IDENTIFICATION OF SUMOYLATED PROTEINS AND INVESTIGATION OF PROTEIN UBIQUITINATION IN THE NF-κB PATHWAY

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IDENTIFICATION OF SUMOYLATED PROTEINS AND INVESTIGATION OF PROTEIN UBIQUITINATION IN THE NF-κB PATHWAY

DISSERTATION

A dissertation submitted in partial fulfillment of the requirements for the degree of Doctor of Philosophy in the College of Medicine at the University of Kentucky

By

Xiaoyan Liu
Lexington, Kentucky
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Lexington, Kentucky
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SUMOylation and ubiquitination are important post-translational modifications. While ubiquitination is well known for targeting proteins for degradation, SUMOylation often regulates the intracellular localization of substrates. In the first project of this dissertation, we developed proteomic strategies to identify novel SUMOylated proteins in mammalian cells. In the second project, we investigated the regulation of protein ubiquitination in the NF-κB signaling pathway in the context of Paget’s disease of bone (PDB).

Identification of SUMOylated proteins has been a challenge because of low abundance of SUMOylation substrates. Here, we utilized a mass spectrometry (MS)-based proteomic approach to identify novel SUMOylated proteins in mammalian cells. Seventy-four unique proteins were commonly identified in the collection of four SUMO-1 plasmids, thus considered candidate SUMOylated proteins. Many of these proteins are associated with the nucleus. The results were validated by confirming SUMOylation of a novel substrate Drebrin and a well known substrate Ran-GAP1. Furthermore, the potential SUMOylation sites in Drebrin have been identified and confirmed using site-directed mutagenesis.

PDB is a disorder characterized by increased bone turnover containing hyperactive osteoclasts. Mutations in Sequestosome 1 (p62) are associated with 40% of familial PDB. P62 is a scaffold protein and plays a critical role in regulating ubiquitination of TRAF family signaling molecules and mediating the activation of NF-κB by RANK and TNFα ligands. P62 also plays a critical role in shuttling substrates for autophagic degradation. The objective of this project is to determine the effects of PDB-associated p62 mutants on NF-κB signaling and autophagy. We compared the effect of wild-type (WT) p62 and PDB mutations (A381V, M404V and P392L) on the TNFα-induced NF-κB signaling using an NF-κB luciferase assay. Our results show that these p62 mutations increased the NF-κB signaling. In addition, we found that the PDB mutations did not change the interaction between p62 and the autophagy marker protein LC3. In summary, the PDB mutations in p62 are likely gain-of-function mutations that
can increase NF-κB signaling and potentially contribute to disease progression. Based on the results, we proposed a model to speculate the synergetic role of p62 PDB mutant on NF-κB signaling and autophagy.

KEYWORDS: SUMOylation; ubiquitination; Paget’s disease of bone (PDB); p62; NF-κB signaling
IDENTIFICATION OF SUMOYLATED PROTEINS AND INVESTIGATION OF PROTEIN UBIQUITINATION IN THE NF-κB PATHWAY

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05/12/2012
To my dear parents Xingfu Liu and Cui-E Zhan
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Chapter 1. Background and introduction

Dissertation overview

SUMOylation and ubiquitination are two important post-translational modifications [1, 2]. This dissertation consists of two parts. In the first part (Chapter 2), we developed a relatively simple proteomic method to identify SUMOylated proteins in mammalian cells. We are also the first group to identify and validate that an actin-binding protein, Drebrin [3], could be SUMOylated. The second part (Chapter 3 and 4) of this dissertation is related to “ubiquitination” for the following reasons. Firstly, p62 is an ubiquitin-binding protein [4-6]. Secondly, most of the PDB-associated p62 mutations are in the ubiquitin-associated (UBA) domain [7]. Thirdly, in Chapter 3, we proposed that p62 PDB mutants increase the NF-κB signaling through increasing TRAF6 polyubiquitination. Fourthly, in Chapter 4, we studied the effect of p62 PDB mutants in autophagy. Autophagy is involved in degradation of polyubiquitinated proteins [8].

Currently, little is known about cellular consequences of PDB-associated p62 mutants currently [9]. To fill in this gap, we focused on studying the cellular consequences of PDB-associated p62 mutants on the NF-κB signaling and autophagy. We found that p62 PDB mutants increased TNFα-induced NF-κB signaling, but not through TRAF6 polyubiquitination (Chapter 3). Additionally, we showed that PDB mutants did not change the interaction between p62 and the autophagy marker protein LC3 (Chapter 4). Finally, an integrated model for the role of PDB mutant p62 in NF-κB signaling and autophagy is proposed (Fig. 5.1).
SUMOylation

**Discovery of SUMO**

The small ubiquitin-like protein modifier (SUMO) was firstly discovered as reversible post-translational modification by several groups in the middle 1990s [10-12]. The first SUMOylated protein identified is Ran-GTPase-activating protein 1 (Ran-GAP1), which had been implicated in both nuclear transport and the control of mitosis [11, 12]. In 1997, the researchers found that cells contain two forms of RanGAP1, 70 kDa and 90 kDa [11]. Further analysis showed that the larger form contained a 97 amino acid protein, which is similar to ubiquitin in its shape, known as SUMO [12, 13].

Generally, three major SUMO paralogues, SUMO-1, SUMO-2 and SUMO-3 are expressed in cells [14]. It is not certain whether SUMO-4 protein is expressed in cells although a gene encoding SUMO-4 has been reported [15]. SUMO-2 and SUMO-3 are often referred to as SUMO-2/3 because they share 98% sequence similarity. However, SUMO-2/3 and SUMO-1 have only approximately 50% sequence identity [10, 16]. Human small ubiquitin-like modifier 1 (hSUMO-1) is a protein of 101 amino acids, similar to ubiquitin in 3D structure, even though they only share 17% homology at the amino acid level [17-19]. The SUMO-1 precursor has to be cleaved by SUMO-specific proteases to expose a C-terminal glycine-glycine (GG) functional group for subsequent SUMO activation and conjugation [1].

SUMOylation often regulates protein intracellular localization, protein-protein interactions or transcription regulator activity [2, 20]. SUMOylation is essential to normal
cellular behavior. Dysregulation of protein SUMOylation has been associated with a number of diseases such as cancer, neurodegenerative disease, viral infection, diabetes and developmental defects [2].

**SUMOylation vs. ubiquitination**

SUMOylation and ubiquitination are two important post-translational modifications in the cells. They share many similarities but they are also different in many aspects [21].

**(1) Similarity**

SUMO and ubiquitin have similar protein size, tertiary structure and a C-terminal di-glycine motif. Both SUMO and ubiquitin target the protein with the help of E1 (activating enzyme), E2 (conjugating enzyme) and E3 ligases. In addition, both SUMO and ubiquitin proteins are synthesized as immature precursors. These precursors are processed by the specific hydrolase for subsequent activation and conjugation [14, 22].

**(2) Difference**

Ubiquitination is well known for targeting substrates for degradation, whereas SUMOylation regulates a substrate’s functions mainly by altering the intracellular localization, protein-protein interaction and transcription factor activity. In addition, the ubiquitin pathway has a large number of E2 s and E3 s, whereas the SUMO pathway only uses a single E2 and a few E3 s [17].

*SUMO conjugation, deSUMOylation and SUMO consensus sequence*
It is well known that ubiquitin conjugation requires E1 ubiquitin-activating enzyme, E2 ubiquitin-conjugating enzyme and E3 ubiquitin ligase [23]. SUMO conjugation is very similar to ubiquitin conjugation (Fig. 1.1). SUMO proteins are synthesized as inactive precursors, which must first undergo a C-terminal cleavage mediated by a family of sentrin/SUMO-specific protease (SENP) enzymes. This cleavage exposes a di-glycine motif, which is available for subsequent activation and conjugation [14]. In each conjugation cycle, SUMO is activated in an ATP-dependent manner by the E1 “activating” enzyme. SUMO is then passed to the active site of the conjugating enzyme Ubc9 (ubiquitin-conjugating 9). Finally, SUMO is covalently attached to lysine residues of the target protein through the isopeptide bond between the terminal glycine residue and the ε-amino group of a lysine residue in the target protein by SUMO E3 ligase [1, 24, 25]. SUMOylation is a highly dynamic process that can be reversed by deconjugating enzymes such as the SENP enzymes [26]

The SUMO-1 consensus sequence is a motif of conserved residues next to the modified lysine residue and is found in many identified SUMO-1 substrates [27]. The sequence is ΨKXE/D, where Ψ is a large hydrophobic residue (such as Val, Ile, Leu, Met, or Phe). K is the lysine to which SUMO-1 is conjugated and X is any amino acid, D is aspartic acid and E is glutamic acid. More than two-thirds of the known substrate proteins have at least one SUMOylation consensus sequence ΨKXE/D [2, 28]. However, SUMOylation can also occur at lysine residues without this consensus motif, such as non-consensus SUMOylation sites. In addition, although not all lysine residues within the ΨKXE/D motif are SUMOylated [1], SUMOylation consensus sequence ΨKXE/D is still generally believed to be helpful for predicting SUMOylation sites.
Classification of SUMOylation substrates

To date, more than 120 mammalian substrate proteins for SUMOylation have been identified [2]. Based on the subcellular localization of identified SUMOylation substrates, they could be classified as nuclear proteins, cytoplasmic proteins and transmembrane proteins. The majority of these substrates are nuclear proteins, indicating that SUMOylation is primarily involved in nuclear functions. However, a growing number of non-nuclear proteins have been identified, suggesting important non-nuclear roles of SUMOylation [29-31].

The nuclear SUMOylation substrates are well studied. These substrates could be further classified into nuclear pore complexes, transcription factors and coregulators, DNA replication and repair proteins, as well as kinetochore and centromere proteins [2, 32-34]. Functions of SUMOylation of these nuclear proteins are summarized in the Fig. 1.2.

A growing number of non-nuclear SUMOylation substrates have been identified, indicating more functions of SUMOylation beyond those related to the nucleus [2]. Ran-GAP1, the first identified SUMOylation substrate is a cytoplasmic protein. SUMOylation is clearly required for targeting Ran-GAP1 to the nuclear pore complex (NPC) [11]. Also, some identified SUMOylation substrates are transmembrane proteins, such as death receptors, Fas and TNFR1. SUMOylation of these receptors inhibits their apoptotic signaling [2, 35]. It is noted that a number of non-nuclear SUMOylation substrates are involved in signal transduction. SUMOylation of these proteins could change the activity, stability, or subcellular distribution of the proteins and eventually alter signaling events.
For example, SUMOylation protects inhibitor of NF-κB (IκB) from ubiquitination and degradation by 26S proteasome [2, 30].

**Functional consequences of SUMOylation**

The major functional consequences of SUMOylation include alteration of protein localization, protein-protein interactions and transcription regulator activity [2, 36].

(1) **Protein localization**

As mentioned earlier, Ran-GAP1 is a cytosolic protein. Only SUMOylated Ran-GAP1 binds Ran-GTP binding protein (RanBP2), which mediates SUMOylated Ran-GAP1 translocation from the cytosol to the nuclear pore [11, 37]. Therefore, SUMOylation is critical for nuclear import of some proteins. In addition, SUMOylation could also target substrate proteins to specific locations within the cytoplasm. For example, DRP1 is a GTPase protein required for mitochondrial fission. SUMOylation of DRP1 facilitates its recruitment from the cytosol to the mitochondrial outer membrane [38].

(2) **Protein-protein interactions**

SUMOylation of Ran-GAP1 is also an example of SUMOylation being involved in protein-protein interactions. As discussed earlier, only SUMOylated Ran-GAP1 can bind RanBP2. It is hypothesized that SUMO may serve as an interaction “hub” that recruits new interacting proteins to the substrate [1, 2]. For another example, it has been demonstrated that SUMOylation of transcription factor Elk1 could recruit the histone deacetylase 2 (HDAC2). This recruitment has been shown to result in decreased histone acetylation of Elk-1-regulated promoters and thus transcriptional repression of Elk-1 target genes [39].
(3) Transcription regulator activity

Most SUMOylation substrates are nuclear proteins. Particularly, the primary nuclear SUMOylation substrates are transcription factors and regulators. In most cases, SUMOylation negatively regulates gene expression by either enhancing the function of transcription repressors or inhibiting the function of transcription activators [34, 40, 41]. However, the opposite occurs occasionally. For example, heat shock factor 1 (HSF1) is SUMOylated in response to stress and HSF1 SUMOylation often leads to activation of its target genes [13, 42].

SUMOylation and disease

As described above, SUMOylation is a dynamic process that could be reversed by deconjugating enzymes such as the SENP enzymes [1, 26]. A delicate balance between SUMOylation and deSUMOylation is essential to normal cell functions (Fig. 1.2). Growing evidence has shown that the loss of this balance in SUMOylation and deSUMOylation can lead to diseases including cancer, diabetes and neurodegenerative diseases such as Alzheimer’s disease, Parkinson’s disease, familial amyotrophic sclerosis (fALS) and Huntington’s disease [2, 14, 43]. For example, a recent study has shown that SUMOylation of amyloid precursor protein (APP) close to the β-secretase cleavage site is associated with a decrease of Aβ aggregates, which is generally believed a probable cause of Alzheimer’s disease [14, 29, 31]. The causative relationships between the deregulation of SUMOylation and pathogeneses of the diseases are still unclear and under active investigation. Studies so far have suggested that SUMO target proteins might be therapeutic targets for treating these diseases [14, 43].
In our study, we used a modified proteomic method to identify SUMOylated proteins in HEK293 cells. We also found that Drebrin is a novel substrate for SUMOylation. Background about Drebrin is described as below.

Discovered, isoforms and domains of Drebrin

Developmentally-regulated brain protein (Drebrin) is an actin-binding protein, involved in the regulation of actin filament organization. Drebrin plays an important role during the formation of neurites and cell protrusions of motile cells [44]. The expression level of Drebrin is very high in the cerebral cortex, hippocampus, thalamus and striatum [45]. Drebrin was originally discovered by Shirao et al. [46].

Drebrin has three isoforms including E1, E2 (embryonic) and A (adult) isoform, which are generated by alternative RNA splicing from a single Drebrin gene [47]. Drebrin E1 and E2 were first identified as developmentally regulated brain proteins by two-dimensional gel electrophoresis in 1985 [46]. Drebrin A was discovered using a monoclonal antibody against Drebrin E in 1986 [48]. A cDNA clone for a common domain of Drebrin E1, E2 and A was first isolated from brains of 10-day chick embryos in 1988 [3, 49]. All three isoforms are strongly expressed in neurons [44]. On SDS-PAGE gels, the molecular weight of Drebrin A is about 125 kDa and the molecular weight of Drebrin E is about 115 kDa [45].

The N-terminal domain of Drebrin is an actin-depolymerizing factor homology (ADF-H) domain which is highly conserved across vertebrate [50]. Also, there is an actin-binding domain close to ADF-H. In the C terminus of Drebrin, there are homer-
binding motifs [44]. Homer proteins are scaffold proteins at the post-synaptic density where they facilitate synaptic signaling and appear to be critical in learning and memory [51].

Drebrin contributes to the formation of filopodia

The formation and maintenance of an appropriate shape is fundamental to cells. It is also important for cells to modulate morphology in response to changing environmental stimuli. The cytoskeleton plays an important role to provide both a rigid scaffold and mechanical forces to move the cell [44]. Also, the cytoskeleton is regulated by many proteins which bind cytoskeletal components such as microtubules and actin filaments. Drebrin is one of these actin-binding protein [44], and growing evidence shows that Drebrin is important for controlling cell shape and function by its interaction with other proteins [44].

The most visualized function of Drebrin is that Drebrin contributes to filopodia formation in neurons and other cell types [52]. In 1992, it was firstly reported that exogenous GFP-Drebrin A accumulated within spines and elongated the length of spine [53-55]. Later, it was reported that overexpression of full length Drebrin, or truncations containing the actin-binding domain, induced the formation of numerous microspikes in fibroblasts and massive spines in cultured hippocampal neurons [44, 56, 57]. Also, it was shown that overexpression of Drebrin E2 in cultured epithelial cells resulted in a phenotype similar to that produced in neurons and fibroblasts [44, 58]. It was also shown that filopodia formation could also be inhibited by a reduced amount of Drebrin [52].
In our study, we used the proteomic method to first show that Drebrin could be SUMOylated. Additionally, we found the potential SUMOylation sites of Drebrin. We also try to investigate the functional consequence of Drebrin SUMOylation.

Next, we will talk about another post-translational modification, ubiquitination, which is similar with SUMOylation described above. Here, we firstly will introduce a disease called Paget’s disease of bone (PDB) because mutation in the gene encoding an ubiquitin-binding protein, p62 is associated with PDB. Then, we will describe the protein p62 in detail. We also will introduce NF-κB signaling, in which ubiquitination of many proteins occur. In the end, we will describe autophagy, which is involved in degradation of polyubiquitinated proteins [8].

**Paget’s disease of bone (PDB)**

**Prevalence and symptoms of PDB**

Paget’s disease of bone is named after Sir James Paget who was a British Surgeon. In 1876, he published a scientific article describing cases of previously unrecognized chronic bone disease, which he called “osteitis deformans”. Over 120 years after Sir Paget’s finding, scientists and clinicians began to make significant progress in understanding the etiology of the condition we now know as Paget’s disease of bone (PDB) [59], which is the second most common bone disease after osteoporosis [60].

PDB is most common in England, Western Europe, and North America. Very few cases have been reported in Asia and Africa [59]. Approximately 3% of individuals aged over 50 years are affected with PDB in Caucasian populations [7, 61]. PDB is not lethal, but a chronic disorder that typically results in deformed bones [9]. The symptoms include
bone pain, susceptibility to pathological fractures, osteoarthritis, headache, deafness and neurological complications [59, 62, 63]. Osteosarcoma often occurs in PDB patients [64]. An elevated level of alkaline phosphatase, bone scans, x-rays help the diagnosis [61]. Currently, the common drug for treating the PDB are bisphosphonates, a class of relatively non-selective compounds that target and induce apoptosis of osteoclasts [7].

**PDB is a disorder of bone remodeling**

Bone mass in human being is controlled by both osteoclasts (bone-resorbing cells) and osteoblasts (bone-forming cells) [65]. The opposing activities of these two cell types ensure that bone is constantly remodeled in a process essential to maintain adult bone structure and function [59].

PDB is characterized by focal areas of increased bone turnover containing enlarged hyperactive osteoclasts [7, 66]. Pagetic lesions contain increased numbers of osteoclasts compared with normal bone, which have increased size and contain more nuclei than normal osteoclasts [7]. The increased osteoclast activity leads subsequently to increases in osteoblast activity [59]. Although bone resorption (osteoclast) initially exceeds formation (osteoblast), bone formation greatly exceeds bone resorption in later stages. Therefore, the overall process of bone formation becomes accelerated and disorganized, ultimately resulting in abnormal bone structure.

**RANK plays an important role in the formation of osteoclast**
Receptor activator of nuclear factor kappa-B ligand (RANKL), also known as TNF-related activation-induced cytokine (TRANCE) or osteoprotegerin ligand (OPGL), is a member of the tumor necrosis factor superfamily. It is most abundantly expressed as a cell surface protein by bone marrow stromal cells [67-69]. In 1998, RANKL (OPGL) was shown to be the main osteoclastogenic cytokine both in vitro and in vivo [67, 70, 71].

Osteoclast precursors are monocyte/macrophages. It was reported that RANKL could transform the macrophages to osteoclasts [67, 70, 71]. RANKL could interact with its receptor RANK (Fig. 1.5) [72]. RANKL and RANK are encoded by Tnfsf11 and Tnfrsf11a genes, respectively [67]. It has been reported that both Tnfsf11-knockout mice (without RANKL) and mice in which Tnfrsf11a has been deleted (RANK –/–) fail to generate osteoclasts [67, 71, 73]. However, Tnfrsf11a knockout mice (RANK –/–) could be rescued by RANK-expressing haematopoietic cells, which suggests that RANK plays an important role in osteoclast formation [67, 73].

**Etiology of PDB: disordered RANKL-induced NF-κB signaling**

It has been generally accepted that disordered osteoclast RANKL-induced NF-κB signaling may be central to disease etiology [9]. RANKL-induced NF-κB signaling plays a role in regulating the transformation of osteoclast to activated osteoclast [67]. Therefore, hyperactivated osteoclasts identified in PDB patient might be due to the up-regulation of RANKL-induced NF-κB signaling [9].

There are other factors that also contribute to PDB. It is suggested that PDB etiology is also involved with slow virus [59]. It has been shown that the infection of osteoclasts with a paramyxovirus is a possible cause of PDB [74]. Exposure to
environmental toxin could be another factor affecting PDB incidence. PDB cases in Lancashire (county of historic origin in the North West of England) identified in a 1974 survey have been linked to the cotton industry. It was proposed that arsenic pesticide from cotton bales might be responsible for the high prevalence of disease [75].

Both viral infection and exposure to environmental toxins such as arsenic may upregulate the expression of SQSTM1 (p62), which is an important protein in RANKL-induced NF-κB signaling pathway [7, 9, 59].

Genetics of PDB: p62 (SQSTM1) mutations

The most common genetic mutations found in classical PDB patients are in the SQSTM1 (sequestosome1) gene, located on chromosome 5 at the PDB3 locus [59, 76]. This gene encodes the SQSTM1 protein (also known as p62), which has diverse functional properties [59, 76]. In osteoclasts, p62 appears to be an important component in the RANKL-induced NF-κB signaling pathway [59, 77]. Mutations in p62 gene are a major cause of PDB, but do not account for all cases of PDB [7]. Mutations in p62 gene have also been associated with familial and sporadic disease in up to 40% of cases [7].

To date, over 20 PDB-associated p62 mutations have been identified. Most of the p62 PDB mutations are either missense or truncating mutations in the ubiquitin-associated (UBA) domain, the C terminus of the p62 protein. A few p62 PDB mutations are outside of UBA domain [9].

Recent studies have supported the idea that p62 PDB mutations including P392L, P384S and K378X, are associated with increased RANKL-induced NF-κB signaling compared with wild-type p62 [9, 78-81]. While p62 mutations are linked to most cases of
PDB, mutations in genes encoding other proteins including VCP and RANK, are linked to PDB-related syndromes [59].

In our study, we are interested in cellular consequences of PDB-associated p62 mutations. Thus, we investigated the impact of these p62 mutations on NF-κB signaling and autophagy.

**P62 (Sequestosome 1)**

*The domain structures of p62*

P62 is also called sequestosome 1 or SQSTM1 [82]. It is a conserved multifunctional protein that is mainly involved in cellular signaling, protein degradation, protein aggregation and apoptosis [5, 83-85]. Plus, p62 is a cellular protein which is found in almost all mammalian cell types [86]. It was identified as a common component of cytoplasmic inclusions in protein aggregation diseases including amyotrophic lateral sclerosis (ALS) [87] and other neurodegenerative diseases [88].

The p62 gene has 8 exons and encodes a protein of 440 amino acids [89]. The diverse functions of p62 could be reflected by its domain structure (Fig. 1.4 A) [90]. Generally, p62 consists of six domains/motifs: Phox and Bem1 (PB1) domain, ZZ-type zinc finger, the SOD1 mutant interaction region (SMIR), TRAF6 binding (TB) motif, the microtubule-associated protein 1 light chain 3B (LC3) interaction region (LIR) and an ubiquitin binding-associated (UBA) domain [5, 83].
(1) **PB1 domain**

The N-terminal Phox and Bem1 (PB1) domains of p62 could form heterodimers with other PB1 domains, and could also form homodimers and homooligomers of p62 [83, 91, 92]. The PB1 domain of p62 interacts with the PB1 domain of a number of proteins including atypical protein kinase C (aPKC), MAPK/ERK kinase 5 (MEK5), extracellular responsive kinase (ERK) and neighbor of BRCA1 gene 1 (NBR1). Particularly, the interaction of p62 and aPKC plays an important role in NF-κB signaling described below [5].

(2) **ZZ-type zinc finger**

The ZZ-type zinc finger mediates the interaction of p62 with receptor-interacting protein kinase 1 (also called RIP, RIP1 or RIPK1) [83, 93]. This interaction also plays an important role in the TNFα-induced NF-κB signaling pathway.

(3) **SMIR motif**

We identified a motif that is essential for the interaction of p62 with mutants of the Cu/Zn superoxide dismutase (SOD1) linked to familial ALS [83]. The SOD1 mutant interaction region (SMIR, residues 178-224) is the actual sequence that interacts with mutant SOD1. In particular, the conserved W184, H190 and the positively charged R183, R186, K187 and K189 residues within the SMIR are critical for the interaction because substitution of these residues with alanine significantly impaired the p62-mutant SOD1 interaction. In addition, oligomerization of p62 via the PB1 domain also plays an indispensable role in the p62-mutant SOD1 interaction [83]. The ubiquitin-independent recognition of misfolded proteins by SMIR is illustrated in Figure 2.

(4) **TB motif**
P62 binds to the TNF receptor-associated factor 6 (TRAF6) through the TRAF6 binding (TB) motif. TRAF6 is an E3 ubiquitin ligase in the RANKL-induced NF-κB signaling pathway. The interaction of p62 with TRAF6 could promote K63 linked polyubiquitination of TRAF6, which could further activate NF-κB [5]. The interaction of p62 with the E3 ubiquitin ligase TRAF6 promotes K63-linked polyubiquitination of TRAF6 and of other substrates such as Trk A and IKKγ [94-96].

(5) LC3 interaction region (LIR)

The microtubule-associated protein 1 light chain 3B (LC3) is a protein essential to autophagosome formation [83, 97]. The LC3 interaction region (LIR) of p62 can directly interact with LC3 [83, 97]. Particularly, one PDB-associated p62 mutation, D335E, is found in this LIR region [98].

(6) Ubiquitin binding-associated (UBA) domain

The C-terminus of ubiquitin binding-associated (UBA) domain of p62 is responsible for ubiquitin binding [82, 99, 100]. It is proposed that p62 interacts with polyubiquitinated proteins through the UBA domain. Once the polyubiquitin chain of a substrate protein binds to the UBA domain of p62, the substrate can be either translocated to the proteasome or autophagosome for degradation (see below).

The role of p62 in protein degradation

P62 plays a critical role in both ubiquitin-proteasome system (UPS) and autophagy, the two major known protein degradation pathways in mammalian cells (Fig. 1.6). P62 was reported to be a shuttling factor to the proteasome [4, 82, 101]. Accumulating evidence suggests that the involvement of p62 in autophagy is likely more important. Moreover, p62 can mediate the cross-talk between the UPS and autophagy. It
was proposed that p62 accumulation after autophagy inhibition could further suppress the clearance of ubiquitinated proteins destined for proteasomal degradation [102].

The p62 protein plays a critical role in autophagy as a cargo receptor. P62 is frequently detected in protein inclusions related to human diseases [88, 103-105]. The depletion of p62 inhibited autophagic degradation of aggregation-prone polyglutamine-expanded huntingtin inclusions while p62 protected cells from cell death induced by polyglutamine-expanded huntingtin [89]. Inhibition of autophagy caused elevated levels of p62 and induced more and larger p62 inclusions [89, 106, 107]. It was found that p62 actually regulates the formation and autophagic removal of protein inclusions [89, 106]. p62 binds directly to LC3 through its “LC3 interaction region” (LIR) [89, 97] that is critical to its ability to shuttle substrates to autophagosomes for degradation [97, 108, 109]. The C-terminal UBA domain can interact with polyubiquitin chains [100]. It was proposed that p62 targets ubiquitinated protein aggregates to autophagy through an interaction between its UBA domain and polyubiquitin [97]. However, our lab found that the UBA domain of p62 was dispensable for the recognition of familial ALS-related mutant SOD1 [83]. Instead, an internal sequence motif, the SMIR plays a critical role in mutant SOD1 recognition, suggesting that p62 might also target protein cargos for autophagic degradation via ubiquitin-independent mechanisms [83].

**PDB-associated p62 mutations and phenotype**

The most common genetic mutations found in classical PDB patients are in the p62 gene, located on chromosome 5 at the PDB3 locus [59, 76]. To date, over 20 PDB-
associated p62 mutations have been identified [9]. Most of the p62 PDB mutations are in
the UBA domain [110]. A few p62 PDB mutations are outside of UBA domain [9].

(1) p62 UBA domain mutations

In 2003, the three-dimensional structure of the p62 UBA domain (residues 387-
436) was determined by protein NMR (Fig. 1.4 B) [99, 111]. Identified p62 UBA domain
mutations include P387L [112], P392L [113, 114], L394X [115], E396X [112], S399P
[116], M404V [115, 117], G411S [115], G425R [115, 117], M404T [116] and others.
Moreover, it was demonstrated that all of these mutations impaired K48-linked
polyubiquitin binding by p62 in vitro [116]. Therefore, it was proposed that the disease
mechanism in PDB involves a common loss of ubiquitin binding of p62. Interestingly,
the mutations found in the UBA domain, P392L, G411S and G425R were also recently
reported in ALS patients [118]. These findings suggest that p62 mutations might
represent a causative or risk factor in ALS too.

(2) p62 non-UBA domain mutations

In recent years, more PDB-associated p62 mutations have been found in the non-
UBA domain. A D335E missense mutation located in the LC3-interacting region (LIR)
of p62, 50 amino acids away from UBA domain was identified [9]. Other non-UBA
mutations include P364S [79], A381V [78, 98], Y383X [98] and others.

In our study, we mainly investigated the effect of PDB-associated p62 mutations
on NF-κB signaling and possible mechanisms. We selected three p62 UBA domain
mutations (P392L, M404V and G411S) and two p62 non-UBA domain mutations (D335E and A381V) in our work.

**NF-κB signaling**

Nuclear factor κB (NF-κB) is a transcription factor found in almost all mammalian cell types [119, 120]. NF-κB is well known for regulation of immune responses and inflammation [119, 121]. Growing studies have shown that NF-κB is also involved in the oncogenesis [121], bone diseases [65] and cell death [122].

Currently, the numerous studies of NF-κB consist of website (www.nf-kb.org), patent and around 25,000 publications [120]. Here, we only focus on introducing RANKL-induced and TNFα-induced NF-κB signaling pathways which are related to our research topic.

**RANKL-induced NF-κB signaling**

RANKL is a cytokine that is highly expressed in bone marrow [67]. RANKL-induced NF-κB activity controls normal osteoclastogenesis and also plays an important role in the bone resorbing function of mature osteoclasts [59, 67, 123]. Therefore, upregulation of RANKL-induced NF-κB signaling could at least in part explain the increase in osteoclastic activity in PDB [59].

The binding of RANKL and RANK receptor induces receptor trimerization and recruitment of TRAF6 to bind RANK receptor [124, 125]. P62 subsequently binds to TRAF6 through its TB motif, and facilitates TRAF6 Lys63-linked polyubiquitination [126], since TRAF6 is an RING domain-containing E3 ligases [127, 128]. In addition,
TRAF6 could catalyze the K63-linked polyubiquitination of TAB1-TAB2-TAK1 complex [129], which activates the IkB kinase (IKK). Moreover, atypical protein kinase C (aPKC) is activated by interaction with p62 through its PB1 domain, and further activates the IKK [95]. Activated IKK further phosphorylates IkB, and IkB will be degraded by 26 S proteasome. Transcription factor NF-κB is then released from IkB and translocates from the cytosol to the nucleus, activating the transcription of target genes related to the osteoclast formation [130]. The pathway is illustrated in Fig. 1.5. In addition, Osteoprotegerin (OPG) negatively regulate RANKL-induced NF-κB signaling by competitive binding of RANKL with RANK receptor [131].

In our study, we studied the impact of PDB-associated p62 mutants on RANKL-induced NF-κB signaling. However, the lack of appropriate cell lines prevented us from further investigation.

**TNFα-induced NF-κB signaling**

Tumor necrosis factor α (TNFα) is an important cytokine involved in inflammation, cellular homeostasis and tumor progression and apoptosis [132, 133]. TNFα-induced NF-κB activation is similar to RANKL-induced NF-κB signaling discussed above, but also has its own characteristics.

TNFα functions through two receptors, TNF-R1 and TNF-R2 [134]. TNF-R2 is exclusively expressed only on endothelial and immune cells. TNF-R1 is universally expressed in many cell types, and has been studied more extensively than TNF-R2 [135]. TNFα binds TNF-R1 and induces receptor trimerization and leading to the recruitment of
the adaptor protein TNF-R1-associated death domain protein (TRADD) which binds to the death domain (DD) of TNF-R1[135]. TRADD protein also recruits TNF receptor-associated factor 2 (TRAF2), a family protein with TRAF6 mentioned above. TRAF2 is an E3 ligase and it could undergo auto-polyubiquitination and ubiquitinates RIP through K63-linked polyubiquitin chains. RIP polyubiquitination binds to TAB2/TAB3 complex, and recruit TAK1, which phosphorylates IKK, leading to the activation of the IKK complex [72]. Moreover, p62 interacts with RIP through its ZZ finger. P62 could also interact with aPKC through PB1 domain, and thereby activate IKK [72]. Activated IKK further phosphorylates IκB, and IκB will be degraded by the 26S proteasome. Transcription factor NF-κB is then released from IκB and translocates from the cytosol to the nucleus, activating the transcription of target genes [135]. TNFα-induced NF-κB signaling is illustrated in the Fig. 1.5.

In our study, we investigated the impact of PDB-associated p62 mutations on TNFα-induced NF-κB signaling. We found that p62 PDB mutants increased TNFα-induced NF-κB signaling compared with WT p62. Additionally, we tried to determine the molecular mechanisms of the role of p62 PDB mutant in signaling.

**Autophagy**

In Greek, “autophagy” means “self-eating” [136]. It is another way of protein degradation, in addition to ubiquitin proteasome system (UPS) [137]. Autophagy is a process for degradation of cellular contents, organelles, misfolded proteins and invading bacteria through the lysosomal machinery [138-143]. It contains several different forms, including macroautophagy, microautophagy and chaperone-mediated autophagy [85].
Autophagy has emerged as a very active area of investigation as it closely regulates many cellular functions. Autophagy is also implicated in many diseases, including alcoholic liver disease, neurodegenerative disease and cancer [138, 144-146].

**Inducers and inhibitors of macroautophagy**

We focus on macroautophagy in our study. The soluble materials and organelles in the cytoplasm are sequested by an isolation membrane (also termed “phagophore”). Autophagosomes are formed by expansion of the isolation membrane. The autophagosome then fuses with the lysosome to become an autophagolysosome (also termed “autolysosome”) where the enclosed substrates are degraded (Fig. 1.7 A) [85]. Currently, several autophagy inducers and inhibitors have been widely used (Fig. 1.7 A) [147]. For example, rapamycin is an inhibitor of the mTOR pathway which negatively regulates autophagy. Therefore, rapamycin is an inducer for autophagy. NH₄Cl and Bafilomycin A inhibit the fusion of lysosome and autophagosome, thereby inhibiting autophagy. Another common strategy to induce autophagy is starvation [148]. The class III phosphatidylinositol 3-kinase (PI3K-III) activates autophagy and 3-MA inhibits PI3K-III. Therefore 3-MA is another inhibitor for autophagy. In contrast, beclin-1 activates PI3K-III. Thus beclin-1 is an inducer for autophagy. Other autophagy inhibitors include E64d and peptatin A, which inhibit the protease activity in the autophagolysosome (Fig. 1.7 A).

**LC3-II is a marker for autophagosome**

LC3 is widely used as an autophagy marker. In yeast, Atg 8 is the homolog of LC3 in human [149, 150]. There are two forms of LC3, LC3-I and LC3-II, in yeast and
mammalian cells. LC3-I is cytosolic, whereas LC3-II is conjugated with phosphatidylethanolamine (PE) and is mainly present in isolation membranes, autophagosomes and much less on autolysosomes. Therefore, LC3-II serves as a marker for autophagosomes. The conversion of LC3-I to LC3-II requires the Atg5-Atg12 complex (Fig. 1.7 B) [151].

It seems that the increase of LC3-II indicates more autophagosome and higher autophagic activity. However, LC3-II is also degraded by autophagy, making it difficult to interpret autophagy activity solely based on LC3-II level. Therefore, lysosomal protease inhibitors (E64d and pepstatin A) and inhibitors for the fusion of lysosome and autophagosomes (NH₄Cl and Bafilomycin A) are commonly used in the studies to help determine whether autophagic activity is truly increased [152]. It is important to compare the amount of LC3-II in the presence and absence of these inhibitors.

Methods for studying autophagy also include counting the number of GFP-LC3 puncta in cells overexpressing GFP-LC3.

**P62 is both a substrate and regulator of autophagy**

P62 is a specific substrate of autophagy [153]. It can bind LC3 on the autophagosome through LIR domain (Fig. 1.4 A) [152]. P62 proteins which have mutations in LIR region are not degraded by autophagy, result in their accumulation followed by inclusion formation [8]. Therefore, it is suggested that p62 is degraded by autophagy through interaction with LC3 directly. In addition, p62 mutations in PB1 domain are defective in oligomerization. Lower autophagic degradation of these p62 mutants indicates that oligomerization of p62 through PB1 are critical for their
degradation by autophagy [8]. Besides, the level of p62 is upregulated in Atg5 -/- MEFs, suggesting that accumulation of p62 could serve as an indicator of autophagy suppression [152]. The steady-state level of p62 has recently been used as a marker of autophagic degradation activity. For instance, an elevated level of p62 would be interpreted as inhibition or failure of autophagic activity [154]. However, this involves the critical assumption that p62 biosynthesis is not itself regulated. It has been reported that p62 can be induced at the transcriptional level by various stresses including oxidative stress [155, 156] or proteasome inhibition [157]. Thus, caution should be exercised when using the p62 level as a marker of autophagic activity.

On the other hand, p62 is also a regulator of autophagy. P62 binds the polyubiquitinated protein aggregates through its UBA domain. P62, which binds polyubiquitinated proteins, could also oligomerize through PB1 domains. It is indicated that the interaction of p62 and LC3 is involved in linking polyubiquitinated protein aggregates to autophagy [8, 106, 158] (Fig. 1.6). Therefore, p62 is not only a substrate for autophagy, but it also regulates the autophagic activity of other proteins.

In our study, we mainly investigated the impact of PDB-associated p62 mutations in autophagy.
**Figure 1.1. SUMOylation conjugation pathway.** SUMO conjugation needs E1 SUMO-activating enzyme, E2 SUMO-conjugating enzyme (Ubc9) and E3 SUMO ligase, which is similar to ubiquitin conjugation. SUMO could also be removed by deconjugating enzymes such as the SENP enzymes. This picture is modified from a review paper written by Wilkinson *et al.* [1].

Figure 1.2. Nuclear SUMOylation substrates and their functions. The well studied nuclear SUMOylation substrates could be further classified into nuclear pore complexes, transcription factors & coregulators, DNA replication & repair proteins and kinetochore & centromere proteins. A delicate balance between SUMOylation and deSUMOylation is essential to normal cell functions. This picture is modified from a review paper written by Zhao [2].
Figure 1.3. PDB is a disorder of bone remodeling. Bone mass in human being is controlled by both osteoclasts (bone-resorbing cells) and osteoblasts (bone-forming cells). The opposing activities of these two cell types ensure bone is constantly remodeled in a process essential for maintaining adult bone structure and function [59]. PDB is characterized by focal areas of increased bone turnover containing enlarged hyperactive osteoclasts [7, 59]. This picture is taken from a review paper written by Layfield (2007) [59]. This picture is used with a license agreement between Xiaoyan Liu and Cambridge University Press, with a license number 2822601263692.
Figure 1.4. Schematic domain structure of p62 and NMR structure of the p62 UBA domain. (A) P62 has different domains which exhibit diverse functions by interacting with a number of key proteins. This picture is modified from a review paper written by Moscat et al. [90]. (B) Surface representation of the p62 UBA domain determined by protein NMR. Several representative p62 PDB mutations are shown here. This picture is taken from a review paper written by Layfield et al. [111]. Fig. 1.4 B is used with a license agreement between Xiaoyan Liu and Springer provided by Copyright Clearance Center, with a license number 2822600493041.
Figure 1.5. RANKL-induced and TNFα-induced NF-κB signaling. Upon RANKL or TNFα stimulation, TNF-R and RANK receptor undergo trimerization and recruit TRAFs to membrane. Briefly, NF-κB signaling is involved in TRAFs polyubiquitination, IKK activation, IκB degradation and NF-κB translocation from cytosol to nucleus and following target genes expressions. This diagram was drawn in light of a number of papers [65, 93-95, 130, 135, 159].
Figure 1.6. P62 is proposed to involve in both ubiquitin proteasome system (UPS) and autophagy pathway. P62 involves in targeting polyubiquitinated proteins for degradation by both UPS and autophagy. This picture is modified from a review paper written by Komatsu et al. [8] with permission. The use of this picture is also permitted by the FEBS Letters.
Figure 1.7. **Autophagy pathway.** (A) The process of autophagy. Autophagy inducers (shown in red) and autophagy inhibitors (shown in blue) are presented here. This picture is modified from Dr. Ping Shi’s dissertation in Dr. Haining Zhu’s lab. (B) The conversion of LC3-I to LC3-II requires the Atg5-Atg12 complex. This picture is modified from a review paper written by Nedelsky *et al.* [151].

(A)

(B)
Chapter 2. Proteomic analysis of SUMOylated proteins in mammalian cells

Introduction

As described in Chapter 1, SUMOylation of proteins are involved in a number of diseases including neurodegenerative diseases, cancer and diabetes [14, 160]. Therefore, it is important to identify the SUMOylation substrates and investigate the functional consequences of SUMOylation of these substrates, which might shed light on finding the therapeutic target for treating these diseases. Identification of SUMOylated proteins has remained a challenge because of the low abundance of SUMOylation substrates, a small portion of SUMOylated proteins, in addition to the high activity of SUMO deconjugating enzymes such as SENP [161]. Quantitative proteomics, using isotope labeling-based methods, have been used to identify SUMOylation substrates [162]. Knuesel et al. [163] showed that the SUMO-1(T95R) mutant can be used for the identification of the SUMOylation site by mass spectrometry in vitro [163]. In light of this study, we introduced a relatively simple proteomic method without isotope labeling for identification of SUMOylated proteins, which had not been previously reported. Furthermore, Knuesel et al. [163] was unable to determine whether this hSUMO-1(T95R) mutant was still functional in vivo. In our current study, we demonstrated whether that this mutant retained its functionality in HEK cells.

Most reported substrates for SUMOylation are nuclear proteins, though a few cytosolic proteins have been shown to be substrates [24, 36, 164]. Given this lack of knowledge about cytosolic SUMOylation targets, we were particularly interested in novel cytosolic protein targets.
In this Chapter, we aim to identify novel SUMOylation substrates in mammalian cells by using our newly developed proteomic method. The most important advantage of our strategy is that, it is an isotope-labeling free method, which is easier compared with isotope-labeling method. Especially, we expect to identify novel cytosolic protein as SUMOylation substrates. For the potential novel SUMOylation substrates identified by mass spectrometry analysis, we should firstly validate SUMOylation of these proteins by other methods, such as immunoprecipitation and Western blotting. Next, we need to find the potential SUMOylation sites of these novel substrates. Finally, we would like to investigate the functional consequences of these novel SUMOylation substrates.

Materials and methods

cDNA cloning of human SUMO-1 gene

Total RNA was extracted from HEK293 cells using Qiagen RNA extraction kit following the manufacturer’s instruction. Human SUMO-1 full-length cDNA (“SUMO-1-FL”) was amplified with the following two primers containing two restriction sites at each end: 5’-GC GGA TCC ATG TCT GAC CAG GAG GCA AAA CC-3’ and 5’-GC GCGGCCGC CTA AAC TGT TGA ATG ACC CCC TCT TTG-3’ using cDNA RT-PCR amplification kit (Invitrogen). Human SUMO-1 cDNA lacking the last four amino acids with GG bases at the C-terminal end, was amplified with the following two primers containing two restriction sites at each end: 5’-GC GGA TCC ATG TCT GAC CAG GAG GCA AAA CC-3’ and 5’-GC GCGGCCGC CTA AAC TGT TGA ATG ACC CCC TCT TTG-3’ using cDNA RT-PCR amplification kit (Invitrogen) (“SUMO-1-GG”). The
amplified PCR products were recovered using a PCR purification kit (Qiagen), digested with HindIII and BamHI and then ligated into the p3xFLAG-CMV-10 expression vector (Sigma, St Louis, MO, USA) digested with HindIII and BamHI. The positive clones containing the correct inserts were sequenced, and were named pCMV-3xFLAG-SUMO-1-FL (“SUMO-1 FL” or “FL”, full length) and pCMV-3xFLAG-SUMO-1-GG (“SUMO1-GG” or “GG”, last four amino acids truncated), respectively.

**Plasmids construction**

SUMO-1 FL plasmid with the T95R mutation was amplified from the pCMV-3xFLAG-SUMO-1-FL construct by PCR using two primers containing the following sequences: 5’-GC AAG CTT ATG TCT GAC CAG GAG GCA AAA CC-3’ and 5’-GC GCGGCCGC CTA AAC TGT TGA ATG ACC CCC TCT TTG-3’. SUMO-1 GG (last four amino acids truncated) with the T95R mutation was amplified from the pCMV-3xFLAG-SUMO-1-GG construct by PCR using two primers containing the following sequences: 5’-GC AAG CTT ATG TCT GAC CAG GAG GCA AAA CC-3’ and 5’-GC GCGGCCGC CTA ACC CCC TCT TTG TTC CTG ATA-3’. The amplified DNA fragments were inserted into the HindIII and NotI sites of the pCMV-3xFLAG-10 vector (Sigma). The positive clones were named “SUMO-1-FL-T95R” (or “FL-T95R”, full length with T95R mutation) and “SUMO-1-GG-T95R” (or “GG-T95R”, last four amino acids truncated, with T95R mutation), respectively.

The original Drebrin construct was a gift from Drs. Tomas Brdickac and Ondrej Hrusaka (Czech Republic) [165]. Drebrin was firstly amplified by PCR by using upper and lower primers GJ764 and GJ765 (Appendix II). The amplified DNA fragments were inserted into the EcoRI and BamHI sites of p3xHA-CMV-10, and the positive clones
containing the correct inserts were sequenced. The positive clones were named p3xHA-
Drebrin (human full-length, WT Drebrin). Drebrin constructs containing various point
mutations were generated using the QuikChange II Site-Directed Mutagenesis Kit
(Stratagene), or Quikchange Multi Mutagenesis Kit (Stratagene) based on p3xHA-
Drebrin (WT). The mutagenic primer sequences are summarized in Appendix II. These
K270/K271R and the Drebrin mutant in which five Lys are mutated to Arg: K185R/K186R/K192R/K270R/K271R (“5-K mutant”).

**Cell culture and transient transfection**

Human embryonic kidney 293 (HEK293) cells were maintained at 37°C with 5%
CO₂ in Dulbecco's modified Eagle's medium (Invitrogen) containing 10% fetal bovine
serum (FBS), 100 units/ml penicillin and 100 µg/ml streptomycin. Chinese hamster ovary
(CHO) cells were cultured in classical liquid media: Hams nutrient mixture F12
( SH3052601;Thermo scientific, HyClone).

For protein identification by mass spectrometry, 50% confluent cells were
transfected with the four different 3xFLAG-SUMO-1 constructs mentioned above, and
the p3xFLAG-CMV-10 control vector using lipofectamine transfection reagent
(Invitrogen) in a 10cm plate format. For Ran-GAP1 SUMOylation verification by FLAG
immunoprecipitation (IP), 70% confluent cells were transfected with FLAG-SUMO-1
(FL) in a 6-well plate format. For Drebrin SUMOylation verification by FLAG and HA
IP, 70% confluent cells were transfected with 3xHA-Drebrin (WT) and the different mutant 3xHA-Drebrin constructs in a 6-well plate format.

In-gel digestion and gel extraction

After FLAG-IP, enriched SUMOylated proteins were subjected to 10% SDS-PAGE. The gel was washed twice with fix buffer (50% methanol and 7% acidic acid), and was then stained with SYPRO Ruby (S-12000; Invitrogen) overnight (Fig. 2.2). The next day, the gel was washed twice with wash buffer (10% methanol and 7% acidic acid). Each lane of gel was cut into seven or eight bands, and these bands were subsequently cut into the small pieces. These gel pieces were then washed three times with 50% ACN, 25mM NH₄HCO₃ (AMBIC) pH 8.0, dried with a SpeedVac and reduced with 10mM DTT/50mM AMBIC at 56°C for 45 minutes. Next, proteins were alkylated with 55mM IAA/50mM AMBIC at room temperature for 30 minutes in the dark. Following the wash with 25mM AMBIC, the gel pieces were dehydrated with 100% ACN and then dried using a SpeedVac for 10 minutes. The gels pieces were then incubated with 10ng/µl trypsin in 25mM AMBIC overnight at 37°C. The resulting peptides were extracted using 200µl 50% ACN and 5% formic acid. The extraction liquid was subsequently transferred into a new 0.5ml low retention tube and concentrated to 20 µl.

Mass spectrometry analysis

The peptides were subjected to Liquid Chromatography/Mass Spectrometry/Mass Spectrometry (LC-MS/MS) analysis, and the electrospray MS/MS data were collected from a Q-Star XL quadruple time-of-flight (TOF) mass spectrometer (ABI/MDS Sciex,
Foster City, CA, USA) using a nano-flow HPLC system (Eksigent, Dublin, CA, USA). For direct infusion electrospray ionisation- mass spectrometry (ESI-MS) analysis, the sample was diluted 10 times with 90% acetonitrile containing 0.1% formic acid and loaded to Au/Pd-coated spray emitter (Proxeon, Odense, Denmark). The electrospray voltage was 2100 V and the mass range of TOF MS was from 350 to 1600 (m/Z). Nano-flow LC-MS/MS was performed by exploiting the nano-HPLC system for sample pick-up and separation, where the desired volume of sample solution was injected by the autosampler, desalted on a trap column (300 µm i.d. x 5 mm; LC Packings), and was then subsequently separated by reverse phase C18 column (75 µm i.d. x 150mm; Vydac) at a flow rate of 200 nL/minutes. The HPLC gradient was linear from 5% to 75% in 55 minutes using mobile phase A (H₂O, 0.1% formic acid) and B (80% acetonitrile, 0.1% formic acid). Data acquisition was performed using information-dependent mode, where each cycle typically consisted of a 1s TOF MS survey from 350 to 1600 (m/z) and two 2s MS/MS scans with mass range of 100-1600 (m/z).

**Protein identification and data analysis**

LC-MS/MS data were subjected to database searches for protein identification using a local MASCOT search engine, and candidate proteins were generated by searching the Swiss-Prot database. LC-MS/MS data were also submitted to the MASCOT server for MS/MS ion search, and the peak lists from the LC-MS/MS spectra were generated by the MASCOT script embedded in the Analyst QS software using the following parameters: no smoothing, charge state determined from the MS scan, precursor ion charge states of 2+ and 3+, centroid MS/MS data, height percentage 50% and merge distance 0.02 Da. The typical parameters used in the MASCOT MS/MS ion
search are: *Homo sapiens*, maximum of three trypsin missed cleavages, “Ubi-GG-Lys” definition, cysteine carbamidomethylation, methionine oxidation, protein N-term Acetylation, a maximum of 100 ppm MS error tolerance and a maximum of 0.5 Da MS/MS error tolerance. For MS/MS ion search, proteins with one peptide ion scoring higher than 20 were considered an unambiguous identification without manual inspection. All other hits were manually verified by confirming the peptide sequences from the MS/MS spectra. Non-specific proteins from empty vector were eliminated from the protein list generated from other four different SUMO-1 samples.

**Gene ontology (GO) database**

The identified common SUMOylated protein list was subjected to the Gene Ontology (GO) database, and these proteins were classified based on their location of the cell. These locations included nucleus, cellular membrane, cytosol, cytoskeleton, chromosome, mitochondrion, extracellular and unannotated location.

**Immunoprecipitation**

Forty-eight hours after transfection, cells were washed with 1x phosphate-buffered saline (PBS) and lysed using 1x radio immunoprecipitation assay (RIPA) buffer (Millipore) supplemented with protease inhibitor cocktail (P-8340; Sigma, St Louis, MO, USA), 0.625 mg/mL N-ethylmaleimide (Sigma), 1 mM sodium o-vanadate (Sigma) and 0.2 mM phenylmethylsulfonyl fluoride (Sigma).
The FLAG immunoprecipitations were performed by anti-FLAG M2 affinity gel (F2426; Sigma) in a final volume of 500 μl containing 1 mg protein extract. The IP samples and the corresponding extracts were subjected to sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE) followed by Western blotting using the following antibodies: anti-Ran-GAP1 (gift from Dr. Kevin Sarge, University of Kentucky), anti-FLAG (A8592; Sigma), anti-actin (sc-1616; Santa Cruz Biotechnology) and anti-HA (mouse, sc-7392; Santa Cruz Biotechnology).

The hemagglutinin (HA) IPs were performed by using 2 μg of a mouse monoclonal anti-HA antibody (mouse, sc-7392; Santa Cruz Biotechnology) and Protein-G Sepharose (17-0618-01; GE Healthcare). The IP samples and the corresponding extracts were then subjected to SDS-PAGE followed by Western blotting using the following antibodies: anti-FLAG (A8592; Sigma), anti-actin (sc-1616; Santa Cruz Biotechnology) and anti-HA (rabbit, sc-805; Santa Cruz Biotechnology).

**Western blotting and quantification**

Nitrocellulose membrane was incubated in blocking solution, 5% milk in Tris-Buffered Saline and Tween 20 (TBST) for one hour. Then the membranes were incubated with primary antibody for more than three hours. After four washes with TBST for five minutes each, the membrane was incubated with the secondary antibody for more than one hour. After four washes again with TBST for five minutes each, proteins of interest were visualized by either normal or dura enhanced chemiluminescent (ECL) substrate (Thermo scientific) for detection of horseradish peroxidase (HRP) enzyme (Thermo scientific). The membrane was covered with the wrapping membrane and an autoradiography film (Denville Scientific) which was exposed to the membrane. The
exposure time varied from one second to 20 minutes depending on the signal intensity. Films were subsequently developed by a Kodak X-OMAT 2000 processor.

Software Image J was used for quantification of Western blotting bands on X-ray films. Since ECL signals of the Western blot were captured on X-ray films which are known to have a narrow linear range of detection, the quantification of the Western blot may be out of linear range for certain experiments. Enhanced chemofluorescence (ECF) substrate and Alkaline Phosphatase (AP)-conjugated secondary antibody are encouraged for use in the future.

**Bioinformatic analysis**

SUMOylation sites were predicted using the online software SUMOsp 2.0 ([http://www.sumosp.biocuckoo.org/prediction.php](http://www.sumosp.biocuckoo.org/prediction.php)). This software was used to analyze all the candidate SUMOylated proteins in the list (Table 2.1). In addition, AlignX Module Vector NTI (Invitrogen) was used to align the predicted SUMOylated region of human (*Homo sapiens*), mouse (*Mus musculus*), rat (*Rattus norvegicus*), rabbit (*Oryctolagus cuniculus*), horse (*Equus caballus*), cattle (*Bos taurus*), african elephant (*Loxodonta africana*), giant panda (*Ailuropoda melanoleuca*), domestic dog (*Canis familiaris*), opossum (*Monodelphis domestica*), chicken (*Gallus gallus*), lizard (*Anolis carolinensis*), african clawed frog (*Xenopus laevis*) and zebrafish (*Danio rerio*) (Fig. 2.7 C).

**Immunostaining and confocal microscopy**
Chinese hamster ovary (CHO) cells at 30%-40% confluency were transfected with HA-Drebrin (WT and mutants) using lipofectamine reagent (Invitrogen) on gelatin-coated coverslips. Twenty-four hours post-transfection, cells were fixed with 4% paraformaldehyde (PFA) at 37 °C for 15 minutes, permeabilized with PBS/0.1% Triton, and blocked with 3% bovine serum albumin (BSA) in PBS for 1 hour. All the primary and secondary antibodies were diluted in 3% BSA/PBS. Cells were first stained with primary antibody HA (mouse, sc-7392; Santa Cruz Biotechnology, 1:300) or Drebrin (ab11068; Abcam, 1:100) for 5.5 hours. The coverslips were then washed with PBS and incubated with 4’,6-diamidino-2-phenylindole (DAPI) (D9542; Sigma, 2mg/ml, 1:2000), Oregon Green 488 phalloidin (Invitrogen, 1:50), secondary antibody Alexa Fluor 594 anti-mouse (A21203; Invitrogen 1:300), or Alexa Fluor 594 anti-rabbit, (A11012; Invitrogen 1:300) for 1 hour. Finally, the coverslips were mounted using Vectashield mounting medium (Vector Laboratories). Fluorescence microscopy was used by a Leica SP5 inverted confocal microscope with a 40X objective.

**Protrusion quantification and statistical analysis**

For each sample, 10 view images with Z-stack were taken by a Leica SP5 inverted confocal microscope with a 40X objective. Each view contained about 20-30 cells. The numbers of protrusions were counted for each cell which has protrusions. The percentage of cells with protrusions was counted. The data were presented as mean with standard deviation (SD) based on these 10 view images. The significant differences in percentage of cells with protrusions between cells expressing WT Drebrin and mutant
Results

Generation of different versions of 3xFLAG-SUMO-1 fusion protein

The arginine residue at the C-terminus of ubiquitin could be recognized and cleaved by trypsin, leaving a diglycine signature peptide for identification of ubiquitination site by mass spectrometry [166, 167]. However, for human SUMO-1, the C-terminal end is “TGG” instead of “RGG”, thus it is impossible to generate the diglycine signature peptide. In addition, the last lysine or arginine residues of SUMO-1 are considerably distant from the C-terminus end, generating a SUMO tag (ELGMEEEDVIEVYQEQTGG) that is too big for detection by tandem mass spectrometry.

To solve this problem, we mutated the Thr95 residue of SUMO-1 to Arg. Thus, if endogenous SENPs in HEK293 cells correctly recognize and process the mutant construct, the last four amino acids of SUMO-1 will be removed. This cleavage of SUMO-1 (“FL-T95R”) would yield a 114.1 Da diglycine signature tag (“GG-tag”) following trypsin digestion, which could be used for identification of SUMOylation site with high confidence. Considering that endogenous SENP might not work efficiently to remove the last four amino acids of SUMO-1, we also removed the last four amino acids manually to generate the “SUMO-1-GG” (“GG”, last four amino acids truncated) and
“SUMO-1-GG-T95R” (“GG-T95R”, last four amino acids truncated, with T95R mutation) (Fig. 2.1).

**Identification and classification of SUMOylated proteins from HEK293 cells overexpressing four different versions of SUMO-1**

SUMOylated proteins were immunoprecipitated from cell lysates of HEK293 cells overexpressing four different FLAG tagged SUMO-1 plasmids including “FL” (full length), “FL-T95R” (full length with T95R mutation), “GG” (last four amino acids truncated) and “GG-T95R” (last four amino acids truncated, with T95R mutation). The enriched SUMOylated proteins were then subjected to in-gel digestion, and the resulting peptides were extracted and subjected to LC-MS/MS analysis. The LC-MS/MS data were subsequently subjected to MASCOT MS/MS ion search.

Proteins with one peptide ion scoring higher than 60 or two peptide ions scoring higher than 30 were considered unambiguous identification without manual inspection. The ion score filter is 20. All other hits were manually verified by confirming the peptide sequences from the MS/MS spectra. Non-specific proteins from control samples were eliminated from all other four samples. The number of proteins identified in cells expressing “FL” SUMO-1 (full length), “FL-T95R” SUMO-1 (full length with T95R mutation), “GG” SUMO-1 (last four amino acids truncated) and “GG-T95R” SUMO-1 (last four amino acids truncated, with T95R mutation) are 129, 213, 217 and 177, respectively (Fig. 2.3 A). There are 74 identified common SUMOylated proteins among these four samples (Table 2.1). In these 74 proteins, two proteins have GG-tag. One is
Ran GTPase-activating protein 1 (Ran-GAP1) (RAGP1_HUMAN), the other is T-complex protein 1 subunit delta (TCP-1-delta) (TCPD_HUMAN) (Table 2.2). Additionally, there are 88 common proteins identified from cells overexpressing “FL” SUMO-1 (full length) and “GG-T95R” SUMO-1 (last four amino acids truncated, with T95R mutation) (Fig. 2.3 B).

As described above, a total of 177 unique proteins were identified from the purified SUMOylated proteins in cells overexpressing “GG-T95R” SUMO-1 (last four amino acids truncated, with T95R mutation) (Fig. 2.3 A). There were 13 proteins identified with GG-tag from cells overexpressing “GG-T95R” SUMO-1 (last four amino acids truncated, with T95R mutation) when cutoff was 20 (Table 2.2).

The above 74 identified common SUMOylated proteins were classified by subcellular location using the Gene Ontology (GO) databases, and these proteins were classified into nucleus, membrane, cytosol, cytoskeleton, chromosome, mitochondrion, extracellular and unannotated (Fig. 2.4). Among these 74 identified common SUMOylated proteins, most protein (51.4%) were located in the nucleus, 28.4% protein were located into membrane, 21.6% protein were located in the cytosol, 16.2% protein were located in the cytoskeleton, 12.2% protein were located in the chromosome, 9.5% protein were located in the mitochondrion, 6.8% protein were extracellular and 6.8% protein were unannotated (Fig. 2.4).

**Validation of Ran-GAP1 SUMOylation by MS/MS spectra and IP**
Ran-GAP1 was the first identified SUMOylated protein [12] and was also in our candidate SUMOylation protein list (Table 2.1). Thus, we have again validated Ran-GAP1 SUMOylation using MS/MS spectra (Fig. 2.5) and IP (Fig. 2.6). The successful validation of Ran-GAP1 could serve as a positive control for our proteomic method.

First, we validated Ran-GAP1 SUMOylation by MS/MS spectra. If Ran-GAP1 is SUMOylated, after trypsin digestion, it will generate a peptide with a GG-tag. Then after fragmentation, it will generate a lysine residue (128.09 Da) with two glycine residue (114.04 Da), a total 242.1379 Da. Indeed, here we found the MS/MS spectra of peptide LLVHMGGLLK*(GG)SEDK derived from digestion of Ran-GAP1. This peptide was fragmented into a, b, c, x, y and z series ions. The molecular weight of y4 ion is 478.1643 and molecular weight of y5 ion is 720.2861. The difference between y4 and y5 ion is 242.1218, which is exactly the molecular weight of lysine and two glycine mentioned above, representing the GG tag generated by trypsin digestion (Fig. 2.5). Therefore, it is shown that Ran-GAP1 is SUMOylated by MS/MS spectra.

Next, HEK293 cells were transfected with “FL” SUMO-1 (full length) or “GG-T95R” SUMO-1 (last four amino acids truncated, with T95R mutation). Forty-eight hours post-transfection, FLAG-SUMO-1 IP was performed, and IP products were subjected to SDS-PAGE. Western blotting was performed using antibodies against Ran-GAP1, FLAG and actin. SUMOylated Ran-GAP1 was shown in IP products from cells overexpressing either “FL” SUMO-1 or “GG-T95R” SUMO-1, but not in the control samples (Fig. 2.6). Therefore, it is again shown that Ran-GAP1 is SUMOylated by IP and Western blotting.
Identification and validation of SUMOylation of a novel substrate named Drebrin

We also used the software SUMOsp 2.0 to predict SUMOylation sites for all the candidate SUMOylation proteins in our list (Table 2.1). Among these 74 common proteins, 40 proteins (54%) have SUMOylation consensus sequences, 61 proteins (82%) have predicted SUMOylation sites including both SUMOylation consensus sites and non-SUMOylation consensus sites (Table 2.1).

Our candidate SUMOylation protein list (Table 2.1) includes a protein called Drebrin. As described in Chapter 1, Drebrin is an actin-binding protein involved in the formation of neurites and cell protrusions [44]. Unlike nuclear proteins that comprise the majority of SUMOylation substrates, Drebrin is a cytosolic actin-binding protein [45, 56]. The SUMOsp2.0 software showed that Drebrin had four predicted SUMOylation sites, in which one is a SUMOylation consensus motif, and the other three are non-consensus type (Table 2.1 and Fig. 2.7 A). We also aligned this predicted SUMOylated region among different species. This region was shown to be highly conserved among vertebrate species, suggesting that this region is functionally important (Fig. 2.7 C). Consistently, a previous report states that the Drebrin protein is highly conserved, especially at the N-terminal (residues 1-315) [44].

We verified Drebrin SUMOylation by both FLAG-SUMO-1 IP and HA-Drebrin IP. In detail, we first verified Drebrin SUMOylation by FLAG-SUMO-1 IP. HEK293 cells were transfected with “FL” SUMO-1 (full length) or “GG-T95R” SUMO-1 (last four amino acids truncated, with T95R mutation). Forty-eight hours post-transfection, FLAG-
SUMO-1 IP was performed and IP products were subjected to SDS-PAGE. Western blotting was performed using antibodies against Drebrin with longer and short exposure time. SUMOylated Drebrin is shown in both cells expressing “FL” SUMO-1 and “GG-T95R” SUMO-1, not in the control cells (Fig. 2.8 A, long exposure), suggesting that Drebrin is SUMOylated by FLAG-SUMO-1 IP. Also, the SUMOylation level of Drebrin is higher in cells expressing “FL” SUMO-1 than “GG-T95R” SUMO-1 (Fig. 2.8 A).

In addition, we verified Drebrin SUMOylation by HA-Drebrin IP. HEK293 cells were transfected with HA-Drebrin (WT) and FLAG-SUMO-1 (full length). Forty-eight hours post-transfection, HA-Drebrin IP was performed and IP products were subjected to SDS-PAGE. Western blotting was performed using antibodies against FLAG, HA and actin. SUMOylated Drebrin was observed in cells expressing both HA-Drebrin and FLAG-SUMO-1, but not in control cells (Fig. 2.8 B), indicating that Drebrin is SUMOylated by HA-Drebrin IP.

**K185, K186, K270 and K271 are potential SUMOylation sites of Drebrin**

We next aimed to find potential SUMOylation sites of Drebrin. According to the SUMOylation consensus sequence ΨKXE/D [1], Drebrin has four predicted SUMOylation sites including one SUMOylation consensus site, AKKE (amino acid 184-187) and three non-consensus sites, which are KKEE (amino acid 185-188), RKEE (amino acid 191-194) and KKSE (amino acid 270-273) (Fig. 2.7 B). These sites are also conserved among a number of different species (Fig. 2.7 C).
By using the site-directed mutagenesis, we generated several Drebrin single mutants through changing the K185, K186, K192, K270 and K271 of Drebrin to R respectively. Also, we generated double mutants including K185R/K186R and K270R/K271R. Additionally, we generated a mutant in which all the five K are mutated to R, K185R/K186R/K192R/K270R/K271R, called “5-K mutant”.

Then, we performed both FLAG-SUMO-1 IP and HA-Drebrin IP in HEK293 cells overexpressing FLAG-SUMO-1 (full length) and a variety of different HA-Drebrin mutants including double mutations K185R/K186R, K270R/K271R, single mutant K192R and “5-K mutant” in which all the five K were mutated to R. IP products were subjected to SDS-PAGE and Western blotting was performed using antibodies against HA, FLAG and actin (Fig. 2.9 A and B). Interestingly, the SUMOylation level of mutant Drebrin (combined mutations K185R/K186R) was much less (50% reduction) than that of WT Drebrin (Fig. 2.9 B, lane 3). Also, the SUMOylation level of mutant Drebrin (combined mutations K270R/K271R) was slightly less (30% reduction) than that of WT Drebrin (Fig. 2.9 B, lane 5).

Therefore, these data suggest that potential Drebrin SUMOylation sites could be K185, K186, K270 and K271. To further clarify which K is SUMOylated, we performed the HA-Drebrin IP using the single Drebrin mutant, our preliminary data showed that no single lysine mutant (K185R, K186R, K270R or K271R) could abolish the SUMOylation level of Drebrin (data not shown). Clarification of this issue requires additional experiments.
Double mutation (K185R/K186R and K270R/K271R) in Drebrin separately did not appear to change protrusion formation

As described in Chapter 1, previous studies have demonstrated that Drebrin plays an important role in the formation of filopodia [53, 54, 57]. Also, we have shown that mutant K185R/K186R and K270R/K271R impair Drebrin SUMOylation compared with WT Drebrin. The next questions would be whether these two mutants also change the formation of filopodia compared with WT Drebrin. In order to address this question, CHO cells were transfected with HA-Drebrin (WT, K185R/K186R or K270R/K271R). Twenty-four hours post-transfection, immunostaining was performed using either HA or Drebrin antibody for staining Drebrin, Oregon Green 488 phalloidin for staining actin and DAPI for staining nucleus. Confocal microscopy was used to observe cellular protrusions (Fig. 2.10 A).

Consistent with the literature [53, 54, 57], we observed obvious cellular protrusions in cells overexpressing Drebrin (WT and mutants) (Fig. 2.10 A). However, after we quantified the percentage of cells with protrusions from 10 view images (each image was selected from five Z-stack images) for each sample, we found that there was no significant change among cells expressing WT Drebrin and Drebrin mutants (Fig. 2.10 B). Therefore, our preliminary results suggested that mutation of these two sites (K185R/K186R and K270R/K271R) in Drebrin separately did not appear to change protrusion formation.
Discussion

An applicable proteomic method for identification of SUMO substrates

Previous proteomic method for identification of SUMO substrates is using isotope-labeling method, which is complex [162]. In this Chapter, we developed a relatively simple proteomic method to identify SUMOylated proteins in HEK293 cells based on previous studies [163, 167]. First, we generated a number of SUMO-1 constructs to facilitate identification of SUMOylated proteins by mass spectrometry. In detail, we mutated the Thr95 of SUMO-1 to Arg (“FL-T95R” SUMO-1, full length, with T95R mutation). Since endogenous SENPs remove the last four amino acids of SUMO-1, it could yield a GG-tag after trypsin digestion and facilitate the identification by mass spectrometry [163, 167]. Considering that SENPs may not work very efficiently, we further removed the last four amino acids manually, and generated “GG-T95R” SUMO-1 (last four amino acids truncated, with T95R mutation). For comparison, we also generated a SUMO-1 plasmid without the last four amino acids but retained Thr 95, called “GG” SUMO-1 (last four amino acids truncated). Together with full length SUMO-1 (“FL”), these four different versions of SUMO-1 are all FLAG tagged (Fig. 2.1), and served for enriching SUMOylated proteins by FLAG-IP. The IP products were subjected to in-gel digestion and mass spectrometry for analysis.

After eliminating the non-specific proteins, 74 common SUMOylated proteins were identified from cells expressing four different SUMO-1 constructs. These 74 proteins are considered as candidate SUMOylated proteins. In this protein list, three proteins were reported before as SUMOylation substrates including Ran-GAP1 [12],
nucleophosmin (NPM) [168] and proliferating cell nuclear antigen (PCNA) [169] (Table 2.1). Also, 16 proteins were found in the articles related to the SUMOylation in PubMed database. In addition, we predicted SUMOylation sites including SUMO consensus sites (ΨKXE/D) and non-consensus sites for all these 74 proteins using software SUMOsp 2.0. The numbers of consensus and non-consensus sites of each protein are shown in Table 2.1. Among these 74 common proteins, 40 proteins (54%) have SUMOylation consensus sequences, and 61 proteins (82%) have predicted SUMOylation sites including SUMOylation consensus sites and non-SUMOylation consensus sites (Table 2.1). Additionally, 51.4% of these 74 identified common SUMOylated proteins are nuclear proteins, consistent with the observations that most identified SUMOylated proteins are located within the nucleus [2]. Moreover, we successfully verified the SUMOylation of Ran-GAP1, the first identified SUMOylation substrate [37] in our system by MS/MS spectra (Fig. 2.5) and IP (Fig. 2.6). The successful validation of Ran-GAP1 could serve as a positive control for our proteomic method. Altogether, we have developed an applicable proteomic method for identification of SUMO substrates.

**A novel substrate for SUMOylation, a cytosolic protein called Drebrin**

Currently, not many cytosolic SUMOylation substrates have been found [24, 36]. To fill this gap, we identified and validated SUMOylation of a cytosolic protein, called Drebrin. In our study, we identified a novel cytosolic SUMO substrate, Drebrin. We verified Drebrin SUMOylation by both FLAG-IP and HA-IP in HEK293 cells overexpressing FLAG-SUMO-1 and HA-Drebrin (Fig. 2.8 A and B). Drebrin has four
predicted SUMOylation sites including one SUMOylation consensus site, AKKE (amino acids 184-187) and three non-consensus sites, which are KKEE (amino acids 185-188), RKEE (amino acids 191-194) and KKSE (amino acids 270-273) (Fig. 2.7 B). These sites are also conserved among a number of different species (Fig. 2.7 C).

In order to find potential SUMOylation sites of Drebrin, we generated several Drebrin single mutants through changing the K185, K186, K192, K270 and K271 to R respectively. Also, we generated double mutants including K185R/K186R and K270R/K271R. Additionally, we generated a mutant in which all the five K are mutated to R, K185R/K186R/K192R/K270R/K271R, called “5-K mutant”. Next, we compared the effect of SUMOylation in cells overexpressing these HA-Drebrin mutants and WT HA-Drebrin by HA-IP. Our data showed that the SUMOylation level of Drebrin K185R/K186R was much less than that of WT Drebrin (Fig. 2.9 B, lane 3), about 50% decrease. Plus, the SUMOylation level of Drebrin K270R/K271R is slightly less than that of WT Drebrin (Fig. 2.9 B, lane 5), about 30% decrease.

We are also curious about the functional consequence of SUMOylated Drebrin. It was reported that overexpressing Drebrin in CHO cells could cause the formation of cellular membrane protrusions [53]. Therefore, we investigated the effect of SUMOylated Drebrin on these CHO cells by overexpressing either WT HA-Drebrin or double mutant HA-Drebrin (K185R/K186R and K270R/K271R). Consistent with the literature [53, 54, 57], we observed obvious membrane protrusions in cells overexpressing Drebrin (WT and mutants). However, after we quantified the percentage of cells with protrusions from 10 view images (each image was selected from five Z-stack images) for each sample, there was no significant change among cells expressing WT Drebrin and Drebrin mutants.
Fig. 2.10 B). Thus, our data suggests that mutation of these two sites (K185R/K186R and K270R/K271R) separately did not appear to change protrusion formation.

**Troubleshooting and further technique development**

Although our proteomic method is applicable as discussed before, several problems still remain. Firstly, among the 177 unique proteins identified from the purified SUMOylated proteins in cells overexpressing SUMO-1 (GG-T95R) (Fig. 2.3 A), there were only 13 protein identified with GG-tag when cutoff was 20 (Table 2.2). Additionally, only two of these 13 proteins were in the 74 common protein list (Table 2.1 and 2.2). Moreover, the evidence of MS/MS spectra for GG-tag was not obvious except for Ran-GAP1 (Fig. 2.5). Therefore, GG-tag might not be easy for identification by current LC-MS/MS settings. It might due to majority identification of the GG-tags are below the current limit of detection. Potential ways to overcome this problem could be increasing the amount of enriched SUMOylated proteins subjected to LC-MS/MS by increasing cultured cells. Also, more advanced mass spectrometry might help to better identify the GG-tag.

Another potential problem of our method is that the FLAG-IP is not very specific for enrichment of SUMOylated proteins. Although proteins in control cells were eliminated, the enriched SUMOylated proteins by FLAG-IP might still be contaminated with proteins containing other post-translational modification, such as ubiquitination, which is very similar to SUMOylation. If so, it is difficult to distinguish SUMOylation
from ubiquitination in the identified proteins containing GG-tags because they both will have GG-tags following tryptic digestion. One way to get around this problem could be using tandem affinity purification (TAP) for better enrichment of SUMOylated proteins.

A third concern is that the “5-K mutant” Drebrin in which all the five Lys are mutated to Arg did not completely abolish SUMOylation compared with WT Drebrin by HA-IP (data not shown). Therefore our data suggest that other Lys outside of the predicted SUMOylation region (Fig. 2.7 B) might be present. Another possibility is that cells have “compensation mechanism”, thus SUMOylation could occur at some random Lys when all the SUMOylated Lys are mutated. In addition, the experiments shown in Fig. 2.9 B were repeated three times. SUMOylation of K185R/K186R mutant Drebrin decreased compared with WT p62 for all three experiments. However, K270R/271R mutant Drebrin did not decrease for all three experiments, suggesting again that SUMOylation sites might be outside of the predicted SUMOylation region. Clarification of this issue requires additional experiments.

Significance of this study and future directions

Based on a previous study [163], we developed a relatively simple proteomic method to identify SUMOylated substrates. We have identified 74 SUMOylated proteins by our method, in which three proteins are reported to be SUMOylation substrates, 16 proteins are related to the SUMOylation substrates, 40 proteins have SUMOylation consensus sequences (ΨKXE/D) and 61 proteins have predicted SUMOylation sites by
software. We also validated the first SUMOylated protein Ran-GAP1 in our system. Therefore, our method is applicable for identifying novel SUMOylation substrates.

In addition, we are the first group to report that an actin-binding protein, Drebrin, is a substrate for SUMOylation. It is interesting that Drebrin is not in the nucleus, and little is known about SUMO substrates located outside of nucleus [24, 36]. These data may expand our knowledge of non-nuclear SUMOylation substrates by studying the functional consequence of Drebrin. In our study, we verified Drebrin SUMOylation by IP and found that K185, K186, K270 and K271 might be Drebrin SUMOylation sites. Also, our data have shown that double mutation (K185R/K186R and K270R/K271R) separately did not appear to change protrusion formation (Fig. 2.10 B). Therefore, we could further explore the functional consequences of Drebrin in the future.

Moreover, in our protein list, there are several proteins that have a relatively high number of predicted SUMOylation sites (highlighted in Table 2.1). These proteins include general transcription factor II-I (four consensus and two non-consensus sites), heterogeneous nuclear ribonucleoproteins C1/C2 (two consensus and eight non-consensus sites), heterogeneous nuclear ribonucleoprotein U (three consensus and five non-consensus sites), heterogeneous nuclear ribonucleoprotein D0 (one consensus and three non-consensus sites), nucleolin (four consensus and four non-consensus sites), poly [ADP-ribose] polymerase 1 (five consensus and 11 non-consensus sites), 60S ribosomal protein L24 (two consensus and two non-consensus sites) and spectrin beta chain, brain 1 (three consensus and 13 non-consensus sites), Ras GTPase-activating-like protein IQGAP1 (four consensus and 11 non-consensus sites), ATP-dependent DNA helicase 2 subunit 1 (two consensus and four non-consensus sites) and myosin-10 (six consensus
and 10 non-consensus sites), another cytosolic protein. Future studies could validate whether these proteins above are SUMOylation substrates by IP and other methods.
Figure 2.1. Schematic diagrams of SUMO-1 constructs in our study. Schematic diagrams of four p3xFLAG-SUMO-1 constructs used in our study including “FL” (full length), “FL-T95R” (full length with T95R mutation), “GG” (last four amino acids truncated) and “GG-T95R” (last four amino acids truncated, with T95R mutation).
Figure 2.2. Sypro Ruby staining SDS-PAGE gel of FLAG-SUMO-1 IP from cells overexpressing four versions of SUMO-1 constructs. HEK293 cells were transfected with four different SUMO-1 constructs including “FL” (full length), “FL-T95R” (full length with T95R mutation), “GG” (last four amino acids truncated) and “GG-T95R” (last four amino acids truncated, with T95R mutation). Forty-eight hours post-transfection, FLAG-SUMO-1 IP was performed and IP products were subjected to SDS-PAGE. The gel was then stained with Sypro Ruby overnight. Each lane was cut into seven or eight bands and the gel was subjected to the in-gel digestion, gel extraction and LC-MS/MS.
Figure 2.3. Number of identified SUMOylated protein from HEK293 cells overexpressing four different versions of SUMO-1 constructs. (A) Numbers of SUMOylated proteins identified with high confidence (ion score filter 20) from HEK293 cell overexpressing four different versions of SUMO-1 constructs including “FL” (full length), “FL-T95R” (full length with T95R mutation), “GG” (last four amino acids truncated) and “GG-T95R” (last four amino acids truncated, with T95R mutation). There are 74 common SUMOylated proteins. Non-specific proteins from control samples were eliminated from identified SUMOylated proteins. (B) Number of SUMOylated proteins identified with high confidence (Ion score filter 20) from HEK293 cell transfected with “FL” (full length) and “GG-T95R” (last four amino acids truncated, with T95R mutation). There are 88 common SUMOylated proteins.

(B)
Figure 2.4. Classification of 74 identified common SUMOylated proteins. The 74 common proteins were classified based on the subcellular locations including nucleus, membrane, cytosol, cytoskeleton, chromosome, mitochondrion, extracellular and unannotated locations by Gene Ontology (GO) databases. The percentage of proteins belonging to each subcellular location is also shown.
Figure 2.5. MS/MS evidence of SUMOylated peptide from Ran-GAP1. MS/MS spectrum of the peptide LLVHMGLLK*(GG)SEDK derived from digestion of Ran GTPase-activating protein 1 (Ran-GAP1). Validation of Ran-GAP1 serves as a positive control for our proteomic method.
Figure 2.6. Validation of SUMOylated Ran-GAP1 by FLAG-SUMO-1 IP followed by Western blotting. HEK293 cells were transfected with “FL” SUMO-1 (full length) or “GG-T95R” SUMO-1 (last four amino acids truncated, with T95R mutation). Forty-eight hours post-transfection, FLAG-SUMO-1 IP was performed and IP products were subjected to SDS-PAGE. Western blotting was performed using antibodies against Ran-GAP1, FLAG and actin.
Figure 2.7. Predicted SUMOylation region of human Drebrin. (A) Predicted SUMOylation region of human Drebrin by software SUMOsp 2.0. (B) The predicted SUMOylation consensus site is highlighted and the predicted non-consensus SUMOylation sites are underlined. The predicted SUMOylated Lys residues are shown in red. (C) The sequence alignment of predicted SUMOylation region of Drebrin among different species including human (Homo sapiens), mouse (Mus musculus), rat (Rattus norvegicus), rabbit (Oryctolagus cuniculus), horse (Equus caballus), cattle (Bos taurus), african elephant (Loxodonta africana), giant panda (Ailuropoda melanoleuca), domestic dog (Canis familiaris), opossum (Monodelphis domestica), chicken (Gallus gallus), lizard (Anolis carolinensis), african clawed frog (Xenopus laevis) and zebrafish (Danio rerio).
Figure 2.8. Validation of SUMOylated Drebrin by FLAG-SUMO-1 IP and HA-Drebrin IP followed by Western blotting. (A) Validation of SUMOylated Drebrin by FLAG-SUMO1 IP. HEK293 cells were transfected with “FL” SUMO-1 (full length) or “GG-T95R” SUMO-1 (last four amino acids truncated, with T95R mutation). Forty-eight hours post-transfection, FLAG-SUMO-1 IP was performed and IP products were subjected to SDS-PAGE. Western blotting was performed using antibodies against Drebrin with longer and short exposure time. (B) Validation of SUMOylated Drebrin by HA-Drebrin IP. HEK293 cells were transfected with FLAG-SUMO-1 (full length) and HA-Drebrin (WT). Forty-eight hours post-transfection, HA-Drebrin IP was performed and IP products were subjected to SDS-PAGE. Western blotting was performed using antibodies against FLAG, HA and actin.
(B)

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**Blot**

**HA-Drebrin IP**

- **kDa**
  - 250
  - 130
  - 130
  - 42

**Extracts**

- **kDa**
  - 250
  - 130
  - 42

**Proteins**

- **FLAG (SUMO-1)**
- **HA (Drebrin)**
- **Actin**
Figure 2.9. Identification of Drebrin SUMOylation sites by FLAG-SUMO-1 IP and HA-Drebrin IP. (A) Identification of Drebrin SUMOylation sites by FLAG-SUMO-1 IP. HEK293 cells were transfected with full length FLAG-SUMO-1 and HA-Drebrin (WT) or mutant HA-Drebrin including K185R/K186R, K192R, K270R/K271R and K185R/K186R/K192R/K270R/K271R (“5-K mutant”). FLAG-SUMO-1 IP was performed and IP products were subjected to SDS-PAGE. Western blotting was performed using antibodies against HA, FLAG and actin. (B) Identification of Drebrin SUMOylation sites by HA-Drebrin IP. HEK293 cells were transfected with full length FLAG-SUMO-1 and HA-Drebrin (WT) or mutant HA-Drebrin including K185R/K186R, K192R and K270R/K271R. HA-Drebrin IP was performed and IP products were subjected to SDS-PAGE. Western blotting was performed using antibodies against HA, FLAG and actin. Quantification of SUMOylated Drebrin/Drebrin was done by Image J software. The relative amount of SUMOylated Drebrin was obtained by normalization of SUMOylated Drebrin to HA-Drebrin and this number for WT Drebrin was set to 1. All other mutants were normalized accordingly. The quantification data is shown in red.
(B)

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**HA-Drebrin IP**

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**Blot**

- FLAG (SUMO-1)

**Re-blot**

- HA(Drebrin)

**SUMOylated Drebrin/Drebrin**

**Extracts**

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- HA(Drebrin)
- Actin
Figure 2.10. Immunostaining of Drebrin and actin filament in CHO cells. CHO cells were transfected with WT Drebrin or mutant Drebrin (K185R/K186R or K270R/K271R) or empty HA vector. Non-transfected CHO cells also were used as control cells. Twenty-four hours post-transfection, cells were fixed with 4% PFA and permeabilized with 0.1% Triton. Primary antibody HA (1:300) or Drebrin (1:50) and secondary antibody Alexa Fluro 594 (mouse or rabbit) (red, 1:300), as well as DAPI (blue, 1:2000), Oregon Green 488 phalloidin (green, 1:50) were used. Confocal microscopy was applied for observation. Scale bars=10 µm. (A) Immunostaining of Drebrin and actin. (B) Quantification of % of protrusions in each sample. The data were presented as mean ± S.D., and one way ANOVA analysis with Tukey’s test was used to analyze the differences between the individual experiments.
(B)

Percentage of cells with protrusions

- WT Drebrin
- K185R/K186R
- K270R/K271R

Percentage of cells with protrusions (%)
Table 2.1. Seventy-four proteins identified in cells expressing four different SUMO-1 constructs. Proteins were identified by nano-LC-MS/MS from in-gel tryptic digestion followed by MASCOT search. The number of SUMOylation consensus sites and predicted non-consensus SUMOylation sites were analyzed by SUMOsp 2.0 software. A number of proteins with higher number of predicted SUMOylation sites (large than four, at least one SUMO consensus site) were highlighted.

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Table 2.2. Thirteen proteins with “GG”-tag identified by mass spectrometry from cells overexpressing SUMO-1 (“GG-T95R”). Proteins were identified by nano-LC-MS/MS from in-gel tryptic digestion followed by MASCOT search. T-complex protein 1 subunit delta and Ran GTPase-activating protein 1 are also in the common protein list (Table 2.1).

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</tr>
<tr>
<td>Lethal(2) giant larvae protein homolog 1 (LLGL)</td>
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<td>T-complex protein 1 subunit delta (TCP-1-delta)</td>
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Chapter 3. The role of PDB-associated p62 mutants in NF-κB signaling pathway

Introduction

PDB-associated p62 mutations are in 40% of the familiar PDB cases, and the upregulation (or defective) NF-κB signaling pathway is linked to PDB [7, 9]. Therefore, it is important to know the functional consequences of p62 PDB mutations [7, 9]. However, not much of the cellular impact of PDB-associated p62 mutations is known. In this Chapter, the focus is on the role of p62 PDB mutants in the NF-κB signaling pathway, which is related to the formation of hyperactivated osteoclasts as described in Chapter 1 [59, 67]. Specifically, we ask two important questions: (1) Do PDB-associated p62 mutations increase the NF-κB signaling? and (2) If so, by which mechanism do p62 PDB mutants increase the signaling?

Previous studies have reported that four mutations in p62 including P364S [79], K378X [81], P392L [79, 80] and E396X [81] increased NF-κB signaling. In this Chapter, whether several other mutants (M404V, G411S, D335E and A381V) also increase the NF-κB signaling was tested. Also, there is no report regarding how p62 PDB mutants increase the NF-κB signaling [9, 59]. Therefore, the possible mechanisms by which PDB-associated p62 mutants increase the NF-κB signaling pathway were tested.

We initiated our study on the impact of p62 PDB mutants using Raw264.7 cells, which are osteoclast-like cells and widely used in the field [170]. To study the impact of PDB-associated p62 mutants on signaling, NF-κB luciferase assays and IκB degradation assays were performed in Raw264.7 cells overexpressing WT p62 and mutant p62
induced by the GST-rRANKL [171]. Due to the high background of endogenous p62 and low transfection efficiency of Raw264.7 cells, it was challenging to draw definite conclusions. TNFα and RANKL belong to the same family [172]. Therefore, we subsequently studied the impact of p62 PDB mutants in TNFα-induced NF-κB signaling using p62 KO MEF cells, which have the advantage of lacking of endogenous p62.

Furthermore, previous studies have reported that p62 facilitates TRAF6 polyubiquitination, and activates the NF-κB signaling pathway [94]. Therefore, we firstly hypothesized that PDB-associated p62 mutants increase NF-κB signaling by increasing TRAF6 polyubiquitination. In this Chapter, we aim to test this hypothesis.

**Materials and methods**

**Cells and reagents**

Raw264.7 cells were a gift from Dr. Lisa Cassis (University of Kentucky). WT MEF and p62 KO MEF cells were kindly provided by Dr. Masaaki Komatsu at Tokyo Metropolitan Institute of Medical Science [106]. The Raw264.7 cell line stably expressing the NF-κB luciferase reporter was generously shared by Dr. Jiake Xu (University of Western Australia, Australia) [170, 173]. HEK 293 cells stably expressing RANK receptor was kindly provided by Dr. Sarah Rea (Sir Charles Gairdner Hospital, Australia) and Dr. Julie Crockett (University of Aberdeen, United Kingdom) [79].

Cells were cultured at 37°C with 5% CO₂. HEK293 cells were cultured in Dulbecco’s Modified Eagle’s Medium (DMEM) supplemented with 10% fetal bovine serum (FBS), 100 units/ml penicillin and 100 µg/ml streptomycin. WT MEF and p62 KO
MEF cells were cultured in DMEM supplemented with FBS, PS, 100 mM Sodium Pyruvate (11360-070; Invitrogen) and 10 mM Non-Essential Amino Acids (11140-050; Invitrogen). Raw264.7 cells were cultured in DMEM supplemented with FBS, PS, Sodium Pyruvate (Invitrogen), Non-Essential Amino Acids (Invitrogen) and 20 mM HEPES (15630-106; Invitrogen). For Raw264.7 cells stably expressing NF-κB luciferase reporter, an additional 400 µg/ml G418 (G8168; Sigma) was required. HEK293 cells stably expressing the RANK receptor were cultured in DMEM supplemented with FBS, PS and 100 µg/ml Hygromycin B (400053; EMD Biosciences).

Additionally, GST-rRANKL, NF-κB luciferase reporter and HA-RANK plasmids were generous gifts from Dr. Jiake Xu (University of Western Australia, Australia) [171, 173]. GST-rRANKL protein was purified with the help of Dr. Weimin Gong (Institute of Biophysics in Beijing, China). HA-Ub constructs including K29R, K48R and K63R were kindly shared by Dr. Marie Wooten (Auburn University). The 3xHA-Ub constructs including WT, “K48 only” and “K63 only” were gifts from Dr. Matthew Gentry (University of Kentucky). The phRL-TK (Renilla) vector used in this study was from Promega. The information of plasmids is summarized in Appendix I.

Plasmid construction

The Myc-p62 (human) plasmid was a gift from Dr. Marie Wooten (Auburn University). The human p62 was amplified by PCR and was inserted among the EcoRI and KpnI sites of the p3xFLAG-CMV10 vector (Sigma). Five PDB mutants, including D335E, A381V, P392L, M404V and G411S were generated by using the QuikChange II
Site-Directed Mutagenesis Kit (Stratagene). The details of these plasmids (Appendix I) and designed primers (Appendix II) for mutagenesis are shown in the Appendix.

**NF-κB luciferase assay**

For the study of RANKL-induced NF-κB signaling, 70% confluent Raw264.7 cells stably expressing NF-κB luciferase reporter were transfected with Renilla control vector and WT p62 or p62 PDB mutants. Forty-eight hours after the transfection, cells were treated with GST-rRANKL (30ng/ml) for 7 hours. Cells were lysed with the Passive Lysis Buffer (PLB) of the Dual Luciferase Reporter Assay System (Promega). Aliquots of the cell lysates were subjected to SDS-PAGE and Western blotting using antibodies including anti-FLAG (A8592; Sigma) and anti-actin (sc-1616; Santa Cruz Biotechnology). The NF-κB luciferase assays were performed using the Dual Luciferase Reporter Assay System (Promega) using Optocomp I luminometer.

For the study of TNFα-induced NF-κB signaling, 70% confluent p62 KO MEF cells or WT MEF cells were transfected with Renilla vector, NF-κB luciferase reporter and WT p62 or p62 PDB mutants. Twenty-four hours after the transfection, cells were starved with DMEM containing only 0.1% BSA for 4 hours. Cells were then treated with 20 ng/ml mouse TNFα (315-01A; PeproTech, Inc.) overnight. The next day, cells were lysed in PLB. Aliquots of the cell lysates were subjected to SDS-PAGE and Western blotting using antibodies including anti-FLAG (A8592; Sigma) and anti-actin (sc-1616; Santa Cruz Biotechnology). NF-κB luciferase assays were performed using the Dual Luciferase Reporter Assay System (Promega).
**Statistical analysis**

Three independent NF-κB luciferase assays in either p62 KO MEF cells or HEK293 cells were performed. All data were presented as mean with standard deviation (SD). The significant differences in NF-κB activity fold increases between control cells and cells expressing WT p62 were analyzed by t-test. The significant differences in NF-κB activity fold increases among cells expressing WT p62 and p62 PDB mutant were analyzed by one-way ANOVA, with Tukey’s post-test using the software GraphPad Prism 5 Demo.

**Monitoring the rate of IκB degradation**

Raw264.7 cells or p62 KO MEF cells were transfected with WT p62 or M404V p62. After 24 hours, the cells were either left untreated or were treated with GST-rRANKL (100 ng/ml) or mouse TNFα (30 ng/ml) for 15 or 45 minutes. Then, the cells were lysed with RIPA buffer (Millipore) supplemented with PMSF, P8340 protease inhibitor cocktail (Sigma), Na₃VO₄ and NEM for 30 minutes. The cell lysates were centrifuged at 1000g for 10 minutes and the supernatants were boiled with 6xSDS loading buffer. Samples were subjected to SDS-PAGE, followed by Western blotting using antibodies including anti-IκBα (mAb #4812; Cell Signaling) and anti-actin (sc-1616; Santa Cruz Biotechnology).
**P65 nuclear translocation**

Raw264.7 cells were left untreated or treated with GST-rRANKL (100 ng/ml) for 30 minutes. The cells were fixed with 4% paraformaldehyde (PFA) at 37°C for 15 minutes, permeabilized with 0.1% Triton in PBS and blocked with 3% bovine serum albumin (BSA) in PBS for 1 hour. All the primary and secondary antibodies were diluted in 3% BSA/PBS. Cells were first stained with p65 antibody (mouse, sc-8008, Santa Cruz Biotechnology, 1:50) overnight. The next day, the coverslips were washed with PBS and incubated with Hoechst (33258; Sigma, 1:1000) and secondary Alexa Fluor 488 anti-mouse antibody (A21202; Invitrogen, 1:300) for 2 hours. Finally, the coverslips were mounted using Vectashield mounting medium (Vector Laboratories). Fluorescence microscopy was performed using a confocal microscope (Olympus FluoView) with a 60x objective.

**P62 siRNA**

Raw264.7 cells or Raw264.7 cells stably expressing NF-κB luciferase reporter were transfected with non-targeting siRNA (Dharmacon, D-001210-01-05) or p62 siRNA (40nM or 80nM, Dharmacon, M-047628-01) using Lipofectamine LTX transfection reagent (Invitrogen) in the Opti-MEM reduced serum medium (Invitrogen). Forty-eight hours after transfection, cells were lysed with RIPA buffer (Millipore). The cell lysates were centrifuged at 1000g for 10 minutes and the supernatants were boiled with 6xSDS loading buffer. Samples were subjected to SDS-PAGE. The knockdown efficiency was checked by Western blotting using antibodies including anti-p62 (H00008878-M01; Abnova) and anti-actin (sc-1616; Santa Cruz Biotechnology).
**TRAF6 ubiquitination assay**

HEK293 cells were transfected with FLAG-TRAF6 and DsRed-p62 (WT or mutant) using the Lipofectamine transfection reagent (Invitrogen). Forty-eight hours after transfection, cells were lysed using either PPHB buffer (50 mM Na₂HPO₄, 1 mM sodium pyrophosphate, 20 mM NaF, 2 mM EDTA, 2 mM EGTA and 1% Triton X-100) or RIPA buffer (Millipore). Cells were starved for 3.5 hours and treated with human TNFα (30ng/ml) for 10 minutes. The cell lysates were centrifuged at 1000g for 10 minutes and pre-cleared with Sepharose 4L-CB (Sigma) beads for 1 hour. The lysates were then incubated with anti-FLAG M2 affinity beads for 2 hours at 4°C (F2426; Sigma). The beads were washed three times with lysis buffer and the immunoprecipitation (IP) products were eluted by boiling in SDS-PAGE loading buffer. The IP products and the extracts were subjected to SDS-PAGE using 4–20% ReadyGel Tris–HCl gradient gels (BioRad), followed by Western blotting with different antibodies including anti-ubiquitin (sc-8017, Santa Cruz Biotechnology), anti-FLAG (A8592; Sigma) and anti-actin (sc-1616, Santa Cruz Biotechnology).

**Western blotting and quantification**

Nitrocellulose membrane was incubated in blocking solution, 5% milk in Tris-Buffered Saline and Tween 20 (TBST) for one hour. Then the membranes were incubated with primary antibody for more than three hours. After four washes with TBST for five minutes each, the membrane was incubated with the secondary antibody for more than
one hour. After four washes again with TBST for five minutes each, proteins of interest were visualized by either normal or dura enhanced chemiluminescent (ECL) substrate (Thermo scientific) for detection of horseradish peroxidase (HRP) enzyme (Thermo scientific). The membrane was covered with the wrapping membrane and an autoradiography film (Denville Scientific) which was exposed to the membrane. The exposure time varied from one second to 20 minutes depending on the signal intensity. Films were subsequently developed by a Kodak X- OMAT 2000 processor.

Software Image J was used for quantification of Western blotting bands on X-ray films. Since ECL signals of the Western blot were captured on X-ray films which are known to have a narrow linear range of detection, the quantification of the Western blot may be out of linear range for certain experiments. Enhanced chemofluorescence (ECF) substrate and Alkaline Phosphatase (AP)-conjugated secondary antibody are encouraged for use in the future.

Results

Effect of the p62 PDB mutation on RANKL-induced NF-κB signaling in Raw264.7 cells

We used three classical techniques, including the NF-κB luciferase assay, IκB degradation assay and p65 nuclear translocation assay, to study the impact of PDB-associated p62 mutants in the RANKL-induced NF-κB signaling.

(1) NF-κB luciferase assay
The schematic diagram of the NF-κB signaling in these studies is shown in Fig. 3.1. After RANKL treatment, a significant increase of the luciferase signal was observed, suggesting that the RANKL reagent and the whole system were working (Fig. 3.2 C). It was also shown that the endogenous level of p62 was high (Fig. 3.2 A). The level of overexpressed p62 was different among WT p62 and mutant p62 (M404V, A381V and P392L) (Fig. 3.2 A). Therefore, it is reasonable to normalize the NF-κB luciferase data using the expression level of p62 (Fig 3.2 B). The original data of Firefly and Renilla were shown in Fig. 3.2 C. Fold increases after treatment were calculated by using the Firefly data divided by the Renilla. These increased ratios were further calibrated by using the data from the overexpressed p62 without treatment (Fig 3.2 B). Final calibrated fold increases were obtained (Fig. 3.2 D). It is shown that mutant M404V and A381V increased the NF-κB luciferase activity compared with WT p62 (Fig. 3.2 D). The high endogenous p62 level in Raw264.7 cells prevented us from further studying the impact of mutant p62 on signaling (Fig. 3.2 A). Also, the luciferase assay in HEK293 cells expressing the RANK receptor induced by RANKL was used, but did not produce a signal (data not shown).

(2) **IκB degradation assay**

The IκB degradation assay in Raw264.7 cells induced by RANKL (Fig. 3.3) was then performed. When stimulated with RANKL, IκB was phosphorylated, ubiquitinated and degraded by the proteasome (Fig. 1.5). Therefore, the lower IκB level indicates increased NF-κB signaling activity. The data showed that the IκB level in cells expressing M404V was always lower than that in cells expressing WT p62 when treated with RANKL for either 15 or 45 minutes (Fig. 3.3 B). This result indicates that M404V
p62 increased the NF-κB signaling. Again, due to the high endogenous p62 level in Raw264.7 cells, it is difficult to reach a definite conclusion.

(3) P65 nuclear translocation

Raw264.7 cells were either treated or untreated with RANKL for 30 minutes. P65 nuclear translocation was observed using confocal microscopy. After stimulation of RANKL in Raw264.7 cells, a significant amount of p65 was observed in the nucleus (Fig. 3.4 upper panel), whereas p65 remained in the cytoplasm without treatment (Fig. 3.4 lower panel). These results indicate that RANKL induced p65 nuclear translocation. Since the high background of p62 is an issue, comparison of the WT p62 and mutant p62 has not been done. Also, a better cell model system is needed to study the impact of p62 PDB mutants in RANKL-induced NF-κB signaling (see “Discussion”).

Attempts to knockdown the endogenous level of p62 of Raw264.7 cells by using the siRNA were also tried. However, the p62 siRNA (40nM and 80nM) did not decrease the endogenous p62 level (Fig. 3.5). It is possible that Raw264.7 cells have low transfection efficiency for siRNA too.

P62 contributes to TNFα-induced NF-κB signaling

Firstly, this study has shown that the Firefly/Renilla ratio after TNFα treatment was higher in WT MEF cells compared with p62 KO MEF cells (Fig. 3.6 A and B). It is shown that overexpression of WT p62 in p62 KO MEF cells increased the signaling compared with control p62 KO MEF cells (Fig. 3.6 C and D). Altogether, this suggests that p62 contributes to TNFα-induced NF-κB signaling.
This conclusion is further supported by three independent TNFα-induced NF-κB luciferase assays in p62 KO MEF cells (Fig. 3.9 A). In order to find the best concentration of TNFα for measurement, cells were treated with different concentrations of TNFα (9 ng/ml, 18 ng/ml and 30 ng/ml) overnight. It is shown that 18 ng/ml performed best which could cause the largest difference of Firefly/Renilla signals between cells overexpressing WT p62 and control cells (Fig. 3.7). Therefore, 20ng/ml TNFα for the following luciferase assay for statistical study was used. It is shown that cells expressing WT p62 had higher fold increases after TNFα treatment compared with control cells to a statistically significant extent, with the p value 0.034 (Fig 3.9 A).

**P62 PDB mutants have a tendency to increase NF-κB signaling compared with WT p62**

Next, the impact of PDB-associated p62 mutants in TNFα-induced NF-κB signaling was examined. Firstly, it is shown that Firefly/Renilla signals were higher in p62 KO MEF cells overexpressing M404V p62 compared with WT p62, suggesting that M404V p62 increased TNFα-induced NF-κB signaling compared with WT p62 (Fig. 3.6 C and D). Then, we performed three independent TNFα-induced NF-κB luciferase assays in p62 KO MEF cells (Fig. 3.8 and 3.9). The expression level of FLAG-p62 was similar among cells overexpressing WT and mutant p62 (Fig. 3.8 A). The expression level of p62 was also used for calibration of fold increases in the analysis (Fig. 3.9 C).

Firefly/Renilla signals with or without TNFα in cells overexpressing WT and mutant p62 are shown in Fig. 3.8 B. The fold increases were calculated by using Firefly/Renilla (with TNFα) divided by Firefly/Renilla (without TNFα). The luciferase data either without considering the p62 expression level (“Non-calibrated fold increases”, 86
Fig. 3.9 B) or considering the p62 level (“Calibrated fold increases”, Fig. 3.9 C) was analyzed. After statistical analysis, it is shown that p62 PDB mutants including M404V, A381V and P392L all increased signaling compared with WT p62 (Fig. 3.9 B and C). However, the increase is not statistically significant with the p value larger than 0.05.

The TNFα-induced NF-κB luciferase assay was also performed in HEK293 cells. The results have shown that p62 PDB mutants had a tendency to increase the signaling compared with WT p62 (Fig. 3.10 B).

**P62 PDB mutant suppresses its binding with polyubiquitinated proteins in HEK293 cells**

Next, we considered mechanisms by which p62 PDB mutants increased signaling. We started to study the cellular consequences of p62 PDB mutants. Because most of PDB mutations are in the UBA domain of p62, which binds the polyubiquitinated proteins [9], we want to know that whether these p62 PDB mutants have effects on the binding of polyubiquitinated proteins.

To address this question, HEK293 cells were transfected with human FLAG-p62 (WT, M404V, A381V, P392L and G411S) or mouse p62 (WT and UBA domain deleted). Forty-eight hours after transfection, FLAG-IP was performed, and it was shown that all of the mutants, especially M404V and P392L p62, impaired the binding of p62 and polyubiquitinated proteins compared with WT p62 (Fig. 3.11). Mouse p62 (UBA domain deleted) also impaired its binding with polyubiquitinated proteins compared with mouse
WT p62, suggesting the validation of negative control (Fig. 3.11). However, for the FLAG-p62 IP shown in Fig. 3.11, whether it was polyubiquitination of p62 or polyubiquitination of other proteins remains investigation.

**Overexpression of p62 in HEK293 cells leads to TRAF6 polyubiquitination in HEK293 cells**

As mentioned above, it was found that the p62 PDB mutations abolish binding with polyubiquitinated proteins in HEK293 cells. It has also been reported that TRAF6 polyubiquitination is regulated by p62 [94]. Therefore, it is possible that p62 PDB mutations could have effects on TRAF6 polyubiquitination. To address this question, the TRAF6 ubiquitination assay in HEK293 cells overexpressing FLAG-TRAF6 and DsRed-p62 (WT, M404V, A381V and P392L) was performed.

It is shown that in the basal condition, the level of TRAF6 polyubiquitination in cells overexpressing WT p62 is much higher than control cells, indicating that overexpression of p62 in HEK293 cells leads to TRAF6 polyubiquitination (Fig. 3.12, 3.13 and 3.14). In addition, it was found that the level of TRAF6 polyubiquitination increases in proportion to the amount of p62 (Fig. 3.13). It is indicated that the level of TRAF6 polyubiquitination is dependent on p62 amount.

**Mutant p62 impaired TRAF6 polyubiquitination compared with WT p62 in HEK293 cells**

From the TRAF6 ubiquitination assay in HEK293 cells above, it was shown that the level of TRAF6 polyubiquitination in cells overexpressing PDB-associated p62
mutants (M404V, A381V and P392L), especially M404V p62 is lower than cells overexpressing WT p62 (Fig. 3.12, 3.13 and 3.14). Therefore, p62 PDB mutants, especially M404V, impaired TRAF6 polyubiquitination compared with WT p62 in the basal condition. However, compared with control cells without overexpression of p62, cells overexpressing p62 PDB mutant still increase TRAF6 polyubiquitination (Fig. 3.12, 3.13 and 3.14), indicating that both the WT and the p62 PDB mutant could facilitate TRAF6 polyubiquitination in HEK293 cells.

**TNFα did not induce TRAF6 polyubiquitination in HEK293 cells**

As mentioned above, it was shown that p62 contributes to TNFα-induced NF-κB signaling (Fig. 3.9 A). It was also shown that PDB-associated p62 mutants increased TNFα-induced NF-κB signaling (Fig. 3.9 and 3.10). Furthermore, it is shown that both WT and mutant p62 facilitate TRAF6 polyubiquitination in HEK293 cells in the basal condition (Fig. 3.12, 3.13 and 3.14). Therefore, it was of interest to determine whether WT p62 and p62 PDB mutants change TRAF6 polyubiquitination upon TNFα treatment compared with basal conditions. In order to address this question, TRAF6 ubiquitination assay was performed in the absence and presence of TNFα in HEK293 cells. It was shown that TNFα was effective because IκB was degraded after TNFα treatment (Fig 3.14). However, the change of TRAF6 polyubiquitination after TNFα treatment (Fig. 3.14) was not observed. Therefore, it is suggested TNFα does not play a role on TRAF6 polyubiquitination in HEK293 cells.
**Overexpression of p62 in HEK293 cells leads to a TRAF6 polyubiquitination chain of different linkages**

Previous work has shown that p62 facilitates K63-linked polyubiquitination of TRAF6 and further activates the NF-κB signaling [95] (Fig. 1.5). In this study, it was found that p62 facilitates TRAF6 polyubiquitination in HEK293 cells. Therefore, is the TRAF6 polyubiquitination observed (Fig. 3.12 and Fig. 3.14) a K63-linked chain? Could the TRAF6 polyubiquitination observed activate the NF-κB signaling upon TNFα treatment?

To address these questions, HEK293 cells were transfected with FLAG-TRAF6, DsRed-p62 (WT) and a variety of different HA-Ub constructs including K29R, K48R, K63R [95], as well as “K48 only” or “K63 only” in which all the Lys of Ub are mutated to Arg except Lys48 or Lys63, respectively. If it is a K63-linked chain, the level of TRAF6 polyubiquitination should decrease in cells overexpressing HA-Ub (K63R) compared with cells overexpressing WT Ub. However, this difference (Fig. 3.15, lane 1 and 4) was not seen. There is no difference among levels of TRAF6 polyubiquitination in cells overexpressing K29R Ub, K48R Ub and K63R Ub (Fig. 3.15, lane 2-4). In addition, there is no difference between levels of TRAF6 polyubiquitination in cells overexpressing Ub (“K63 only”) and Ub (“K48 only”) (Fig. 3.15, lane 5 and 6). Altogether, it is suggested that the TRAF6 polyubiquitination observed is not a K63-linked chain, but a mixture of K48-linked, K63-linked and K29-linked chains.

Moreover, from the TRAF6 polyubiquitination assay in HEK293 cells with and without TNFα treatment (Fig. 3.14), it is shown that overexpression of p62 could not
activate the NF-κB signaling because IκB level is similar in the absence and presence of p62 (Fig. 3.14). Therefore, the TRAF6 polyubiquitination observed by overexpressing p62 in HEK293 cells could not activate the NF-κB signaling (Fig. 3.14).

**Discussion**

Does p62 really facilitate TRAF6 to form a K63-linked polyubiquitin chain which activates NF-κB signaling?

Wooten et al. [95] and Moscat et al. [93] have shown that p62 could activate the NF-κB signaling in HEK293 cells in the basal condition. Our study not only showed that p62 activates NF-κB signaling basally (Fig. 3.6 C), but also upon TNFα treatment to a statistically significant degree in p62 KO MEF cells (Fig. 3.9 A) and HEK293 cells (Fig. 3.10 A).

Wooten et al. [95] have reported that the loss of p62 completely abolishes ubiquitination of TRAF6 by performing TRAF6 immunoprecipitation from lysates of the brains from p62 WT or knock-out mice. In this study, it was shown that p62 facilitates TRAF6 polyubiquitination in HEK293 cells (Fig. 3.12 and 3.14). In their study, they have shown that the TRAF6 polyubiquitination chain is a K63-linked chain by using a variety of different Ub constructs. We also used these different Ub mutants and performed the TRAF6 ubiquitination assay as done previously [95]. However, it was found that the TRAF6 polyubiquitination chain is not K63-linked, but a mixture of K29-linked, K48-linked and K63 linked. It was further confirmed by using two other different Ub mutants, “K48 only” and “K63 only”. The contradictory results between this study and Dr.
Wooten’s study might be due to the specific experiment details. In their study, although they have shown that TRAF6 polyubiquitination is a K63-linked chain, they did not show that this chain is sufficient to induce the NF-κB signaling. In this work, it was shown that the TRAF6 polyubiquitination observed is not sufficient to induce NF-κB signaling by comparing the IκB level in cells with and without overexpressing p62 (Fig. 3.14).

These data have shown that p62 contributes to NF-κB activation (Fig. 3.9 A). However, for the TRAF6 polyubiquitination assay, the IκB level remains similar in HEK293 cells with and without overexpressing p62, which seems to contradict that p62 contributes to NF-κB signaling. To clarify this point in the future, more signaling proteins such as IKK and phospho-IκB need to be examined.

**Do p62 PDB mutants impair the TRAF6 polyubiquitination upon cytokine treatment?**

In this Chapter, it was shown that M404V p62 impairs TRAF6 polyubiquitination compared with WT p62 in the basal condition (Fig. 3.12 and 3.14). Also, it has been shown that M404V p62 had a tendency to increase TNFα-induced NF-κB signaling (Fig. 3.10 B). Because TRAF6 polyubiquitination facilitates NF-κB signaling [125], these two pieces of data seem to be contradictory. However, it is suggested in the literature that, it is “K63 linked” polyubiquitination that facilitates NF-κB signaling. Was the “K63 linked” polyubiquitination chain in our FLAG-TRAF6 IP observed? Actually it did not. These data have shown that the TRAF6 polyubiquitination chain is a mixture of polyubiquitination chains which contain K29 linked, K48 linked and K63 linked chains.
(Fig. 3.15). Therefore, overexpression of p62 could not facilitate TRAF6 to form the K63-linked chain.

In addition, overexpression of p62 does not appear to activate NF-κB signaling because the IκB levels were similar in the absence and presence of p62 (Fig. 3.14). It is suggested that TRAF6 K63-linked polyubiquitination might require the stimulation of cytokines, such as TNFα or RANKL. It has also been shown that TNFα did not induce the TRAF6 polyubiquitination (Fig. 3.14). Therefore, it is indicated that TRAF6 K63-linked polyubiquitination could be observed by stimulation with other cytokines, such as RANKL.

Moreover, these data showed that TNFα did not increase TRAF6 polyubiquitination (Fig. 3.14). Previous studies reported that TNFα increased TRAF2 polyubiquitination [23, 93, 135], but they did not test whether TNFα increased TRAF6 polyubiquitination. Thus, our study expands the knowledge of TNFα. In addition, Fuanakoshi-Tago et al. [174] have reported that TRAF6 negatively regulates TNFα-induced NF-κB signaling. They found that IKK activation and IκB degradation were enhanced in TRAF6-deficient MEFs compared with WT MEFs [174]. Although our data and their data are not directly related, all these studies showed at least that TNFα-induced NF-κB signaling is not activated through TRAF6.

**Three workable cell models for studying the impact of PDB-associated p62 mutants on NF-κB signaling**

*(I) Raw264.7 cells*
Previously, Raw264.7 cells stably expressing the NF-κB luciferase reporter were used as a cell model to study the effect of the p62 on the RANKL-induced NF-κB signaling. It was thought that Raw cells constituted a good cell model because they are osteoclast-like cells [170], which are widely used in the field studying PDB. However, for this project, the goal was to examine the effect of PDB-associated p62 mutations on NF-κB signaling. Because all these p62 mutations are single point mutations, it was recognized that it is very important to rule out the effect of the endogenous p62 of Raw cells. It turned out that Raw cells have a high background of endogenous p62 (Fig. 3.2 A), whereas the transfection efficiency for Raw cells are also low. The use of p62 siRNA to knockdown the endogenous p62 of Raw cells was attempted, but p62 siRNA did not work; this might also be due to the low transfection efficiency of Raw cells (Fig. 3.5). Therefore, other cell models to study the signaling were considered.

(2) p62 KO MEF cells

P62 KO MEF cells shared by Dr. Masaaki Komatsu (Japan) [106] were used for this study. The advantage of p62 KO MEF cells is that there is no endogenous p62 in this cell line. Therefore, it is advantageous to compare the effect of WT p62 and p62 PDB mutant using this p62 KO MEF cell line. The disadvantage of this cell line is that the transfection efficiency for the cell is also low. In addition, p62 KO MEF cells did not respond to the RANKL (data not shown). HA-RANK receptor (shared by Dr. Jiake Xu, Australia) was transfected into p62 KO MEF cells, and these cells again did not respond to RANKL (data not shown). However, p62 KO MEF cells did respond to TNFα (Fig. 3.6). Therefore, the study shifted to the effect of PDB-associated p62 mutations on TNFα-induced NF-κB signaling. The TNFα-induced NF-κB luciferase assay was
performed in p62 KO MEF cells overexpressing either WT p62 or p62 PDB mutant for more than three times (Fig. 3.8 and 3.9). These data have shown a tendency for mutant p62 to increase TNFα-induced NF-κB signaling compared with WT p62, which is not statistically significant in p62 KO MEF cells. Additionally, it is shown that even without p62, there is still basal TNFα-induced NF-κB signaling (Fig. 3.9). Also, p62 increases the TNFα-induced NF-κB signaling pathway to a statistically significant degree (p<0.05) (Fig. 3.9 A). Therefore, p62 PDB mutants increase NF-κB signaling in p62 KO MEF cells, but the increase is so subtle that it is difficult to detect with NF-κB luciferase assay.

(3) HEK293 cells

In addition, the TNFα-induced NF-κB assay in HEK293 cells was studied. HEK293 cells could also respond to TNFα signals (Fig. 3.10). In addition, the increased folds in cells expressing p62 (the number is over 50) are much higher than control cells, suggesting that it might be a good model to compare the effect of WT p62 and mutant p62 on TNFα-induced NF-κB signaling.

It appears to be challenging to study the RANKL-induced NF-κB signaling in this study. An ideal cell model should have little background of endogenous p62, high transfection efficiency and response after stimuli of RANKL. While it is difficult to find such a perfect cell model for study, some ways to optimize our existent cell lines are considered. The strategies are described below and summarized in Table 3.1.

(1) HEK293 cells stably expressing RANK receptor

Rea et al. [79] reported that P364S and P392L increased RANKL-induced NF-κB signaling by NF-κB luciferase assay in HEK293 cells stably expressing RANK receptor.
We also obtained the above stable HEK293 cells from them. Although we tried the same assay as described in their paper, the NF-κB signaling induced by GST-rRANKL [171] was not detected (data not shown).

(2) p62 KO MEF cells stably expressing HA-RANK receptor

HA-RANK receptor was also kindly shared by Dr. Jiake Xu (Australia). We also tried transient transfection of HA-RANK into p62 KO MEF cells, but again cells did not respond to GST-rRANKL [171] (data not shown). Because currently no report regarding using p62 KO MEF cells expressing HA-RANK receptor, there is a need to change the parameters for experiment optimization. If it works in the future, this approach might generate the p62 KO MEF cells stably expressing HA-RANK receptor.

(3) Raw264.7 cells with p62 shRNA

Since Raw264.7 cells have low transfection efficiency, lentivirus-delivered p62 shRNA and exogenous p62 (WT or mutant) could be applied. It will be necessary to be careful about the design of p62 shRNA, which should only interfere with endogenous p62, not exogenous p62. Biosafety issues also need to be considered.

Significance of this study and future directions

The significance of this study in this Chapter is that this is the first example using the p62 KO MEF cells as a model for studying the effect of p62 PDB mutants on NF-κB signaling. Also, this is the first study to show that p62 not only increases TNFα-induced NF-κB signaling basally, but also upon TNFα treatments in p62 KO MEF cells, which was not reported. Moreover, these data have shown that several PDB-associated p62
mutants had tendency to increase TNFα-induced NF-κB signaling, which was not reported before. Additionally, it excluded the possibility that these p62 PDB mutants increase signaling through increasing TRAF6 polyubiquitination, suggesting the existence of other mechanisms.

Future directions include examining other possible mechanisms by which p62 PDB mutants increase TNFα-induced NF-κB signaling. Another direction is to find a better model to study the impact of PDB-associated p62 mutants on RANKL-induced NF-κB signaling (see Chapter 5). Additionally, previous studies have shown that p62 interacts with RIP and is involved in the TRAF2 polyubiquitination [93] in the TNFα-induced NF-κB signaling, which raises the possibility that PDB-associated p62 mutations increase TNFα-induced NF-κB signaling through increasing TRAF2 polyubiquitination. This hypothesis could be investigated in the future.
Figure 3.1. A schematic diagram of NF-κB luciferase assay.
Figure 3.2. RANKL-induced NF-κB luciferase assay in Raw264.7 cells expressing WT p62 and p62 PDB mutants. Raw264.7 cells stably expressing NF-κB luciferase reporter were transfected with FLAG-p62 (WT, M404V, A381V and P392L) and Renilla. Forty-eight hours after transfection, cells were treated with GST-rRANKL (100ng/ml) for 7 hours. Cells were harvested with passive lysis buffer and NF-κB luciferase assay was performed. Cell lysates were also subjected to SDS-PAGE and Western blotting was performed using antibodies against FLAG and actin. (A) Western blotting of overexpressed p62 and endogenous p62. (B) Quantification of FLAG-p62/Actin by Image J. (C) Original Firefly/Renilla ratio with or without RANKL treatment. (D) Normalized fold increases after RANKL treatment according to the expression level of p62.
Figure 3.3. Comparison of the rate of IκB degradation in Raw264.7 cells expressing WT and mutant p62 induced by GST-rRANKL. (A) IκB degradation in Raw264.7 cells overexpressing WT or M404V p62 induced by RANKL at 10 or 45 minutes. (B) Quantification of IκB/Actin by Image J software.
Figure 3.4. Immunostaining of p65 in Raw264.7 cells induced by GST-rRANKL. Raw 264.7 cells were treated with and without GST-rRANKL (100ng/ml) for 30 minutes. Cells were fixed with 4% PFA and permeabilized with 0.1% Triton. Primary antibody p65 (1:50), secondary antibody Alexa Fluro 488 (mouse) (green, 1:300) and Hoechst (blue, 1:1000) were used. Confocal microscopy was applied for observation. Scale bars=5µm.
Figure 3.5. Knockdown of the endogenous p62 in Raw264.7 cells by p62 siRNA. Raw264.7 cells stably expressing NF-κB luciferase reporter were transfected with non-targeting siRNA or p62 siRNA (40nM, 80nM) using the Lipofectamine LTX. Forty-eight hours after transfection, cells were lysed with RIPA buffer. Samples were subjected to SDS-PAGE. The knockdown efficiency was checked by Western blotting using antibodies against p62 and actin.
Figure 3.6. TNFα-induced NF-κB luciferase assays in WT MEF and p62 KO MEF cells. (A) Firefly/Renilla ratio induced by TNFα in MEF cells. MEF cells were transfected with NF-κB luciferase reporter and Renilla. Cells were treated with TNFα (30ng/ml) for 7 hours. (B) Fold increases after TNFα treatment in p62 KO MEF and WT MEF cells. (C) Firefly/Renilla ratio at basal condition. MEF cells were transfected with FLAG-p62 (WT or M404V), NF-κB luciferase reporter and Renilla. (D) Firefly/Renilla ratio with or without TNFα treatment. Cells were treated with TNFα (30ng/ml) for 7 hours.
Figure 3.7. NF-κB luciferase assay induced by different concentration of TNFα in p62 KO MEF cells. (A) The Firefly/Renilla ratio was obtained at different concentration of TNFα (9ng/ml, 18ng/ml and 30ng/ml). The cells were starved with DMEM (0.1% BSA) for 4 hours, and then treated with TNFα overnight. (B) Fold increases by different concentration of TNFα.
Figure 3.8. TNFα-induced NF-κB luciferase assay in p62 KO MEF cells expressing WT p62 or p62 PDB mutants. P62 KO MEF cells were transfected with FLAG-p62 (WT, M404V, A381V and P392L), NF-κB luciferase reporter and Renilla. Twenty-eight hours after transfection, cells were starved for 3.5 hours. Cells were treated with TNFα (20ng/ml) overnight. Cells were harvested with passive lysis buffer and NF-κB luciferase assay was performed. Cell lysates were also subjected to SDS-PAGE and Western blotting was performed using antibodies against FLAG and actin. Three independent experiments were performed. (A) Representative Western blotting of overexpressed p62 and quantification of FLAG-p62/Actin by Image J. (B) Representative Firefly/Renilla with and without TNFα.
(B)

Firefly/Renilla with or without TNFα

Increasing folds = \frac{Firefly/Renilla (+TNFα)}{Firefly/Renilla (-TNFα)}

N=3
Figure 3.9. Statistical analysis of fold increases in the TNFα-induced NF-κB luciferase assay in p62 KO MEF cells. TNFα-induced NF-κB luciferase assay were performed three times. (A) Fold increases of Firefly/Renilla in control cells and cells expressing WT p62. The data were presented as mean ± S.D., and *-test was used to analyze the differences between the individual experiments. *: P < 0.05. (B) Fold increases of Firefly/Renilla in cells expressing WT p62 or p62 PDB mutant. The data were presented as mean ± S.D., and one way ANOVA analysis with Tukey’s test was used to analyze the differences between the individual experiments. (C) Fold increases of Firefly/Renilla in cells expressing WT p62 or p62 PDB mutant were calibrated according to the expression level of WT or mutant p62. The data were presented as mean ± S.D., and one way ANOVA analysis with Tukey’s test was used to analyze the differences between the individual experiments.

(A)
Figure 3.10. Statistical analysis of fold increases in the TNFα-induced NF-κB luciferase assay in HEK293 cells. TNFα-induced NF-κB luciferase assay were performed three times. HEK293 cells were transfected with FLAG-p62 (WT, M404V, A381V and P392L), NF-κB luciferase reporter and Renilla. Thirty hours after transfection, cells were starved overnight. Cells were treated with TNFα (30ng/ml) for 4 hours. (A) Fold increases of Firefly/Renilla in control cells and cells expressing WT p62. The data were presented as mean ± S.D., and t-test was used to analyze the differences between the individual experiments. *: P < 0.05. (B) Fold increases of Firefly/Renilla in cells expressing WT p62 or p62 PDB mutant. The data were presented as mean ± S.D., and one way ANOVA analysis with Tukey’s test was used to analyze the differences between the individual experiments. *: P < 0.05.
Fold increases after TNFα

* p<0.05
Figure 3.11. Comparison of immunoprecipitated polyubiquitinated proteins in HEK293 cells overexpressing WT and mutant p62. HEK293 cells were transfected with FLAG-TRAF6 (WT, A381V, P392L and M404V). Forty-eight hours after transfection, FLAG-IP was performed. Samples were subjected to 4–20% gradient gels. Western blotting was performed using antibody against Ub.
Figure 3.12. Comparison of TRAF6 polyubiquitination in HEK293 cells overexpressing WT and mutant p62. HEK293 cells were co-transfected with FLAG-TRAF6 and DsRed-p62 (WT, A381V, P392L or M404V). Forty-eight hours after transfection, FLAG-IP was performed. Samples were subjected to 4–20% gradient gels. Western blotting was performed using antibodies against Ub, FLAG and actin.
Figure 3.13. Comparison of the level of TRAF6 polyubiquitination in HEK293 cells overexpressing different amount of p62. HEK293 cells were co-transfected with FLAG-TRAF6 and different amounts (0.5µg, 0.25 µg and 0.1 µg) of DsRed-p62 (WT or M404V). Forty-eight hours after transfection, FLAG-IP was performed. Samples were subjected to 4–20% gradient gels. Western blotting was performed using antibodies against Ub and FLAG.
Figure 3.14. Comparison of TRAF6 polyubiquitination in HEK293 cells overexpressing WT and mutant p62 with and without TNFα treatment. HEK293 cells were co-transfected with FLAG-TRAF6 and DsRed-p62 (WT, M404V, A381V or P392L). Forty-eight hours after transfection, FLAG-IP was performed. Samples were subjected to 4–20% gradient gels. Western blotting was performed using antibodies against Ub, FLAG, IκB and actin.
Figure 3.15. Comparison of TRAF6 polyubiquitination in HEK293 cells overexpressing p62 and a variety of different ubiquitin constructs. HEK293 cells were co-transfected with FLAG-TRAF6, DsRed-p62 (WT) and different HA-Ub constructs including K29R, K48R, K63R, K48 only and K63 only. Forty-eight hours after transfection, FLAG-IP was performed. Samples were subjected to 4–20% gradient gels. Western blotting was performed using antibodies against Ub, FLAG, DsRed and HA.
Table 3.1. Comparison of three workable cell models for our study.

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<th>Advantage</th>
<th>Disadvantage</th>
<th>Strategy</th>
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<tbody>
<tr>
<td>Raw cell</td>
<td>Osteoclast-like cell Have RANK receptor</td>
<td>Low transfection efficiency High background of p62</td>
<td>p62 siRNA p62 shRNA</td>
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<tr>
<td>p62 KO MEF</td>
<td>No p62</td>
<td>Not Osteoclast-like cell Low transfection efficiency Need to transfec RANK</td>
<td>Transfect RANK receptor</td>
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<tr>
<td>HEK 293</td>
<td>High transfection efficiency</td>
<td>Not Osteoclast-like cell Background of p62 Need to transfec RANK</td>
<td>p62 siRNA Transfect RANK receptor</td>
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Chapter 4. The role of PDB-associated p62 mutants in autophagy

Introduction

It was reported that the p62 levels were elevated in PDB patients, suggesting a defect in autophagy [80, 175], which may play a role in the disease [7, 9]. Little is known, however, about the role of autophagy in PDB. As mentioned earlier, p62 mutations are linked to about 40% familiar PDB cases. Therefore, the effect of PDB-associated p62 mutations in autophagy was investigated in our study.

There are two important questions regarding the role of p62 in PDB. (1) P62 is a regulator of autophagy [6, 90], therefore do p62 PDB mutants increase or decrease autophagic activity compared with WT p62? and (2) P62 is also a substrate of autophagy [82, 150], therefore does the rate of autophagic degradation of p62 PDB mutant increase or decrease compared with WT p62?

In this Chapter, we mainly try to address the first question. We compared the effect of WT p62 and p62 PDB mutants on the interaction of LC3, a marker for autophagosome. Then, we also compared the GFP-LC3 puncta among cells overexpressing WT p62 and mutant p62. In addition, we compared the LC3-II levels after rapamycin (autophagy inducer) and/or NH₄Cl (autophagy inhibitor) treatment in cells expressing WT p62 and mutant p62.
Methods and materials

Cells and reagents

P62 KO MEF cells were a gift from Dr. Masaaki Komatsu at Tokyo Metropolitan Institute of Medical Science [106]. Both rapamycin (autophagy inducer, R0395; Sigma) and NH₄Cl (autophagy inhibitor, 254134-25G; Sigma) were used in this study.

Plasmids construction

The human p62 open reading frame was amplified by PCR and was inserted among the EcoRI and KpnI sites of the p3xFLAG-CMV10 vector (Sigma). Five PDB mutants, including D335E, A381V, P392L, M404V and G411S were generated by site-directed mutagenesis (in Chapter 3). In addition, the FLAG-p62 (WT, A381V, P392L and M404V) was inserted between the EcoRI and KpnI sites of the pDsRed monomer C1, generating four human DsRed-p62 (WT, A381V, P392L and M404V). The mutagenic primer sequences of these DsRed-p62 constructs are listed in Appendix II.

Immunoprecipitation

HEK293 cells were co-transfected with FLAG-p62 (WT, M404V, A381V and P392L) and EGFP-LC3 using the Lipofectamine reagent (Invitrogen). After 48 hours, the cells were lysed with the RIPA buffer with the PMSF (Sigma), P8340 (Sigma), Na₃VO₄ and NEM for 30 minutes. The cell lysates were centrifuged at 1000g for 10 minutes.
Then 1000 μg cell lysates were pre-cleared with the Sepharose 4L-CB beads for 1 hour and subsequently incubated with anti-FLAG M2 affinity beads (F2426; Sigma). The beads were washed with RIPA buffer and resuspended in SDS loading buffer. The IP products and the extracts were subjected to SDS-PAGE followed by Western blotting with antibodies including anti-LC3 (PM036; MBL), anti-FLAG (A8592; Sigma) and anti-actin (sc-1616; Santa Cruz Biotechnology).

**Immunofluorescence and confocal microscopy**

P62 KO MEF cells were transfected with FLAG-p62 (WT or M404V) and EGFP-LC3. Twenty-four hours after transfection, the cells were fixed with 4% paraformaldehyde (PFA) at 37 °C for 15 minutes, permeabilized with PBS/0.1% Triton, and blocked with 3% bovine serum albumin (BSA) in PBS for 1 hour. All the primary and secondary antibodies were diluted in 3% BSA/PBS. Cells were firstly incubated with anti-FLAG primary antibody (mouse, F3165; Sigma, 1:230) overnight. The next day, the cells on the coverslips were incubated with Hoechst (33258; Sigma, 1:1000) and Alexa Fluor 488 anti-mouse (A21202; Invitrogen, 1:300) for 2 hours. Finally, the coverslips were mounted using Vectashield mounting medium (Vector Laboratories). Confocal microscope (Olympus FluoView) was applied with a 60X objective.

**Fractionation**

P62 KO MEF cells were transfected with FLAG-p62 (WT or M404V) using the Lipofectamine LTX (Invitrogen). The cells were treated with NH_4Cl (10mM, 16 hours) or rapamycin (200nM, 16 hours). Forty-eight hours after transfection, the cells were
harvested with RIPA buffer with PMSF (Sigma), P8340 (Sigma), Na₃VO₄ and NEM for 30 minutes. The cell lysates were centrifuged at 1000g for 10 minutes and the supernatant (S1) was collected. S1 was further centrifuged at 20,000g for 50 minutes, generating supernatant (S2) and pellet (P2). The samples were added to the 6xSDS loading buffer and subjected to the SDS-PAGE, followed by Western blotting using antibodies including anti-LC3 (PM036; MBL), anti-FLAG (A8592; Sigma) and anti-actin (sc-1616; Santa Cruz Biotechnology).

**Results**

**Effect of the p62 PDB mutation on interaction with LC3**

LC3-II is an active marker of the autophagosome [8], and p62 binds to LC3 through the LIR domain of p62 [83]. The effect of PDB-associated p62 mutation on this interaction was tested in this study. HEK293 cells were co-transfected with FLAG-p62 (WT, A381V, P392L and M404V) and EGFP-LC3. FLAG-IP was performed followed by SDS-PAGE and Western blotting. We used mouse LIR domain deleted p62 as a negative control, and as expected, it showed no interaction with GFP-LC3 (Fig. 4.1 A, lane 5). The p62 PDB mutation did not, however, change the interaction with LC3 (Fig. 4.1). The interaction of LC3 and p62 bearing a mutation (D335E) in LIR domain was also tested. Interestingly, this mutation also did not alter the interaction with LC3 (data not shown).

**Effect of the p62 PDB mutation on GFP-LC3 puncta formation**
In Chapter 3, we showed that M404V p62 impaired TRAF6 polyubiquitination compared with WT p62 in HEK293 cells (Fig. 3.12 and 3.14). Therefore, we chose this mutant for the study. P62 KO MEF cells were transfected with FLAG-p62 (WT or M404V) and EGFP-LC3. After 24 hours, the cells were fixed with 4% PFA and incubated with anti-FLAG antibody (1:230) and Hoechst (1:1000). Confocal microscopy was used to observe the GFP-LC3 puncta.

It was found that WT FLAG-p62 colocalized with GFP-LC3 (Fig. 4.2), which is consistent with reports in the literature [89]. The same colocalization was found for GFP-LC3 and M404V p62 (Fig. 4.2). In addition, it was found that p62 and LC3 were exclusively located in the cytosol (Fig. 4.2). For quantification, nine Z-stack pictures for each sample were taken (data not shown). Attempts were made to count and compare the GFP-LC3 puncta in cells overexpressing WT and M404V p62 using available software. However, a larger number of cells and better software are needed for more precise quantification.

In the Fig. 4.2, it is shown that GFP-LC3 was in the nucleus. The nuclear staining of GFP-LC3 is likely due to high levels of overexpression.

**Effect of the p62 PDB mutant on LC3-II levels after rapamycin and NH₄Cl treatment**

In Chapter 1, I introduced several available autophagy inducers and inhibitors. Here, we used rapamycin, which inhibits the mTOR pathway, and activates autophagy. Also we used NH₄Cl to inhibit the fusion of lysosomes and autophagosomes, thus
inhibiting autophagy (Fig. 1.7). The level of LC3-II was monitored after treatment with these compounds.

P62 KO MEF cells were transfected with FLAG-p62 (WT or M404V). Cells were then treated with rapamycin for 16 hours. Alternatively, cells were treated with rapamycin and NH₄Cl together for 16 hours. Forty-eight hours after transfection, cells were lysed with RIPA buffer. Cell lysates were centrifuged at 1000g for 10 minutes. The supernatant (S1) was further centrifuged 20,000g for 50 minutes, generating supernatant (S2) and pellet (P2). The S1, S2 and P2 were subjected to SDS-PAGE followed by Western blotting.

Since rapamycin is an autophagy inducer, we expected to see LC3-II levels to increase after rapamycin treatment as more autophagosomes form. For cells without transfection (NT), LC3-II did increase slightly in S1 and S2, and decreased slightly in P2 after rapamycin treatment (Fig. 4.3 A, lane 1 and 5). For cells transfected with empty vector or FLAG-p62 (WT or M404V), LC3-II remained similar in S1, and slightly decreased in S2 and P2 (Fig. 4.3 A, lane 2-4 and lane 6-8). Since LC3-II decreased in P2 after rapamycin treatment in the control cells (Fig. 4.3 A, lane 2 and 6), it was indicated that rapamycin not only increased the formation of autophagosome, but also increased the enzymatic activity in the autophagolysosome, therefore more LC3-II was degraded. Based on Western blotting (Fig. 4.3 A), we quantified the LC3-II/Actin in S1 from cells overexpressing WT p62 or M404V p62 with and without rapamycin (Fig. 4.3 B). There was no significant difference between WT p62 and M404V p62 in terms of the change of LC3-II levels after rapamycin. LC3-II decreased after rapamycin in cells transfected with empty FLAG (Fig. 4.3 B), suggesting that rapamycin increased the enzymatic activity in
the autophagolysosome, thereby degrading LC3-II more rapidly. In cells overexpressing WT p62, the LC3-II levels decreased less compared with the control (7.9% vs. 23.2%) (Fig. 4.3 B), suggesting that WT p62 attenuated the autophagic activity and the rate of LC3-II degradation in the autophagolysosome. However, in cells overexpressing M404V p62, the LC3-II levels decreased more compared with cells overexpressing WT p62 (18.3% vs. 7.9%), suggesting that PDB mutant p62 (M404V p62) may increase the autophagic activity compared with WT p62 because LC3-II was degraded at a faster rate.

NH₄Cl decreases autophagy by inhibiting the fusion of lysosomes and autophagosomes. We expected, therefore, to see LC3-II levels increase after NH₄Cl treatment. As expected, LC3-II increased dramatically in S1, S2 and P2 after NH₄Cl treatment alone (data not shown). Rapamycin was applied with NH₄Cl together with the expectation that this double treatment would raise LC3-II levels even further. Treatment with both compounds (Fig. 4.3 A, lane 9-12), however, did not further increase LC3-II, suggesting LC3-II reached maximal levels after treatment with NH₄Cl alone (data not shown). Based on the Western blotting (Fig. 4.3 A), we quantified the LC3-II/Actin in S1 from cells overexpressing WT p62 or M404V p62 with and without NH₄Cl and rapamycin together (Fig. 4.3 C). It is shown that after double treatment, LC3-II levels in cells overexpressing WT p62 were higher than control (Fig. 4.3 C), suggesting that p62 may facilitate the autophagic activity, which appears contradictory with Fig. 4.3 B. Also, it was shown that after double treatment, LC3-II levels in cells overexpressing M404V p62 were higher than those in cells overexpressing WT p62, suggesting that M404V p62 may increase autophagic activity, which is consistent with Fig 4.3 B. However, since
LC3-II reached maximal capacity after treatment with NH$_4$Cl, a solid conclusion regarding the impact of p62 PDB mutant on autophagy remains unclear.

Discussion

Challenges of current methods for studying the effect of PDB-associated p62 mutants on autophagy

In this Chapter, we mainly used p62 KO MEF cells to study the effect of PDB-associated mutants on autophagy. We used two common methods for the study. One is comparing the GFP-LC3 puncta in cells overexpressing WT or mutant p62, the other is comparing the LC3-II levels in cells overexpressing WT or mutant p62 after treatment with an autophagy inducer and/or inhibitor.

P62 KO MEF cells have the advantage of producing no endogenous p62. However, it is not an osteoclast-like cell. For quantitative study of GFP-LC3 puncta, since there will be a large number of cells and puncta for counting, a better program or software is needed to count the puncta using the same criteria for cells over-expressing WT or mutant p62.

Another challenge is that p62 is also a substrate of autophagy. Therefore it is a little complicated to explain the regulatory role of p62. There are two possibilities. If mutant p62 increases autophagic activity, the levels of the mutant p62 will decrease with time, self limiting the stimulation of autophagy. If mutant p62 decreases autophagic activity, the levels of the mutant p62 will accumulate through the time. Thus, autophagic activity
will continue to decrease with time. The second scenario might be the case, so the central hypothesis of our study would be: PDB-associated p62 mutants impair the autophagic activity, leading to accumulation of p62, which then further impairs the autophagic activity. Meanwhile, accumulated p62 mutants further increase the NF-κB signaling and the formation of osteoclasts, which leads to PDB (Fig. 5.1).

**Caution should be exercised when explaining LC3-II data**

LC3-II is widely used to estimate autophagy activity [152]. However, we need to very carefully explain LC3-II data. When autophagic activities increase, LC3-II levels increase because more autophagosomes form. However, LC3-II itself is degraded by autophagy, the increased autophagic activity also lead to more degradation of LC3-II, therefore decreasing the LC3-II levels. Therefore, it is important to measure LC3-II in the presence and absence of lysosomal protease inhibitors (e.g. E64d and pepstatin A) or inhibitors which block the fusion of lysosomes and autophagosomes (e.g. NH₄Cl and Bafilomycin A1). If mutant p62 really increases the autophagic activity, we will see LC3-II levels increases after treatment of either lysosomal protease inhibitors or inhibitors blocking the formation of autophagolysosome in cells overexpressing mutant p62 compared with cells overexpressing WT p62.

In our study, we used rapamycin as an inducer of autophagy and NH₄Cl as an inhibitor. We expected to see LC3-II levels increase upon rapamycin treatment because more autophagosomes form. However, LC3-II decreased in P2 in the control cells after rapamycin treatment (Fig. 4.3 A, lane 2 and 6), suggesting that while rapamycin may
increase the formation of autophagosome, this may ultimately decrease LC3-II levels through increased degradation. In addition, cells treated with NH₄Cl generated more LC3-II (Fig. 4.3 A, lane 9-12), suggesting that NH₄Cl is a good inhibitor for blocking the fusion of lysosome and autophagosome. Also, it seems that LC3-II reached maximal capacity after treatment with NH₄Cl (Fig. 4.3 A, lane 9-12). In future, it is possible that using lower NH₄Cl levels may allow useful measurements of changes in LC3-II levels.

**Significance of this study and future directions**

In this Chapter, we examined the effect of PDB-associated p62 mutants on interaction with LC3, GFP-LC3 puncta formation and LC3-II levels upon autophagy inducer and/or inhibitor treatment. Our data have shown that p62 PDB mutants did not change interaction with LC3 compared with WT p62. In addition, we speculate that mutant p62 might impair autophagic activity, leading to the accumulation of p62.

For future studies, in order to better answer whether p62 PDB mutants change autophagic activity compared with WT p62, methods must be developed to quantify GFP-LC3 puncta. Also, for the LC3-II study, other autophagy inducers (Beclin-1, starvation) and autophagy inhibitors (protease inhibitors E64d and pepstatin A) could be used. In addition, the question of whether autophagy change the rate of degradation of p62 PDB mutant compared with WT p62 has not been addressed. In order to address this question, the half life of p62 could be monitored.
Figure 4.1. Comparison of the effect of WT p62 or PDB mutant p62 on the interaction of LC3. (A) HEK293 cells were transfected with FLAG-p62 (WT, A381V, P392L and M404V) and EGFP-LC3. FLAG-IP was performed, followed by the SDS-PAGE. Western blotting was performed using the antibodies against the EGFP and FLAG. (B) Cell extracts of different samples.
Figure 4.2. Immunostaining of p62 and LC3 in p62 KO MEF cells overexpressing p62 and GFP-LC3. P62 KO MEF cells were transfected with FLAG-p62 (WT or M404V) and EGFP-LC3. Twenty-four hours after transfection, cells were fixed with 4% PFA and permeabilized with 0.1% Triton. Primary antibody FLAG (1:230) and secondary antibody Hoechst (blue, 1:1000), Alexa Fluor 594 (mouse) (red, 1:300) were used. Confocal microscopy was applied for observation. Scale bars=20μm. The nuclear staining of GFP-LC3 is likely due to high levels of overexpression.
Figure 4.3. Comparison of the effect of WT p62 and p62 PDB mutant on LC3-II level after rapamycin or rapamycin/NH₄Cl treatment. (A) Western blotting of LC3-II in S1, S2 and P2 after fractionation in p62 KO MEF cells overexpressing WT or M404V p62. P62 KO MEF cells were transfected with FLAG-p62 (WT or M404V). The cells were treated with rapamycin (200nM) alone for 16 hours or NH₄Cl (10mM) and rapamycin (200nM) together for 16 hours. Forty-eight hours post transfection, the cells were harvested with RIPA buffer. The cell lysates were fractionated into S1, S2 and P2. The samples were subjected to the SDS-PAGE, followed by Western blotting using the antibodies against the LC3, FLAG and actin. NT: non-transfection. (B) Quantification of LC3-II/Actin in S1 treated with rapamycin alone by Image J. The value of WT p62 is normalized to 100%. Percentage of decrease after rapamycin is shown in red. (C) Quantification of LC3-II/Actin in S1 treated with rapamycin/NH₄Cl by Image J. The value of WT p62 is normalized to 100%. Percentage of increase after rapamycin/NH₄Cl is shown in red.
Significance of this research

In Chapter 2, we introduced a relatively simple proteomic method to identify novel SUMOylation substrates. Seventy-four SUMOylated proteins were identified by our proteomic analysis (Table 2.1). Among these proteins, Ran GTPase-activating protein 1 (RanGAP1) [12], proliferating cell nuclear antigen (PCNA) [169] and nucleophosmin (NPM) [168] have been reported to be SUMOylation substrates. In addition, 61 proteins have predicted SUMOylation sites and 40 proteins have SUMOylation consensus sequences (ΨKXE/D) (Table 2.1). Over 50% of these proteins are nuclear proteins (Fig. 2.4), which is consistent with literature that most SUMOylation substrates are in the nucleus [2]. Altogether, this suggests that our proteomic method is applicable for identification of SUMOylated substrates.

Moreover, we are the first group to identify a novel SUMOylation substrate called Drebrin, an actin-binding protein located in the cytosol [3, 45]. We verified Drebrin SUMOylation by both FLAG and HA immunoprecipitation (Fig. 2.8 A and B). Furthermore, by using site-directed mutagenesis, we found K185, K186, K270 and K271 might be SUMOylation sites for Drebrin (Fig. 2.9 B). These lysines are conserved among vertebrate Drebrins by alignment of this region among different species (Fig. 2.7 C), suggesting that they are functionally important. At last, we observed the protrusion formation in CHO cells overexpressing Drebrin (WT and mutants) (Fig. 2.10 A), which is consistent with the literature [53, 54, 57]. However, mutation of these two sites
(K185R/K186R and K270R/K271R) in Drebrin separately did not appear to change protrusion formation (Fig. 2.10 B).

Additionally, we also identified several nuclear proteins with multiple predicted SUMOylation sites (highlighted in Table 2.1), such as general transcription factor II-I, heterogeneous nuclear ribonucleoproteins C1/C2, nucleolin and 60S ribosomal protein L24. We could further validate SUMOylation of these proteins by IP and other methods in the future.

In Chapters 3 and 4, our studies were related to “ubiquitination”, a similar modification of “SUMOylation”. We focused on an ubiquitin-binding protein, p62. We studied the effect of PDB-associated p62 mutants on NF-κB signaling and autophagy. In Chapter 3, we are the first group to use p62 KO MEF cells as a model for studying the effect of p62 PDB mutant on TNFα-induced NF-κB signaling. Wooten et al. [95] and Moscat et al. [93] have shown that p62 activated the NF-κB signaling in HEK293 cells basally. Our study not only showed that p62 activated NF-κB signaling basally (Fig. 3.6 C), but also upon TNFα treatment in p62 KO MEF cells (Fig. 3.9 A) and HEK293 cells (Fig. 3.10 A). In addition, we also have shown that p62 PDB mutants have a tendency to increase TNFα-induced NF-κB signaling compared with WT p62 in p62 KO MEF cells (Fig. 3.9 B and C) and HEK293 cells (Fig. 3.10 B). In addition, our data indicated that p62 did not increase TNFα-induced NF-κB signaling through increasing TRAF6 polyubiquitination (Fig. 3.14), suggesting other mechanisms may exist (see below).

In Chapter 4, we studied the impact of PDB-associated p62 mutants on autophagy. We found that PDB mutations did not change the interaction between p62 and the
autophagy marker protein LC3 (Fig. 4.1 A). Whether p62 PDB mutants change the autophagic activity is not certain until we optimize our experiments by using autophagy inducer/inhibitor appropriately.

In summary, my work has expanded our knowledge of the role of PDB-associated p62 mutants on NF-κB signaling and autophagy, as well as provided insights into several possible mechanisms, which are shown in an integrated model in Fig. 5.1.

**Functional consequences of SUMOylation of Drebrin and other proteins**

Although we detected SUMOylation of Drebrin, the modification does not appear to be involved in protrusion formation in CHO cells (Fig. 2.10 B). So what are the functional consequences of Drebrin SUMOylation? In Chapter 1, we introduced that SUMOylation modification of certain protein could bind other proteins only when SUMOylation is present. For example, only SUMOylated Ran-GAP1 binds RanBP2 [12]. Therefore, we hypothesized that Drebrin SUMOylation might also bind some proteins only when SUMOylation is present. So how to identify these target proteins which bind SUMOylated Drebrin, not non-SUMOylated Drebrin? We could take advantage of mass spectrometry to do the quantification for Drebrin interacting proteins from cells overexpressing WT Drebrin and mutant Drebrin. We will expect to see the amounts of some Drebrin interacting proteins are significantly higher in cells overexpressing WT Drebrin than Drebrin mutants, which impair Drebrin SUMOylation. These proteins are potential Drebrin interacting proteins only when Drebrin is SUMOylated. We will further verify their interaction by immunoprecipitation and Western blotting.
In addition, we could further validate the SUMOylation of proteins with higher predicted SUMOylation sites, which are highlighted in Table 2.1 by IP and other methods. We could also study the functional consequences of these proteins in the future.

**Mechanisms that PDB-associated p62 mutations increase NF-κB signaling**

In Chapter 3, we have shown that PDB-associated p62 mutations (M404V, A381V and P392L) had a tendency to increase TNFα-induced NF-κB signaling in p62 KO MEF cells (Fig. 3.9) and HEK293 cells (Fig. 3.10). Previously, we hypothesized that p62 PDB mutant increased TNFα-induced NF-κB signaling through increasing TRAF6 polyubiquitination. However, we showed that TNFα did not increase TRAF6 polyubiquitination in HEK293 cells overexpressing WT and mutant p62 (Fig. 3.14), which did not support our hypothesis. Therefore, by what other mechanisms do p62 PDB mutants increase NF-κB signaling? Several possibilities are described below and summarized in Fig. 5.1.

1. **TRAFs or RIP polyubiquitination**

Moscat *et al.* [93] showed that p62 could interact with RIP. They also showed that p62 interacted with TRAF2 in the presence of RIP by immunoprecipitation experiment in HEK293 cells transfected with p62, RIP and TRAF2 constructs. In their study, they also showed that upon TNFα treatment, p62 and RIP could be co-immunoprecipitated with TNF-R1. This suggested that the interaction of p62 and RIP plays a role in the TNFα-induced NF-κB signaling. In Chapter 1, we also introduced that upon TNFα treatment,
TRAF2 undergoes auto-polyubiquitination (K63-linked), which catalyzes RIP K63-linked polyubiquitination and activates the TNFα signaling [93] (Fig. 1.5). Our data showed that p62 contributed to TNFα-induced NF-κB signaling in p62 KO MEF cells (Fig. 3.9 A) and HEK293 cells (Fig. 3.10 A) by luciferase assay. Therefore, we hypothesize that WT p62 increases TNFα-induced NF-κB signaling by increasing TRAF2 or RIP polyubiquitination. Based on our data that p62 PDB mutants had a tendency to increase TNFα-induced NF-κB signaling (Fig. 3.9 and 3.10), we also hypothesize that p62 PDB mutant increases TNFα-induced NF-κB signaling by increasing TRAF2 or RIP polyubiquitination compared with WT p62.

In order to test our hypothesis, we should first set up a model system in which TRAF2 or RIP polyubiquitination increases upon TNFα, and the signaling proteins IKK increase, IκB decreases, phospho-IκB increases after TNFα treatment. Then, we will test whether TRAF2 or RIP polyubiquitination increases in the presence of p62. If so, we will have demonstrated that p62 involves in TRAF2 and RIP polyubiquitination. Next, we will compare the effect of WT p62 and p62 PDB mutant on TRAF2 and RIP polyubiquitination, see if mutant p62 could further increase polyubiquitination. If these experiments have results as expected, we could further examine TRAF2 or RIP polyubiquitination using a variety of different Ub constructs. If we find that TRAF2 or RIP polyubiquitination is a mixed linkage polyubiquitin chain, not a K63-linked polyubiquitin chain as we found for TRAF6 (Fig. 3.15), it will be a new finding to the field, that the mixed linkage polyubiquitin chain could also successfully activate the signaling.
In addition, when the model for studying RANKL-induced NF-κB singnaling is available in the future, we could test whether PDB-associated p62 mutations increase RANKL-induced NF-κB signaling through increasing TRAF6 polyubiquitination.

(2) aPKC

In the NF-κB signaling pathway (Fig. 1.5), it is shown that p62 directly interacts with aPKC through its PB1 domain, and further activates IKK, leading to NF-κB activation. Therefore, we propose that PDB-associated p62 mutants increase the interaction with aPKC or increase the activity of aPKC compared with WT p62, and activate the signaling.

Duran et al. [77] demonstrated RANKL ligand-dependent interactions between p62 and aPKC by immunoprecipitating endogenous p62 in Raw264.7 cells. However, they did not show negative controls in their IP. In our study, we performed the same experiment as they did, however our negative control was not working, since HA antibody could also pull down the endogenous aPKC in the absence and presence of RANKL (data not shown). Also, the IP efficiency is only about 13%. In the long run, we could use better IP buffer or better HA antibody to improve the IP efficiency and diminish the effect of non-specific interaction. We could also use fusion protein p62 or aPKC (GST, GFP, Myc, FLAG, or DsRed tagged protein) to study the interaction. Once the experiment is set up, we could introduce WT p62 and p62 PDB mutants, and test if interactions with aPKC are changed. We could also monitor signaling proteins, such as IKK, IκB and phospho-IκB, and see if the activity of aPKC increases in the presence of mutant p62.
(3) CYLD

Jin et al. [176] reported that a deubiquitinating enzyme, CYLD, negatively regulates osteoclastogenesis and RANK signaling. They have shown that RANKL induced a much higher number of ostoclasts from the bone marrow-derived macrophages (BMDMs) in CYLD knockout mice than control mice. They also showed that ubiquitinated TRAF6 was accumulated in BMDMs from CYLD knockout mice. In addition, they showed that CYLD could interact with WT p62, but not UBA-deleted p62 by IP, indicating that the interaction requires the UBA domain. Also, they showed that the interaction of CYLD and TRAF6 is dependent on p62 and its UBA domain by IP.

In light of their work, we hypothesized that p62 PDB UBA mutant decreases interaction with CYLD, leading to decreased deubiquitination of polyubiquitinated TRAF6, further increasing NF-κB signaling. To test this hypothesis, HEK293 cells were transfected with human HA-CYLD [177] and human FLAG-p62 (WT and mutants) or mouse p62 (WT and UBA domain deleted). Forty-eight hours after transfection, FLAG-p62 IP was performed. In contrast to Jin et al. [176], we found that CYLD co-immunoprecipitated with mouse UBA-deleted p62 (data not shown). In their study, they used the EGFP-p62, and we used FLAG-p62 in our study. In the future, to rule out non-specific interactions, we could further change other tags (GST, GFP, Myc and DsRed), or use the human UBA-deleted p62, not mouse, as a better negative control.

A recent study reported that the de-ubiquitination enzyme CYLD interacted with wild-type and a non-UBA mutant A381V p62 in osteoclast progenitor cells, but not to the UBA mutant P392L p62 [178]. Expression of p62 P392L also resulted in increased levels
of polyubiquitinated TRAF6 and phospho-IκB during osteoclast differentiation. These findings suggest that at least some p62 PDB mutations might perturb NF-κB signaling by altering CYLD activity and TRAF6 polyubiquitination.

(4) Other signaling proteins

Our data showed that p62 PDB mutants (M404V, P392L and G411S) suppressed its association with polyubiquitinated proteins (Fig. 3.11). Layfield et al. [116] also reported that p62 PDB mutants impaired K48-linked polyubiquitin binding in vitro. In Chapter 1, we introduced that p62 could bind polyubiquitinated proteins and target them for degradation by both UPS and autophagy (Fig. 1.6) [8]. Therefore, p62 PDB mutations may impair their binding with polyubiquitinated proteins that are active signal proteins in the NF-κB signaling pathway, leading to decreased protein degradation of these proteins, which might increase the NF-κB signaling. To test this hypothesis in the future, mass spectrometry could be applied for identification of these important targeting signaling proteins. In detail, we could use mass spectrometry to do the quantification for the amount of proteins in cells overexpressing WT p62 and p62 PDB mutant (Fig. 3.11). We will expect to see that the amounts of some signaling proteins are significantly higher in cells overexpressing p62 PDB mutant than WT p62. These proteins could be candidate signaling proteins regulated by p62. When these certain proteins are polyubiquitinated, p62 PDB mutants fail to bind with these polyubiquitinated proteins, leading to less capable of sequestering these proteins for degradation. We would confirm these proteins by IP in cells overexpressing Ub and individual candidate proteins. In addition, p62 PDB
mutants could also impair autophagy activity leading to accumulation of important
signaling proteins, and further activating NF-κB signaling (see below).

**Autophagy in PDB**

In a recent autophagy review, it is suggested that autophagy may be related to the
development of bone diseases, although the physiological roles of autophagy in bone are
still mostly unknown [85]. In addition, the presence of inclusion bodies in osteoclasts
seems to link PDB to autophagy [7]. Moreover, it is found that p62 colocalizes with
Autophagy-Linked FYVE-domain containing protein (ALFY) in osteoclasts [7]. While
this suggests that autophagy is linked to PDB [179], little is known about the role of
autophagy in PDB so far. Therefore, the role of autophagy is clearly an area that merits
investigation in the future.

Here, I offer some speculation of the role of autophagy based on the literature and
our preliminary data in Chapter 4. P62 accumulation has been reported in French PDB
patients with and without p62 mutations [175], suggesting that autophagy is impaired in
PDB since p62 is a substrate for autophagy [147]. If impairment of autophagy is
confirmed in a large number of PDB patients, what are the mechanisms?

Defective autophagy in PDB patients could be related to PDB-associated p62
mutants. As described in Chapter 1, p62 mutations are found in about 40% of familiar
PDB [7]. Additionally, p62 is both a substrate and regulator of autophagy [8], therefore
do PDB-associated p62 mutations contribute to the impairment of autophagy in PDB? In
Chapter 4, we tried to use p62 KO MEF cells and several autophagy inducers and/or
inhibitors to test this hypothesis (Fig. 4.3). In preliminary experiments with cells overexpressing WT p62, the LC3-II level decreased less than the control (7.90% vs. 23.20%) after rapamycin treatment (Fig. 4.3 B), suggesting that overexpression of WT p62 attenuates the autophagic activity. Here, we propose that p62 PDB mutants could decrease the autophagic activity compared with WT p62, leading to accumulation of p62, which will further decrease the autophagic activity. In the future, this hypothesis could be tested by better experiment design, including better software to analyze the GFP-LC3 puncta, more effective and appropriate usage of autophagy inducer and/or inhibitor and monitoring the endogenous and exogenous p62 levels as indicators of autophagy.

A recent review paper Goode et al. [9] suggests that defective autophagy and dysregulated NF-κB signaling in PDB may be linked. A number of signaling intermediaries of the RANKL-induced NF-κB signaling, such as IKK, are targets of ubiquitination [9, 62], which might be degraded by autophagy. Therefore, it would be interesting to determine the effect of PDB mutations on p62-mediated autophagic protein degradation of signaling intermediaries in the NF-κB signaling. In order to address this question, we could monitor the signaling proteins, including IKK, IκB and phospho-IκB, in cells treated with autophagy inhibitor/inducer in the absence and presence of RANK or TNFα.

Considering our findings of p62 PDB mutant on NF-κB signaling, a new proposed model showing that PDB-associated p62 mutants increase NF-κB signaling by several of the mechanisms is described in Fig. 5.1. P62 PDB mutants could also impair autophagy activity leading to accumulation of important signaling proteins, and further activating NF-κB signaling. Meanwhile, defective autophagy could lead to accumulation
of mutant p62, which would further impair autophagy and upregulate the NF-κB signaling, creating a feed-forward pathological cycle. This combined model is presented in Fig. 5.1.

Other methods and models for studying the PDB

In Chapter 3, we compared cell models for studying the impact of PDB-associated p62 mutants on NF-κB signaling (Table 3.1). Among them, Raw264.7 cells might be the best one since they are osteoclast-like cells [170]. The main technique we used is the NF-κB luciferase assay.

Aside from Raw264.7 cells and NF-κB luciferase assays, people used other methods and materials for studying PDB [80, 170]. Additional methods include bone resorption pit assay [80, 170], in vitro osteoclastogenesis assay [170], osteoclast apoptosis [80] and others. In addition, besides NF-κB luciferase assays, people used other methods to examine the NF-κB signaling. These methods include electrophoretic mobility shift assay (EMSA) [180, 181], nuclear:cytoplasmic ratios of NF-κB [182], monitoring the rate of IκB degradation [80] and p65 nuclear translocation [183]. In our study, we monitored the rate of IκB degradation (Fig. 3.3). However, our assay needs to be optimized by co-tranfection of HA-IκB (or other tags) with the FLAG-p62, in order to monitor the IκB in cells expressing FLAG-p62 only. In addition, we also tried p65 nuclear translocation in our study (Fig. 3.4). Again, we need to determine how to quantify p65 nuclear translocation between cells expressing WT p62 and mutant p62 in the future.
Mouse models including mice expressing human P392L p62 [184] and P394L mutant mice [185] were reported previously. Isolated bone marrow macrophages (BMM) from WT and p62 mutant mice [170], isolated osteoclast precursors from healthy donors and PDB patients [184], cord blood monocyte (CBM) and peripheral blood mononuclear cells (PBMC) isolated from blood provided by healthy donors and PDB patients [80] were also applied in other people’s work.

Compared with our mutant p62 overexpressing in cells, using cells from PDB patient has its advantage of natural background of the PDB-associated p62 mutants. In the future, we may learn from these other methods and models.

SUMOylation and NF-κB signaling

In Chapter 2, we mainly focused on identification of novel SUMOylation substrates. In Chapter 3, we investigated the PDB-associated p62 mutants in NF-κB signaling, which involves TRAFs polyubiquitination. The linkage of Chapter 2 and Chapter 3 seems to be that both SUMOylation and ubiquitination are important post-translational modifications which also share similarities [2].

In addition, several signaling proteins in the NF-κB signaling pathway are reported to be SUMOylation substrates, such as TNF-R1 [35], IκB [30] and NF-κB essential modulator (NEMO) [186], the IKK regulatory subunit. Therefore, does SUMOylation play a role in the NF-κB signaling related to PDB? What is the interplay of SUMOylation and ubiquitination in the NF-κB signaling related to PDB? Could SUMOylation be
involved in the mechanisms of PDB? We could expand our knowledge of PDB by answering these questions.

**An integrated model: speculation about the role of PDB-associated p62 mutation**

In light of a recent review paper Goode *et al.* [9], as well as our work and the literature, I proposed a model integrating the role of PDB-associated p62 mutants on both NF-κB signaling and autophagy (Fig. 5.1).

PDB-associated p62 mutants increase NF-κB signaling through several mechanisms including increasing aPKC activity, decreasing CYLD activity, increasing TRAF6/TRAF2 polyubiquitination, as well as increasing activity of unknown signaling proteins. Autophagy deficiency was detected in PDB patients, leading to the accumulation of mutant p62, which is proposed to further impair autophagy (dash line). P62 accelerates disease progression by upregulating NF-κB signaling through one or several of the mechanisms. Therefore, PDB-associated p62 mutants play a synergistic role in disease progression by affecting both NF-κB signaling and autophagy. The impairment of autophagy in PDB patient results in accumulation of mutant p62, which accelerates disease progression by increasing the NF-κB signaling. SUMOylation was found on several NF-κB signaling proteins including IκB [30], TNF-R1[35] and NEMO [186]. SUMOylation would therefore be involved in other mechanisms of PDB.
A proposed model for the role of PDB mutant p62 in NF-κB signaling and autophagy. PDB-associated p62 mutants increase NF-κB signaling through several mechanisms including increasing aPKC activity, decreasing CYLD activity, increasing TRAF6/TRAF2 polyubiquitination, as well as increasing activity of unknown signaling proteins. Autophagy deficiency was detected in PDB patients, leading to the accumulation of mutant p62, which is proposed to further impair autophagy (dash line). P62 accelerates disease progression by upregulating NF-κB signaling through several mechanisms. Therefore, PDB-associated p62 mutants play a synergistic role in disease progression by affecting both NF-κB signaling and autophagy. The impairment of autophagy in PDB patient results in accumulation of mutant p62, which accelerates disease progression by increasing the NF-κB signaling. In addition, SUMOylation would be involved in other mechanisms of PDB.
APPENDICES

Appendix I: List of all constructs

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<td>Human Drebrin K185R/K186R/K192R/K270R/K271R</td>
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<td>pcDNA-MYC</td>
<td>Dr. Marie Wooten</td>
</tr>
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<td>p3xFLAG-CMV10</td>
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</tr>
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<td>p3xFLAG-p62 (mouse), delta UBA</td>
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<td>Chapter 4, made by Dr. Jozsef Gal [83]</td>
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<tr>
<td>p3xFLAG-TRAF6</td>
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<td>Mouse</td>
<td>p3xFLAG-CMV10</td>
<td>Chapter 3 and 4 Made by Dr. Jozsef Gal</td>
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<td>NF-κB luciferase reporter</td>
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<td>pCMV</td>
<td>Chapter 3 and 4, Dr. Jiake Xu [170, 173].</td>
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<tr>
<td>GST-rRANKL</td>
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<td>pCMV</td>
<td>Chapter 3, Dr. Jiake Xu [171]</td>
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<th>Mutation</th>
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<th>Chapter, Maker</th>
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<td>pDEST-HA</td>
<td>Addgene Plasmid 15506 [177].</td>
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## Appendix II: List of amplification primers and mutagenic primer sequences.

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<tr>
<th>Number</th>
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<tr>
<td>GJ464</td>
<td>T375A reversion primer in the Wooten human p62 clone</td>
<td>5’-CCTTCAGCCCTGTGGGTCCCTCCTCCTCTTG-3’</td>
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<td>GJ465</td>
<td>T375A reversion primer in the Wooten human p62 clone</td>
<td>5’-CAGGAGGGACCCACAGGGCTGAAGG-3’</td>
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<td>GJ468</td>
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<td>5’-GCTGGAATTCCGCCTCGCCTACCCTGAAG-3’</td>
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<td>GJ469</td>
<td>Human p62 downstream, cont. STOP (KpnI)</td>
<td>5’-CGTCGGAATTCACCAACGGGCGGGGATG-3’</td>
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<td>GJ470</td>
<td>Human p62 internal seq. primer</td>
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<td>Human p62 D335E QuikchangeII primer, upper strand</td>
<td>5’-ACTGTTGCCAGGAGGAAGATGACTGGACCCCATC-3’</td>
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<td>GJ473</td>
<td>Human p62 A381V QuikchangeII primer, upper strand</td>
<td>5’-GGCTGAAGGAAGCCGATTCCACATCT-3’</td>
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<td>GJ474</td>
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<td>Human p62 M404V QuikchangeII primer, upper strand</td>
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<tr>
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<td>5'-CCTGGTGAGCCAGCTGCTCATCAGAGA-3'</td>
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<td>hDrebrin Quikchange Multi, K185R K186R, upper strand</td>
<td>5'-GTTCTGGAGCAAGCCAGGGGAAGAAGAGCTGC-3'</td>
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<td>GJ758</td>
<td>hDrebrin Quikchange Multi, K185R K186R, lower strand</td>
<td>5'-GCAAGCTTCTTCTCTCCCTGCGCTCAGAAGAC-3'</td>
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<td>GJ759</td>
<td>hDrebrin Quikchange Multi, K192R, upper strand</td>
<td>5'-AGAAGAGCTGAGGAGGGGAAGAGGC-3'</td>
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<td>hDrebrin Quikchange Multi, K192R, lower strand</td>
<td>5'-CGCTCCTCCTCCCTCCGCAGCCTTCT-3'</td>
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<td>GJ761</td>
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<td>5'-GAAGAGACCACATGAGGGAGTCAGAGTCGGAGGTG-3'</td>
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<td>GJ762</td>
<td>hDrebrin Quikchange Multi, K270R K271R, lower strand</td>
<td>5'-CACCTCCGACTCTGACCTGAGAGTCAGCC-3'</td>
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<td>GJ763</td>
<td>hDrebrin internal sequencing primer</td>
<td>5'-AAGACGGATGCAGCTGTGGA-3'</td>
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<td>GJ764</td>
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<td>5'-CGTCGAAGTTCTGCCGCGTCAGCTTCAG-3'</td>
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<td>GJ765</td>
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<td>5'-CTGGGAGCAGGCCAGGAAGAAGAGAC-3'</td>
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<td>GJ770</td>
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<tr>
<td>GJ771</td>
<td>hDrebrin K186R Quikchange II, upper strand</td>
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<tr>
<td>GJ772</td>
<td>hDrebrin K186R Quikchange II, lower strand</td>
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<tr>
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<td>5'-GAAGAGACCCACATGAGAGTCAGAGTCGGAG-3'</td>
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<td>GJ774</td>
<td>hDrebrin K270R Quikchange II, lower strand</td>
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<td>GJ775</td>
<td>hDrebrin K271R Quikchange II, upper strand</td>
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<tr>
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<td>5'-CACCTCCGACTCTGACCTCTTCATGTGGTTC-3'</td>
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Appendix III: List of abbreviations

ALS: Amyotrophic lateral sclerosis
aPKC: atypical protein kinase C
CHO: Chinese hamster ovary
Drebrin: Developmentally regulated brain protein
EMSA: Electrophoretic mobility shift assay
ERK: Extracellular responsive kinase
FL: Full-length
GFP: Green fluorescent protein
HA: Hemagglutinin
HEK: Human embryonic kidney
HRP: Horseradish peroxidase
hSUMO-1: human SUMO-1
IKK: IκB kinase
IκB: Inhibitor of NF-κB
KO: Knockout
LC3: Microtubule-associated protein light chain 3
LIR: LC3 interaction region
MEK5: MAPK/ERK kinase 5
MEF: Mouse Fibroblast
NBR1: Neighbor of BRCA1 gene 1
NEMO: NF-κB essential modulator
NF-κB: Nuclear factor κB
NPC: Nuclear pore complex
IP: Immunoprecipitation
PDB: Paget’s disease of bone
PB1 domain: Phox and Bem1 domain
PE: Phosphatidylethanolamine
PML: Promyelocytic leukemia
RANK: Receptor Activator of Nuclear Factor κB
Ran-GAP1: Ran-GTPase-activating protein 1
RIP: Receptor-interacting protein
SDS-PAGE: Sodium dodecyl sulfate polyacrylamide gel electrophoresis
SENP: Sentrin/SUMO-specific protease
SUMO: Small Ubiquitin-like Modifier protein
TB domain: TRAF6 binding domain
TBST: Tris-Buffered Saline and Tween 20
TNFα: Tumor Necrosis Factor-alpha
TRADD: TNFR1-associated death domain protein
TRAF6: TNF receptor associated factor 6
TrkA: Tropomyosin-receptor- kinase
UBA domain: Ubiquitin-association domain
UPS: Ubiquitin-proteasome system
WT: Wild-type
REFERENCES


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EDUCATION
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1995.9-1999.6 NO. 45 Middle School of Hubei Province, Wuhan, Hubei, China

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Graduate
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Undergraduate
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2004 “Excellent Academic Students” for NO.1 student in my grade. 3000 RMB
2003 “Excellent Academic Students” for NO.1 student in my grade. 3000 RMB

PUBLICATIONS

Graduate


XiaoYan Liu, Fujian Zhang, Jianjun Zhai and Haining Zhu. Identification of novel SUMOylated proteins in mammalian cells (ready to submit).


Undergraduate


TEACHING
2008 Spring Semester
Biochemistry 401G FUNDAMENTALS OF BIOCHEMISTRY: DNA Replication, Repair and Recombination (undergraduate)

ACTIVITY
- Chairperson of oral presentation sessions for five years at Biochemistry Departmental Retreat (2007-2011)
  2007-2008 Pine Mtn. State Resort Park, Pineville, Kentucky
  2009-2011 Cumberland Falls State Park, Corbin, Kentucky
- Organizer for the Ice Cream Social for Biochemistry Department, July, 2010
- Organizer for the Holiday Food Drive Donation (2009, 2011)
MEMBERSHIP
2010-Present  Member of American Society for Mass Spectrometry (ASMS)

PRESENTATIONS

One invited talk

Four student seminars of department


One invited data club
Xiaoyan Liu. Data Club. BBSRB B231, Feb 27th, 2009

Five posters in the department summer retreat


disease of bone. Cumberland Falls State Park, Corbin, Kentucky; May, 2009.


Six posters in the conferences


Xiaoyan Liu, Fujian Zhang, Haining Zhu. Proteomic Analysis of SUMOylated Protein in Mammalian Cells. ASBMB Conference, San Diego, California; April, 2008.


