candidate for a Cul3 and Klhl3 substrate as it has been shown by others that it can be degraded in an ubiquitin-dependent manner (Yoshimura et al. 2008). In order to explore this idea further, ATP1β1 will first need to be cloned into a mammalian expression vector so that it can be checked for binding to Klhl3 and Cul3 as well as used in in vitro ubiquitination assays. As this transporter has been shown to be important in brain as well as kidney, future experiments may seek to investigate the role of Klhl3 or other BTB-Kelch proteins in neuronal development and functioning.

MATERIALS AND METHODS

Cell culture and transfections: Cells (mDCT, HeLa, HEK293) were grown in DMEM supplemented with ten percent FBS and penicillin/streptomycin as well as l-Glutamine. Transfections were conducted using calcium phosphate precipitation. Between 1 and 10 micrograms of DNA was used for each transfection and cells were seeded onto 6cm plates. Cells were harvested between 36 and 48 hours post-transfection.

Yeast two-hybrid screens: The two hybrid screens were performed according to the instructions from the Clontech Matchmaker two-hybrid kit (testis and liver screens) and the Clontech Yeast two Hybrid Gold system (Clontech Cat #630489) for the kidney screen. Full length Klhl3 was cloned into the appropriate bait vector (pGilda for Matchmaker and pGBK47 for Matchmaker Gold). Human testis and fetal liver pB4AD cDNA libraries were purchased from Clontech and screened with the pGilda-Klhl3 bait for the Matchmaker screens. The pGBK47-Klhl3 bait was used in
the Matchmaker Gold screen with a pACT2 human kidney cDNA library (Clontech). The sequences of both Klhl3 bait clones were confirmed by Sanger sequencing prior to transformation. The bait vector was then transformed into the appropriate *Saccharomyces cerevisiae* strain (strain EGY48 for the ‘Matchmaker’ screens and yeast strain Y2HGold for the ‘Matchmaker Gold’ screens) followed by transformation of the library. The bait vectors were transformed using electroporation and all library transformations were completed using a lithium acetate transformation procedure. The efficiency of each library transformation (number of clones screened, Table 3.1) was determined by calculating a library titer. Each library plasmid contains a unique cDNA sequence fused to a sequence coding for a DNA activation domain. When a library plasmid interacts with the bait plasmid (Klhl3), then the colony will grow on selective media and turn blue in the presence of a beta galactosidase indicator. Using this system, three screens were performed using human cDNA libraries; Human Matchmaker Kidney, Human Fetal Liver MATCHMAKER LexA cDNA Library, and Human Testis MATCHMAKER LexA cDNA library (Clontech).

**Immunofluorescence:** Immunofluorescence was performed as previously described in Chapter two as well as previous work from our lab (Cummings et al. 2009).

**Western blots and Immunoprecipitations:** These were performed as previously described in Chapter 2 as well as previous work from our lab (Wimuttisuk et al. 2014).
**Cloning of p50 and Klhl3:** Klhl3 was cloned into the yeast two-hybrid vectors pGilda and pGBK7 (Clontech) via PCR from a mammalian expression vector followed by restriction digests and ligation into the appropriate yeast ‘bait’ vector. P50 was cloned by PCR from a human testis cDNA library and inserted into an HA-tagged CS2+ mammalian expression vector. Later, p50 was also sub-cloned from the HA-tagged vector into an S-tagged expression vector (CS2+) using restriction digests followed by ligation into the new vector.

**REFERENCES**


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CHAPTER 4: Conclusions, Discussion, and Future Directions
The work presented here highlights the multitude of cellular processes that are dependent on Cul3-based ubiquitin ligase complexes. The first chapter summarizes what is known about the Cul3 ubiquitin ligase including its structure, cellular functions, and the diseases in which Cul3 is known to be involved. In the body of this dissertation has contributed to a greater understanding of the cellular processes that are governed by Cul3. Chapter two describes the mechanism utilized by Cul3 to target cyclin E for degradation while also helping to establish a role for Cul3 in cancer, as low molecular weight cyclin E that cannot be degraded by Cul3 is associated with tumorigenesis. Chapter three explains our understanding of Cul3’s role in the kidney where it is involved in ion homeostasis and the maintenance of blood pressure. These findings have led to several new research questions worthy of exploration. This final chapter will help to explain the significance of this work and propose new questions and ideas for future experiments relating to these projects. These projects have led to five new research questions, which will be addressed in this chapter:

1. **What is the nature of the interactions between Cul3, Cdk3, and the LRR protein MUF1?**

2. **How does loss of Cul3 affect breast cancer cells?**

3. **What other posttranslational modifications might affect cyclin E in vivo?**

4. **How do BTB proteins affect the binding of E2 enzymes to Cul3-based complexes?**
5. Do changes in Nrf2 activity affect regulation of AQP2 in Cul3 conditional knockout or Cul3Δ403-459 mice?

RESEARCH QUESTION 1: What is the nature of the interactions between Cul3, Cdk2, and the LRR protein MUF1?

BACKGROUND: LRR proteins and Cul3

Cul3 is a member of the Cullin-RING family of ubiquitin ligases, and as such has many structural similarities to other members of the Cullin family. Examples include modification by the ubiquitin-like protein Nedd8, an N-terminal domain which interacts with substrate adaptors, and a C-terminal domain which binds the RING finger protein, Rbx1, which is believed to be responsible for interacting with the E2 enzyme to facilitate substrate ubiquitination (Petroski and Deshaies 2005). Each cullin interacts with a different set of substrate adaptors in order to recognize substrates (Petroski and Deshaies 2005). For example, Cul1 is known to require two substrate adaptor proteins: the Skp1 linker protein, which binds to both Cul1 and the F-box containing substrate adaptor, for example Fbxw7. Unlike Cul1, Cul3 substrate adaptors are only known to consist of one BTB domain containing protein (Refer to Figure 1.3 in Chapter 1 for a visual comparison) (Xu et al. 2003). For the sake of clarity, it is helpful to note that many of the findings included in this section as background information have been published (Wimuttisuk et al. 2014).

In order to increase our understanding of the Cul3-based complex and identify proteins that might interact with Cul3, a proteomics screen was performed. This screen identified a variety of potential Cul3 interacting proteins including ten proteins which contain leucine-rich repeat (LRR) domains (Wimuttisuk et al. 2014).
The identification of many LRR proteins in the screen led to the hypothesis that they may play a structural role in Cul3-based complexes in a similar manner that the LRR protein Skp1 plays in Cul1-based complexes (Wimuttisuk et al. 2014). The proteins identified are involved in a variety of cellular processes, from extracellular matrix construction to neuronal structure and function. The information that is known about these ten proteins is summarized in the list below:

1. **LRR1/LRRC8B/ TA-LRRP**: This protein is a member of the LRRC8 family. It is expressed in many tissues (Kubota et al. 2004).

2. **LRR2/PRAME family member 8**: Members of the PRAME family are considered to be Cancer testis-antigens based on their expression profiles and can be found in different types of tumors (Wadelin et al. 2010). PRAME-like proteins can be expressed in normal tissues as well (Wadelin et al. 2010).

3. **LRR3/SALM-1**: This protein is found in axons and dendrites as well as the synaptic membrane (Morimura et al. 2006). SALM 1 (Synaptic adhesion like molecule-1) is a member of the SALM/Lrfn family of neuronal LRR proteins. This protein has been shown to be involved in the formation homo and heteromeric complexes (Nam et al. 2011). Another member of the SALM family, SALM5, may be involved in autism as well as schizophrenia (Nam et al. 2011).

4. **LRR4/Caspase recruitment domain protein 7**: This protein contains a caspase recruitment domain (CARD). The CARD domain is a member of the death-fold superfamily, which is involved in the formation of signaling
complexes resulting in the activation of caspases and kinases (Kersse et al. 2011).

5. **LRR5/Fibromodulin**: Fibromodulin is a component of the extracellular matrix. It is expressed in B-cell chronic lymphocytic leukemia and mantle cell lymphoma, but not other blood cancers (Mikaelsson et al. 2005).

6. **LRR6/MHC Class II Transactivator (CIITA)**: LRR6/CIITA regulates the activation of MHC II and also transcriptionally regulates many target genes (Wu et al. 2009). This protein is believed to be degraded by the ubiquitin-proteasome system, but the E3 ligase that may degrade it is unknown (Wu et al. 2009).

7. **LRR7/ Reticulon-4 receptor-like 1 precursor, Nogo66 receptor homolog 2, Nogo66 receptor (NgR) related protein 3**: LRR7 is related to the Nogo-66 receptor, which is involved in axonal growth. However, LRR7 does not bind the same ligands as NgR and its function is unknown (Barton et al. 2003).

8. **LRR8/Erbin(ErB2 interacting protein)**: Erbin is expressed in sciatic nerves (Tao et al. 2009). Erbin is another neuronal LRR protein expressed in neurons. It has been shown to be involved in myelination (Tao et al. 2009).

9. **LRR9/LRRK1 (Leucine rich repeat kinase 1)**: LRRK1 is a homolog to LRRK2, a protein shown to be involved in Parkinson's disease. In rats, LRRK1 is expressed in different parts of the rat body and nervous system throughout development (Westerlund et al. 2008).
**10. LRR10/Densin-180** : Densin 180 is a scaffolding protein found in the postsynaptic densities of neurons (Thalhammer et al. 2009). LRR10/Densin-180 and LRR8/Erbin have many structural similarities.

**BACKGROUND: LRR5 (FMOD), LRR3 (SALM1) and Cul3**

In order to determine which, if any, Cul3-dependent processes these proteins might play a role, they were cloned into mammalian expression vectors for transfection into cells. Two LRR proteins which were of interest are LRR5 (FMOD or fibromodulin) and LRR3 (also called SALM1) as both were found to interact with BTB domain containing proteins (Wimuttisuk et al. 2014). As fibromodulin is a component of the extracellular matrix (ECM), we sought to determine its localization, as Cul3 is not an extracellular protein. Immunofluorescence of HA-tagged LRR5 (FMOD) and Flag-Cul3 shows that the two proteins can co-localize in the cytoplasm, indicating that it is possible for FMOD to play a role inside the cell in addition to its established role in the ECM (Figure 4.1, rows 4 and 5). A second LRR protein, LRR3, can also co-localize with Cul3 in the cytoplasm (Figure 4.1, rows 2 and 3). LRR3 (SALM1) is a neuronal protein that is normally found in neurons and is involved in synapse formation as well as axonal outgrowth (Nam et al. 2011). Cul3 has been associated with neuronal functioning and several neuronal disorders (Codina-Sola et al. 2015; Lin et al. 2015; Wang et al. 2016), so it is reasonable to speculate that Cul3 and LRR3 (SALM1) may serve a function in the synapse that remains unknown. This hypothesis may provide a useful basis for future
experimentation, but has not been further explored. All further information in this section will focus on LRR5 (fibromodulin) and seek to determine the potential interactions that might occur between LRR5 and the Cul3 complex within living cells.

**Figure 4.1:** *Cul3 localizes with LRR proteins.* Transfected Flag-Cul3 co-localizes in the cytoplasm with two transfected HA-tagged LRR-domain containing proteins: LRR3 (L3, second row) and LRR5 (L5, bottom row). Antibodies to Flag (Cul3) or HA (LRR proteins) were used for this experiment.

**BACKGROUND: LRR5 (FMOD) interacts with the BTB protein FAZF**

In addition to their ability to interact with Cul3, LRR3 and LRR5 can interact with BTB-domain containing proteins (Wimuttisuk et al. 2014). To elucidate the mechanism by which the Cul3 complex might interact with LRR5 within the cell, a
A yeast two-hybrid screen was performed where LRR5 (FMOD) was used as bait and screened against a human testis cDNA library in order to identify potential LRR5 interacting partners. The only protein that was discovered in this screen was FAZF, a BTB and zinc-finger domain containing protein (Wimuttisuk et al. 2014). FAZF is a transcriptional repressor that is involved in blood cell development and Fanconi anemia, a blood disorder associated with defects in hematopoiesis and bone marrow failure (Dai et al. 2002).

Structurally, FAZF is closely related to PLZF, another DNA binding protein that contains both a BTB domain and a zinc-finger domain (Hoatlin et al. 1999). Further work by our lab revealed that LRR5 can bind FAZF but not PLZF (Wimuttisuk et al. 2014). One structural difference between FAZF and PLZF that we hypothesize might be responsible for the interaction with LRR5 is the presence of a proline-rich region in FAZF which is located just upstream of the BTB domain (Figure 4.2) (Wimuttisuk et al. 2014). This region is not present in PLZF (Wimuttisuk et al. 2014). Analysis of other BTB-domain containing proteins reveals proline-rich regions near the BTB domain in Mayven (Klh12), RhoBTB3, Ctbr73, SPOP, and actinfilin (Klh17). Other proteins such as PLZF and Klhl3 do not contain this region. We hypothesize that this proline-rich region that is located in some BTB proteins may be involved in interactions with LRR-domain containing proteins and that the LRR proteins are not Cul3 substrates and instead might function as a part of the Cul3 complex, possibly as substrate adaptors (Wimuttisuk et al. 2014). Further work will be necessary to determine if proline-rich regions in BTB proteins play a role in any significant interaction between BTB proteins and LRR domain proteins.
containing proteins. In order to further explore this idea, several BTB proteins that contain proline rich regions were identified by searching the amino acid sequences of several BTB proteins by eye (Figure 4.2). The identification of these proteins leads us to hypothesize that other pairs of BTB and LRR proteins might exist. One possible pair consists of the BTB protein RhoBTB3 and the LRR protein MUF1.

**Figure 4.2:** Proline-rich regions in BTB proteins. Domain structures of several different BTB proteins show BTB domains located near proline-rich regions. Many BTB proteins contain a proline-rich region, which is located near the N-terminal portion of the BTB domain. Proline-rich regions were identified by looking through the amino acid sequence of each protein to find areas where there are at least 3 prolines located within a ten residue region.

**RESULTS:** Exploring the interactions between the LRR protein MUF1, Cul3, and Cdk2

RhoBTB3, a BTB protein with a proline-rich region and two BTB domains (Schenkova et al. 2012), is a member of the RhoBTB family of proteins and also a
Cul3 substrate adaptor (Schenkova et al. 2012; Lu and Pfeffer 2013; Zhang et al. 2015). RhoBTB3 is of interest as a potential interacting partner for LRR proteins as it has been recently demonstrated that RhoBTB family members, including RhoBTB3, can interact with the LRR protein MUF1 (Schenkova et al. 2012). MUF1, also called LRRC41, is a nuclear protein whose function is largely unknown (Schenkova et al. 2012). MUF1 is highly conserved in higher eukaryotes, but has not been the focus of extensive research. Others believe that MUF1 may be a substrate of Cul3 and RhoBTB3 (Schenkova et al. 2012).

MUF1 contains an N-terminal B/C box and is believed to interact with Cul5, and a C-terminal leucine rich repeat (LRR) domain (Kamura et al. 2001). The presence of the B/C box suggests that MUF1 is a Cul5 substrate adaptor, but no substrates of MUF1 are known (Kamura et al. 2001). As RhoBTB family members contain proline-rich regions (Figure 4.2), and have been shown to associate with MUF1, we hypothesized that MUF1 and RhoBTB proteins may interact in a similar way as LRR5 and FAZF. In order to investigate this idea, a FLAG-tagged MUF1 construct was obtained for future experimental use (Kamura et al. 2001).

Two nuclear-localized proteins, cyclin E and MUF1, have been shown to interact with the Golgi-associated RhoBTB3 (Schenkova et al. 2012; Lu and Pfeffer 2013). It has been suggested that both proteins are ubiquitinated by RhoBTB3 at the Golgi, which is peculiar since both proteins are normally localized to the nucleus (Schenkova et al. 2012; Lu and Pfeffer 2013). The amino acid sequences of cyclin E and MUF1 were compared in order to identify any possible similarities between the two proteins (Figure 4.3). Cyclin E contains an MRAIL sequence, which is located
within the cyclin homology domain, and also has the sequence EEIYP further downstream. Both of these sequences are involved in cyclin E’s interaction with Cdk2 (Honda et al. 2005). Interestingly, MUF1 contains the sequence RAIV and EEIP further C-terminal, which resembled the Cdk2 interaction region in cyclin E and suggested that MUF1 might also be able to interact with Cdk2. Additionally, MUF1 also contains three possible Cdk2 phosphorylation sites, which suggest that MUF1 may be a substrate of Cdk2 (Figure 4.3).

**Figure 4.3**: Sequences of MUF1 and cyclin E. MUF1 has a B/C box, which interacts with Cul5, shown in red. The MRAIL and EEYIP sequences in cyclin E, which interact with Cdk2, are shown in yellow. Similar sequences in MUF1 are also shown in yellow. MUF1 also contains several possible Cdk2 phosphorylation sites (green). The regions shown in pink, located between the RAIL and EEYIP sequences, appear to differ between cyclin E and MUF1. The MUF1 LRR domain is shown in blue and the B/C box in red.
MUF1 binds Cdk2

The similarities between MUF1 and cyclin E have led us to wonder if MUF1 might be involved in cell cycle regulation. We hypothesized that MUF1 might bind to Cdk2. Binding between Cdk2 and MUF1 was analyzed via immunoprecipitations of transfected protein (Figure 4.4). The results show that MUF1 binds to Cdk2, indicating that Cdk2 and MUF1 might bind in vivo. Further work will be necessary to determine if and how this binding affects the functioning of the cell.

Levels of MUF1 and cyclin E

There are two possible explanations for the binding between MUF1 and Cdk2: First, it is possible that MUF1 behaves similarly to a cyclin and works with Cdk2 to fulfill a function in the cell, and second, MUF1 may be a substrate of Cdk2. One or both of these situations maybe true. There is a third possibility that can be eliminated: Cdk2 may be a substrate of the Cul5/MUF1 ubiquitin ligase. This is extremely unlikely as Cdk2 is not degraded in a ubiquitin-dependent manner as it is regulated by its cyclin subunits which are rapidly degraded (Felix et al. 1989). Therefore, it is most likely that MUF1 acts as a Cdk2 substrate or binding partner. As cyclin E, a known partner of Cdk2, is an unstable protein, it led us to wonder if cyclin E and MUF1 might compete for Cdk2. We hypothesized that co-transfection of cyclin E and MUF1 may cause changes in the stability of one or both proteins (Figure 4.5).

The results of this experiment show that MUF1 and cyclin E both appear to have diminished levels when co-transfected in the absence of excess Cdk2 (Figure
4.5, lane 2 compared to others). When Cdk2 is co-transfected and all three proteins are present, however, cyclin E seems to show increased stability and MUF1 shows decreased stability (Figure 4.5 lanes 5 and 6). When HA-Cul3 is co-transfected with Flag-MUF1, it appears that Flag-MUF1 levels may be slightly elevated, which sometimes also seems to be the case when HA-Cdk2 and Flag-MUF1 are co-transfected (Figure 4.5 compare lanes 1, 2, and 3). Taken together, these results indicate an interaction between MUF1, cyclin E, Cdk2, and the Cul3 complex maybe taking place which is altering the stability of MUF1 and cyclin E. As MUF1 contains several possible Cdk2 phosphorylation sites (Figure 4.3), it remains a strong possibility that MUF1 may be a Cdk2 substrate. This interaction may be responsible for regulating levels of MUF1 within the cell. Further experimentation will be necessary to determine if this is the case, and if so the nature of any interaction between MUF1, cyclin E, Cdk2, or Cul3 which may occur.
Figure 4.4: *MUF1 binds Cdk2*. 293 cells were transfected with CDK2 in the presence and absence of Flag-MUF1. The upper blot shows the immunoprecipitation results. The lower blots show protein expression in the cell extracts.
Expression levels of transfected MUF1 in the presence of different proteins. Co-transfection and western blotting for Flag-MUF1 or Myc-cyclin was utilized to determine their expression levels in the presence or absence of each other as well as HA-tagged Cul3 or HA-tagged Cdk2. MUF1 is shown in the top panel. Cul3, Myc-cyclin E, and HA-Cdk2 are shown in the three lower blots.

DISCUSSION:

Previous work from our lab has identified LRR-domain proteins as potential components of the Cul3 complex which might serve as part of multi-subunit substrate adaptors, similar to the system employed by Cul1-based ligases (Wimuttisuk et al. 2014). Interactions have been reported between MUF1 and members of the RhoBTB family including RhoBTB3 (Schenkova et al. 2012). It is possible that these interactions might fit within our published model suggesting that
LRR proteins can be involved in the formation of Cul3-based complexes via an interaction with BTB domain proteins (Wimuttisuk et al. 2014). This information has led us to our first hypothesis that MUF1 might interact with the RhoBTB proteins via their proline-rich domains (Figure 4.6 panel A).

Although preliminary, the data presented here regarding interactions between MUF1 and the cyclin-dependent kinase Cdk2 suggest that it may play an important role in the cell cycle and merits further study. The information in the literature which demonstrates the ability of RhoBTB3 and related proteins to bind MUF1 as well as the data presented above demonstrating the ability of MUF1 to bind Cdk2 presents a second hypothesis to describe how MUF1 might interact with Cul3 and possibly cyclin E: MUF1 might be a substrate of Cdk2 (Figure 4.6 panel B). MUF1 is a highly conserved protein in mammals (Kamura et al. 2001), indicating that it might be responsible for an important but as of yet unknown function within the cell.

**Figure 4.6:** Model showing two new hypotheses regarding the potential involvement of MUF1 in regulation of Cul3-based complexes, (A) and/or as a possible substrate of Cdk2 (B). MUF1 interacts with proteins in the RhoBTB family, indicated it might associate with Cul3-based complexes via its B/C box (A). Other LRR proteins associate with Cul3 in this way (Wimuttisuk, 2014). MUF1 contains two putative Cdk2 phosphorylation sites and may be regulated by Cdk2 (B).

There are many potential future directions for this project as it is a novel idea and still in its infancy. Further study of MUF1’s structure and function will require
experiments to investigate the cellular role of MUF1. MUF1 is a poorly described protein and much information about MUF1 remains unknown, including its tertiary structure and topology. As MUF1 is believed to be a Cul5 substrate adaptor (Kamura et al. 2001), but none of its substrates are known, a yeast two-hybrid screen using MUF1 as bait would be an effective way to identify proteins that might interact with MUF1 in vivo. It is possible that a substrate for MUF1 may be identified by this method. Cell cycle proteins such as Cdk2 and cyclin E may be identified in such a screen. Deletion of MUF1 from mammalian cells would be a second useful method to help to determine its function. This could be accomplished using either siRNA or CRISP-R technology. However, before MUF1 can be deleted it will be necessary to obtain or produce an antibody that can detect endogenous MUF1 and determine which cell types produce MUF1 protein.

In order to address the hypothesis regarding the role of MUF1 and Cdk2, it will be important to identify sequences in MUF1 that facilitate binding to Cdk2. Mutants of the RAIV and EEIP sequences in MUF1 may be useful to determine if MUF1 interacts with Cdk2 in a similar manner as cyclin E. Truncation mutants of MUF1 should be cloned in order to identify which region on MUF1 interacts with Cdk2. As MUF1 also contains several putative Cdk2 phosphorylation sites (Figure 4.3, shown in green), these sites should be mutated to alanines in order to determine if MUF1 is a substrate of Cdk2. Mutants can then be transfected into cells to be checked for binding to Cdk2 and also to help determine if any of the mutants have notable phenotypes. This could be accomplished via fluorescent microscopy and also transfection assays similar to those utilized with cyclin E in chapter two.
Our original hypothesis, which first led me to MUF1, was that LRR proteins interact with BTB proteins via a proline-rich region on the BTB protein which is located N-terminally of the BTB domain (Figure 4.2). RhoBTB family members possess such a region and have been shown by others to interact with MUF1 (Schenkova et al. 2012). We currently possess a clone of RhoBTB3 in lab, but in order to further investigate this idea it will be necessary to clone RhoBTB1 and RhoBTB2. After cloning all three proteins, the proline rich region should be mutated to alanines in each protein and the resulting mutants should be checked for binding with MUF1 either via immunoprecipitation or yeast two-hybrid screen. These experiments will help to shed light on MUF1’s cellular role as well as any involvement it might have in processes that are regulated by Cul3.

**RESEARCH QUESTION 2: How does loss of Cul3 affect breast cancer cells?**

Chapter two demonstrated that Cul3 binds cyclin E directly and ubiquitinates its N-terminal domain. This finding is significant given the association between N-terminally truncated cyclin E variants and tumor progression. The next step for this project will be to determine how loss of Cul3 and overexpression of Cul3 affects breast cancer cells to determine how loss of Cul3 might affect cancer progression in humans. Based on the results in chapter two, we hypothesize that breast cancer cells that express LMW cyclin E will have similar amounts of LMW cyclin E protein in cells that contain Cul3 and cells that lack Cul3. In order to test this, human breast cancer cell lines should be edited using the CRISPR Cas-9 system to delete the Cul3 gene in the same manner as it was deleted in the 293 cells that were utilized in
Chapter two (Ibeawuchi et al. 2015). Three human breast cell lines should be chosen for this experiment; one normal, one cancerous and producing LMW cyclin E, and the third cancerous but not producing LMW cyclin E. Once these three cell lines have been produced, they will be valuable tools and can be compared to their counterparts with wild-type levels of Cul3. The breast cancer cell line MDA-MB-157 has been used by others for the study of LMW cyclin E, so these cells would be an ideal choice (Porter et al. 2001). Based on our own observations the breast cancer cell lines, Sk-br-3, MCF7, and MDA-MB-231 might also be useful for these experiments (Cummings and Singer, unpublished results).

**RESEAERCH QUESTION 3: What other modifications may affect cyclin E in vivo?**

Continued work with cyclin E revealed the presence of a band visible in 293 cells and detected with the HE12 cyclin E antibody, which is double the size of the endogenous cyclin E band (Figure 4.7). This band appears to be more prevalent in the Cul3 KO 293 cells (Figure 4.7). Cyclin E is known to undergo several post-translational modifications, which suggests the possibility that this 90 kDa band might result from an unknown modification. One modification that might result in such a large product is modification by the transglutaminase enzyme TG2. TG2 is known to cross-link glutamine residues with lysine residues in the same protein or lysine residues with lysine residues in other proteins which results in large products and has been previously demonstrated to modify pRb, the retinoblastoma protein, upon cellular stress (Oliverio et al. 1997; Boehm et al. 2002; Mishra et al. 2007).
Transglutaminase is believed to play a role in apoptosis and also cancer (Mishra et al. 2007).

The cyclin E product that we observed in the 293 cells is large, so we hypothesized that it could be the result of cyclin E modification by TG2. In order to test this, a sample of Cul3 WT and Cul3 KO 293 cells was harvested in lysis buffer and then incubated with varying amounts of TG2 (Figure 4.8). A western blot for cyclin E reveals that with increasing amounts of TG2, both the endogenous 50kDa and 90kDa cyclin E bands appear to disappear from the gel, whereas levels of endogenous cyclin A remain unchanged (Figure 4.8). This result indicates the possibility that TG is modifying cyclin E but not A. It is possible that the modified cyclin E resulted in a product too large in order to be observed in the ten percent resolving gel that was used for this experiment. In order to determine if cyclin E can be modified in a TG2-dependent manner, this experiment will need to be repeated and the incubated extracts will need to be run on a gradient gel in order to identify any large products that might form. If cyclin E does appear modified in this assay, it will then be necessary to determine if this modification differs between the Cul3 WT and KO cells.

Next, in order to determine if cyclin E can be modified by transfected TG2, either WT or KO 293 cells were transfected with a Myc-tagged human TG2 construct in the presence or absence of the transglutaminase inhibitor dansyl cadaverine (Oliverio et al. 1997). The results of this experiment reveal a 72 kDa band that is detectible with cyclin E antibody, in addition to a smear at the very top of the resolving gel (Figure 4.9). It is possible that this band results from TG2 modification.
of cyclin E, but this experiment will need to be repeated to verify these results. Additionally, the dansyl cadaverine did not appear to completely inhibit TG2 activity in this assay, so it will be necessary to titrate the amount of drug that should be added in order to optimize TG2 inhibition (Figure 4.9 lanes 2 and 6).

The results presented here are preliminary and these experiments will need to be optimized and repeated in order to determine if cyclin E is modified in a transglutaminase-dependent manner. Others have shown the Rb protein is modified by transglutaminase during apoptosis (Boehm et al. 2002), so further experimentation suggests that cyclin E is a transglutaminase substrate, it would be necessary to determine if cyclin E is modified during apoptosis as well.

![Western blot showing endogenous levels of Cul3 (top) and cyclin E (bottom) in WT and KO 293 cells. The cyclin E antibody HE12 was used for this experiment.](image)

**Figure 4.7:** A heavy cyclin E band is present in Cul3 KO 293 cells. Western blot showing endogenous levels of Cul3 (top) and cyclin E (bottom) in WT and KO 293 cells. The cyclin E antibody HE12 was used for this experiment.
Figure 4.8: The heavy cyclin E band disappears after incubation in transglutaminase. 293 lysates were harvested and incubated in different amounts of transglutaminase for an hour followed by blotting for endogenous cyclin E (top) and cyclin A (bottom).
Figure 4.9: *The effects of transfected transglutaminase on cyclin E*. Either WT (lanes 1 through 4) or Cul3 KO (lanes 5 through 8) 293 cells were transfected with a myc-tagged transglutaminase (TG2) construct in the presence or absence of the TG inhibitor dansyl cadaverine. A western blot for endogenous cyclin E is shown.

RESEARCH QUESTION 4: How do BTB proteins affect the binding of E2 enzymes to Cul3-based complexes?

The data presented in chapter two demonstrated that RhoBTB3, which is believed to be the substrate adaptor for cyclin E, can bind the E2 ubiquitin conjugating enzyme UbE2E1 in the absence of Cul3 (Figure 2.14). Additionally, it was also shown that Cul3Δ51-67, which does not bind BTB proteins, cannot interact with UbE2E1 (Figure 2.15). Together, these findings suggest that BTB proteins may be involved in recruitment and binding of E2 enzymes to the Cul3 complex. In order to determine if this is the case, it will be necessary to investigate the interactions between different E2s and BTB proteins.
One method that will be necessary to explore these interactions is to check the binding of other BTB proteins that are known to result in the degradation of substrates, such as Klhl3 and Keap1, to UbE2E1, as it is known to make K48-linked (degradative) ubiquitin chains (Plafker et al. 2009). This experiment can be conducted by co-transfecting and immunoprecipitating the BTBs with the E2 in both WT and Cul3 KO 293 cells as demonstrated with RhoBTB3 and UbE2E1 (Chapter 2, Figure 2.14). This method would allow for quick detection of interactions between BTB and E2 pairs.

A second method to determine interactions between E2s and BTB proteins would be to perform a yeast two-hybrid screen in which each E2 has been cloned into the ‘prey’ library vector and each BTB protein is cloned into the ‘bait’ vector. Once the constructs are completed, they can be transformed into *S. cerevisiae* and checked for interactions using the Matchmaker Gold two-hybrid system (Clontech). This method would be useful as it would be cost-effective and allow for the screening of many potential BTB-E2 pairs simultaneously.

Next, it will be necessary to explore the interactions between wild-type Cul3 and Cul3Δ51-67 with the different E2 enzymes. This can be done easily by using immunoprecipitations to check binding between thee Cul3 constructs and different E2 enzymes. It would be best to complete these experiments using the Cul3 KO 293 cells in order to eliminate the endogenous Cul3 from the binding assays. These experiments will help to further describe the binding between BTB proteins and different E2s as well as helping to identify which E2s might be utilized by Cul3 to facilitate ubiquitination in vivo. This information will help to describe a novel
mechanism by which BTB proteins help to determine the type of ubiquitin chain that is attached to various Cul3 substrates.

**RESEARCH QUESTION 5: Do changes in Nrf2 activity affect regulation of AQP2 in Cul3 conditional knockout or Cul3Δ403-459 mice?**

One well studied Cul3 substrate is Nrf2 which is involved in the oxidative stress response (Kim and Keum 2016). A recent study utilized a mouse model to show that Nrf2 is responsible for regulating AQP2 abundance in the kidney during development and increased Nrf2 activity during embryonic development results in misregulation of AQP2 and diabetes in adult animals (Suzuki et al. 2017). This is of interest as we found that when Cul3 deletion in the kidneys of adult mice resulted in reduced AQP2 levels (McCormick et al. 2014). As discussed in chapter three, the animals in which we deleted Cul3 were adults, so it is unclear if Nrf2 played a role in the decreased AQP2 expression in this instance. In chapter three, possible roles of Cul3 in AQP2 regulation were discussed, including the effects that increased Nrf2 activity during development might have on AQP2 regulation. This section will focus on a new, two-part, research question related to this topic: How does the hypertension-causing Cul3 mutation (Cul3Δ403-459) affect the levels and activity of the Cul3 substrate Nrf2 during kidney development and AQP2 expression during adulthood?

In order to answer this question, it will be necessary to track kidney development and Nrf2 levels in mouse embryos that are either wild-type for Cul3, deleted or hypomorphic for Cul3, or contain the hypertension-associated mutation. Before measuring the effects of the hypertension mutation (Cul3Δ403-459 in
humans), it will be necessary to track normal kidney development and Nrf2 levels in comparison to embryos that are deficient for Cul3. This extra step will be necessary as the work associating the misregulation of Nrf2 in kidney development was done using a Keap1 (BTB protein) mouse model (Suzuki et al. 2017), so no direct effect of Cul3 Nrf2 in embryonic kidneys has yet been demonstrated.

We have shown that Cul3Δ403-459 is capable of binding BTB proteins and ubiquitinating substrates and may be a gain-of function Cul3 mutant (McCormick et al. 2014). However, the effects of Cul3Δ403-459 on individual substrates in vivo remain mostly unknown. The mechanism by which Cul3 and the BTB-Kelch protein Keap1 ubiquitinate Nrf2 has been well described, providing a strong framework for a mechanistic comparison between wild-type Cul3 and Cul3Δ403-459. A Cre-inducible mouse model for the Cul3Δ403-459 (also called Cul3Δ9) has been developed (Agbor et al. 2016). This mouse model could be used to induce expression of Cul3Δ9 in utero to track kidney development in the knock-in mice in comparison to wild-type. Levels of Nrf2 could be measured in order to determine if Cul3-mediated degradation of Nrf2 remains unchanged during development in the Cul3Δ9 animals. AQP2 expression in the adult animals would also need to be quantified. If successful, these experiments could shed light on not only the potential role of Cul3Δ403-459 in AQP2 regulation in humans but also help to describe the ability of Cul3Δ403-459 to ubiquitinate substrates when compared to wild-type Cul3. As previously mentioned, it will be necessary to determine the effects of loss of Cul3 on Nrf2 in mice before conducting determining the effects of the Cul3 Δ403-459 mutation. As Nrf2 is known to be involved in a variety of
pathological conditions including cancer (Harder et al. 2015; Holmstrom et al. 2016; Kim and Keum 2016; Suzuki et al. 2017), these results could help to understand the disease phenotype that is present in individuals carrying the Cul3\(\Delta403-459\) mutation.

**FINAL THOUGHTS**

Cul3-based E3 ligase complexes are responsible for the regulation of a variety of cellular pathways, many of which are known to have profound effects on the proper function of multicellular organisms. Moving forward, it will be necessary to further clarify the role of Cul3 in these processes in order to better understand the effects that Cul3 has on human physiology. Chapter two describes a degron recognized by Cul3, which is located in the N-terminal region of cyclin E. Future work should attempt to locate Cul3 degrons in other substrates in order to determine if those degrons resemble the one found in cyclin E.

**METHODS:**

**Yeast two-hybrid screen:** The yeast two hybrid screen was performed using LRR5 (fibromodulin) as bait with the Matchmaker yeast two hybrid system, the same system used for one of the Klhl3 screens in Chapter three. The human testis library (also used for chapter 3) was used for this screen.

**Immunofluorescence:** Immunofluorescence was performed as previously described in both chapter two and previous work from our lab (Cummings et al. 2009).
Cell culture, transfections, western blots and immunoprecipitations: All cell culture work utilizing HeLa and 293 cell lines was performed as described in Chapters two and three. The Myc-tagged TG construct was a gift from Dr. Robert Sheaff at the University of Tulsa. Dansyl-cadaverine was added to the appropriate cells (Figure 4.9) at a final concentration of 1mM to inhibit TG activity. For cyclin E experiments involving TG2, the monoclonal cyclin E antibody HE12 (Santa Cruz Biotechnology) was utilized.

Transglutaminase incubation: Untransfected WT and Cul3 KO 293 cells were harvested in Lysis buffer and sonicated. Lysates were then incubated for 15 minutes in a 37 degree water bath with varying amounts of commercial, food-grade transgluaminase enzyme at a concentration of 50mg/mL. Following incubation, the lysates were mixed with sample buffer, boiled, and ran on 10 percent polyacrylamide gels for western blotting. Western blots were probed using either the monoclonal cyclin E antibody HE-12 (Santa Cruz Biotechnology) or a polyclonal cyclin A antibody (H-432, Santa Cruz Biotechnology). We would like to thank Dr. Robert Sheaff at University of Tulsa for the TG enzyme and protocol.

REFERENCES


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