

Review Article

Cycle on Wheels: Is APP Key to the AppBp1 Pathway?

Chen Y^{1,2*}, Neve RN⁴, Zheng H³, Griffin WST^{1,2}, Barger SW^{1,2} and Mrak RE⁵¹Department of Geriatrics, University of Arkansas for Medical Sciences, USA²Department of Neurobiology and Developmental Sciences, University of Arkansas for Medical Sciences, USA³Huffington Center on Aging, Baylor College of Medicine, USA⁴Department of Brain and Cognitive Sciences, Massachusetts Institute of Technology, USA⁵Department of Pathology, University of Toledo Health Sciences Campus, USA

*Corresponding author: Chen Y, Department of Geriatrics and Department of Neurobiology and Developmental Sciences, University of Arkansas for Medical Sciences, Little Rock, AR 72205, USA

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Abbreviations

APP: Amyloid Precursor Protein; AICD: APP Intracellular Domain; A β : β -Amyloid; NAE: Nedd8 Activating Enzyme, i.e. AppBp1 and Uba3 heterodimer; CCM: Cell Cycle Marker; C57: APP's C-Terminal 57 amino acids; AppBp1: APP-Binding Protein-1, also known as APP-BP1 or NAE1; DDB1: UV-Damaged, DNA-Binding protein 1; DCAF: DDB1-Cul4-associated factor

Introduction

The Amyloid Precursor Protein (APP) is central to understanding Alzheimer's disease (AD) pathogenesis due to its genetic, biochemical and neuropathological connections with AD. First, APP is the source of β -amyloid (A β), a major component of senile plaques in AD brains. Secondly, Genetic mutations of APP cause familial AD [1,2]. Furthermore, an increase in APP gene dosage also causes A β deposition and related dementia [3-5]. In the case of trisomy of chromosome 21, AD neuropathology develops universally due to an extra copy of APP [6]. Therefore, elucidating the function of APP may give insights into disease prevention and treatment strategies.

APP is a Type-1 transmembrane receptor [7-10], which is cleaved by multiple proteases. Sequential cleavage by β -secretase BACE1 at the extra-cellular domain [11-13] followed by γ -secretase cleavage inside the membrane [14-16] generates secreted APP N-terminal fragment sAPP β , A β peptides of various length, and APP intracellular domain (AICD) [17-19]. Alternatively, cleavage of APP by the α -secretase within the A β domain [20,21] followed by γ -secretase cleavage generates sAPP α , p3, and AICD. Despite intensive studies of APP [22-24] and its cleavage products, the function of APP remains poorly understood.

Molecular basis for APP in cell cycle regulation

In an effort to understand APP as a potential signaling receptor,

Abstract

Alzheimer's disease (AD) is the gradual loss of the cognitive function due to neuronal death. Currently no therapy is available to slow down, reverse or prevent the disease. Here we analyze the existing data in literature and hypothesize that the physiological function of the Amyloid Precursor Protein (APP) is activating the AppBp1 pathway and this function is gradually lost during the progression of AD pathogenesis. The AppBp1 pathway, also known as the neddylation pathway, activates the small ubiquitin-like protein nedd8, which covalently modifies and switches on Cullin ubiquitin ligases, which are essential in the turnover of cell cycle proteins. Here we discuss how APP may activate the AppBp1 pathway, which downregulates cell cycle markers and protects genome integrity. More investigation of this mechanism-driven hypothesis may provide insights into disease treatment and prevention strategies.

Keywords: APP; Alzheimer's disease; Ubiquitination; Neddylation; Cell cycle

at least 18 proteins have been identified to bind AICD [25-27]. Among them is APP-binding protein-1 (AppBp1), which binds APP's C-terminal 57 amino acids (C57) [28]. Co-immunoprecipitation experiments further defined AppBp1's binding site to two segments of C57: one is adjacent to the membrane including three lysine residues and the other in the C-terminal 31 amino acids [29,30] (Figure 1A). The function of AppBp1 was unknown when it was cloned as an APP-binding protein. The significance of the interaction emerged when AppBp1 was discovered as a cell cycle protein. The first evidence was obtained from hamster ts41 cells, which harbor a temperature-sensitive mutant of AppBp1's homologue, ts41 [31,32]. At the non-permissive temperature of 40°C, ts41 cells undergo apoptosis after

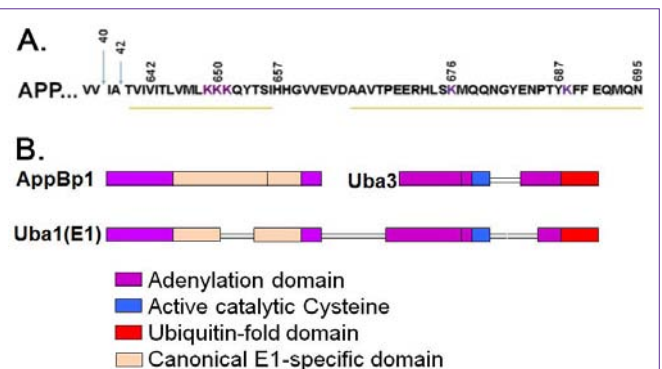


Figure 1: APP binds to AppBp1 in a region that mediates adenylation and nedd8 binding to NAE. A. C-terminal sequence of APP695 showing the C-terminus of A β 40 and A β 42, lysine residues (purple) that can be neddylated and the AppBp1 binding sites in APP AICD region (yellow lines). B. Diagram of conserved domains between NAE and Uba1. The APP-binding site, AppBp1 (145-251) as indicated was identified by Chow et al. [28]. The domains in AppBp1, Uba3, and Uba1 are drawn based on Schulman and Harper's review [38].

successive DNA synthesis without cell division [32]. Transfection of the human homologue AppBp1 into ts41 cells restores normal cell cycle at the non-permissive temperature [33]. These data establish AppBp1 as a key player in cell cycle progression across the S-M checkpoint.

The cell cycle is tightly regulated by ubiquitination through an enzymatic cascade that transfers ubiquitin to selected proteins for proteasomal degradation. The first clue to the function of AppBp1 in the cell cycle is that it is highly homologous to the N-terminus of the ubiquitin-activating enzyme Uba1 [28] (Figure 1B). However, AppBp1 lacks the C-terminal conserved cysteine residue necessary for the formation of a thioester bond with ubiquitin [28,34]. AppBp1 was soon shown to bind Uba3, which is highly homologous to the C-terminus of Uba1 and has the corresponding active site cysteine [35,36] (Figure 1B). Together, AppBp1 and Uba3 form a bipartite Nedd8 Activating Enzyme (NAE) for the ubiquitin-like protein nedd8 (see reviews [37,38]). In the enzymatic cascade that activates nedd8, AppBp1 is upstream of Uba3 since Uba3 is not able to restore ts41 cell growth when AppBp1 is inactivated by non-permissive temperature [33]. Mutation of Uba3 in *C. elegans* also profoundly affects mitosis, presumably by affecting the same nedd8-activation pathway [39].

By now it is understood that NAE activates nedd8 by an ATP-dependent mechanism analogous to the activation of ubiquitin by Uba1 [38,40,41] (Figure 2). Activation of nedd8 results in the fully loaded NAE complex containing two nedd8 molecules, covalently

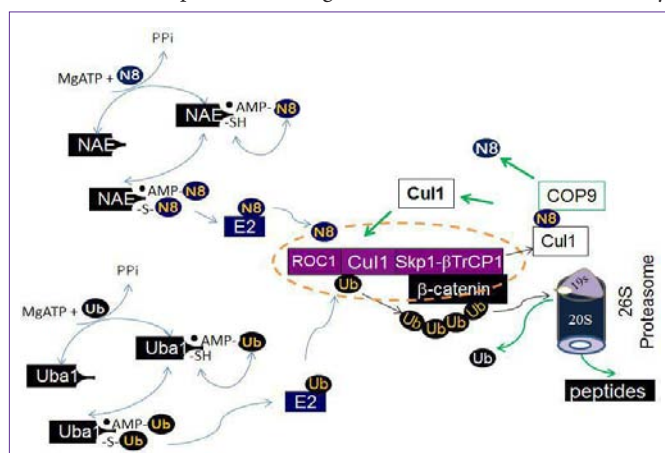


Figure 2: Schematic of neddylation, ubiquitination and proteasomal degradation. In the neddylation pathway, $Mg^{++}ATP$ and nedd8 bind NAE nucleotide binding site, react to yield nedd8-AMP, and release inorganic pyrophosphate. Nedd8-AMP then react with the thiol of the NAE active site cysteine, form the NAE-nedd8 thioester, and release AMP. A second ATP and nedd8 bind AppBp1/Uba3 to form nedd8-AMP, resulting in the fully loaded complex containing two nedd8 molecules, covalently bound nedd8 thioester and nedd8-AMP that occupies the NAE adenylation domain. This form of NAE activates the transfer of nedd8 to the conjugating enzyme by a transthioylation reaction. Subsequently, nedd8 is covalently attached to a Cullin in a lysine residue in the conserved Cullin domain. Neddylation of the Cullin activates the Cullin ubiquitin ligase. In the ubiquitination pathway, Uba1 functions similarly to NAE, but activates ubiquitin, which is ligated to target proteins when Cullin ubiquitin ligase are activated by neddylation. A Cul1 ubiquitin ligase complex (purple) known as SCF complex, is given as an example. The SCF complex consists of the zinc-finger protein, which binds to the C-terminus of Cul1, the adaptor protein Skp1, which binds to the N-terminus of Cul1. The substrate β -catenin is recruited by a specific F-box protein β TrCP1. The ubiquitin chain is recognized by the 26S proteasome, for proteasomal degradation. Cul1 is deneddylated by COP9 signalosome.

bound nedd8 thioester and nedd8-AMP that occupies the NAE adenylation domain. This form of NAE activates the transfer of nedd8 to the nedd8-conjugating enzyme by a transthioylation reaction. Eventually, nedd8 is covalently attached to a Cullin in a lysine residue in the conserved Cullin domain. Neddylation of the Cullin induces a conformational change that couples the ubiquitin-charged ubiquitin-conjugating enzyme with the substrate for ubiquitination. Many Cullin substrates are involved in cell cycle control and often elevated in cancer cells [42]. Due to its role in activating Cullin ubiquitin ligase, the neddylation pathway is being targeted for cancer growth inhibition [41].

The potential function of APP in regulating AppBp1 is first examined in studies using molecular and cellular biology techniques. APP's binding site in AppBp1 includes amino acids 145-251 (Figure 1B) [28]. This AppBp1 fragment was initially discovered by screening a brain cDNA library with GST-C100 (GST fused to APP C-terminal 100 amino acids) as the bait for APP-binding proteins [28]. Further investigations by yeast two-hybrid and co-immunoprecipitation demonstrate that this APP-interacting fragment does not bind Uba3, which does bind AppBp1's C-terminal 443-534 amino acids [33]. These data suggest that a complex consisting of APP, AppBp1 and Uba3 may assemble in cells. Subsequent structural analyses suggest that AppBp1 (145-251) is essential in the activity of NAE because this region overlaps with the active site that mediates adenylation and also the interaction site between nedd8 and AppBp1. In the structural analyses, NAE is superimposed with MoeB (an ancient form of E1 in bacteria), which shows that the adenylation domain comprises AppBp1's residues 6-168 and 486-534, and Uba3's residues 12-210 and 290-347 [43,44]. In addition, AppBp1's residues 178-280 form a portion of the catalytic cysteine domain with a charged surface that contacts nedd8's acidic face [45]. These observations strongly suggest that APP plays a role in regulating adenylation and/or nedd8 activation by NAE (Figure 2). Therefore, it is very important to understand the role of APP in the AppBp1 pathway.

AD pathogenesis suggests a gradual loss of APP's function in cell cycle control

Cell Cycle Markers (CCMs) are often ectopically expressed in AD brain neurons (see review [46]). Table 1 lists examples of the CCMs that are regulated by the AppBp1 pathway and also increased or activated in AD brains. Another CCM Ki67, not expressed in G0 cells, is also significantly increased in AD brain neurons and often co-localizes with neurofibrillary tangles [47,48]. Besides cell cycle proteins, DNA replication in post mitotic neurons is another major CCM. A higher percentage of AD hippocampal neurons enter the S-phase and undergo full or partial DNA re-replication [49,50]. Furthermore, AD neurons may proceed to nuclear division [51]. Such cell cycle events may compromise neuronal function and survival in AD brains [46]. Abnormal CCMs are not limited to neurons. Compared to the controls, AD patient-derived fibroblasts have a two-fold increase in trisomy 21, a mitotic defect due to unequal chromosome segregation [52]. In addition, AD patient-derived lymphocytes are impaired in G1/S checkpoint because they do not respond to cell cycle arrest agent [53]. These pathological changes suggest that the mechanism that prevents ectopic CCM expression is impaired or lost in AD.

Table 1: Examples of the cell cycle proteins degraded by NAE-activated Cullin ubiquitin ligases.

NAE Targets	Function in Cell Cycle	Cullin Ubiquitin Ligase involved	Protein changes in AD brains
Cyclin B	Essential in entry into mitosis [93]	Cul1/ SCF [94]	Increased [95-99]
Cyclin D	Promotes G1-S transition [100]	Cul1/SCF(Fbxw8) [101] Cul1/SCF(Fbx4)[102]	Increased [96,99]
Cyclin E	Endoreplication in placenta trophoblast giant cells [103]	Cul3 [104] Cul1/SCF (Skp2) [105-107] Cul1/SCF(Fbw7) [108]	Increased [97]
Cdc25a	Promote G2-M transition [109]	Cul1/SCF(β -TrCP) [110]	Activated [111]
p27(kip1)	Nuclear p27 inhibit G1 Phase progression [112,113] Cytoplasmic p27 negatively regulates migration [114,115]	Cul1/SCF [105,116] Cul4 [117]	Increased [118]

What is the protective mechanism that prevents neurons from de-differentiation while they actively respond to all kinds of stimuli? Due to the interaction between APP and AppBp1, the AppBp1 pathway is a good candidate molecular pathway for neuronal survival and function. In one set of studies, APP was overexpressed in primary neurons via the herpes simplex virus vector, which is a valuable model system for investigating potential early triggering events in the development of AD [54]. Overexpression of AppBp1 in primary neurons induces apoptosis through the neddylation pathway [33]. Similarly, overexpression of APP in primary neurons also induces apoptosis by increasing AppBp1 levels [29,55]. In both cases, elevation of AppBp1 levels may increase its insolubility, which forms a potential negative feedback loop that inhibits AppBp1's activity, causing neuronal death.

Another strategy to determine whether APP regulates AppBp1's activity is to assay whether APP affects the levels of the substrates downregulated by the AppBp1 pathway. For this purpose, it was first examined whether APP overexpression in primary neurons affected β -catenin, a target of Cullin-1 ubiquitin ligase [56] and it accumulates in Uba3-deficient cells [57] (Figure 2). In primary neurons, a tripling of APP expression via the viral vector leads to a reduction of β -catenin to approximately half of that in the control. Conversely, suppression of the endogenous APP by shRNAs results in a significant increase of total β -catenin compared to the control. The effect of APP on β -catenin was then determined *in vivo* [56,58,59]. APP knockout neurons have much higher levels of β -catenin in cell bodies and processes than in wild type cells. When β -catenin is stabilized, it is known to translocate to the nucleus where it activates transcription of genes such as cyclin D1. Indeed, cyclin D1 protein levels are also dramatically increased in APP knockout granule cells. These data suggest that APP is an essential component of AppBp1 pathway.

Unlike in primary neuronal cultures where protein expression can be significantly increased, which may mimic an early event in the development of AD, APP levels do not necessarily increase in postmortem brains. Some reports show no overall elevation of APP protein or mRNA in AD brain homogenates [60,61] and others have even reported a decrease [62,63]. One recent study shows that in cognitively intact adult brains, APP accumulates with age, but in AD brains, the protein is not elevated in neurons adjacent to mature plaques [64]. This study also shows that neurons adjacent to mature plaques have dramatically lower levels of APP than those remote from such plaques [64]. APP deficiency can cause age-related cognitive deficits and impaired long-term potentiation in mouse models *in vivo* [65]. Furthermore, a moderate elevation of APP is neuroprotective *in vivo* [66]. Similarly, a physiological level of AppBp1 is likely beneficial since it is expressed in some hippocampal neurons *in vivo* [28,37]. In

addition, AppBp1 inhibits amyloid genesis since its suppression by shRNAs results in a dramatic increase of A β 42 [30].

If the activation of the AppBp1 pathway is a physiological function of APP, the lack of APP in neurons near the plaques in late stage AD may result in a functional deficit of AppBp1. Indeed, pathological changes such as elevation of CCMs observed in postmortem AD brains suggest that the AppBp1 pathway does not function properly (Table 1). AppBp1 activates neddylation of Cullins and neddylated Cullins normally reside in the nucleus [67]. However, in AD hippocampal neurons, almost all nedd8 is present in the cytoplasm, different from the nuclear localization of nedd8 in control neurons [29]. The cytoplasmic localization of nedd8 in AD neurons also suggests that AppBp1 fails to function, which in turn prevents neddylation and activation of Cullin ubiquitin ligases, leading to the accumulation of CCMs in post mitotic neurons. Another pathological change is that more AppBp1 became Triton-insoluble in AD brains [29], which does not occur during normal aging [30]. Similar to the inactivation of AppBp1 in ts41 cells by the non-permissive temperature, cortical precursors from APP-deficient E15 cerebral cortex have defect in crossing G2-M phases [68], further suggesting that APP plays a role in cell cycle transit controlled by AppBp1. Together, these data suggest that the development of AD involves a gradual loss of APP's function in the AppBp1 pathway, which may be caused by negative feedbacks that warrants further investigation.

APP in DNA replication and genome integrity

Besides AppBp1, APP also interacts with UV-damaged DNA-binding protein 1 (DDB1) [69], which suggests a role of APP in DNA replication control. DDB1 is the common adaptor subunit in Cul4A and Cul4B (Cul4) ubiquitin ligases [70-73]. DDB1 connects Cul4 with the variable substrate receptor subunit called DDB1-Cul4-Associated Factor (DCAF) [74]. Cul4 ubiquitin ligases are responsible for DNA replication-dependent destruction of Cdt1 in the S-phase of the cell cycle [75-77] and in DNA damage-induced repair synthesis [78,79]. Degradation of Cdt1 in the S-phase prevents DNA rereplication [80] whereas Cdt1 overexpression causes extensive numerical and structural chromosomal aberrations [81]. It has been found that ubiquitination of Cdt1 during the S-phase is coupled to the process of DNA replication [80]. Upon DNA damage, ubiquitination of Cdt1 also localizes to chromatin together with Cul4 [77,82,83]. Together, the evidence suggests that in post mitotic neurons, APP protects genome integrity through DDB1-Cul4 ligase-mediated Cdt1 degradation, but this potential function is gradually lost during disease progression. In fact, the accumulation of unrepaired DNA damage has long been postulated as the cause of neurodegeneration [84-86].

As outlined above, DNA synthesis is coupled to the quality control of genome integrity involving the DNA replication licensing factor Cdt1. Human cells treated with the small molecule inhibitor of NAE, MLN4924, undergo continuous DNA synthesis and have a significant rise in Cdt1 levels among other CCMs [87]. In neurons where overexpression of wild type APP causes DNA synthesis although it has less effect than familial AD APP mutants [29,55], APP overexpression also downregulates Cdt1 levels (Chen's unpublished observation). These data suggest that a major function of APP in neurons is downregulating Cdt1 when it assembles with AppBp1 and DDB1. Increases in DNA replication in post mitotic neurons from AD brains [46,88] probably represent early events since overexpressing APP or AppBp1 in primary neurons induce DNA syntheses before apoptosis [29]. Together, the data suggest that APP plays a key role in downregulating Cdt1 among others through the AppBp1-activated neddylation pathway, which is likely crucial for DNA damage-induced repair synthesis, which is crucial for the genome integrity of long-lived neurons.

Conclusion

In this review, we analyze the function of APP and hypothesize that APP plays a key role in AppBp1-activated neddylation pathway. In this role, APP protects genome integrity in DNA-damage response and promotes cell cycle progression through the S-M checkpoint. Based on this new hypothesis, the decline of APP's function is the driving force for the development of AD, resulting in the accumulation of CCMs and neurodegeneration. Repair synthesis in mature neurons is essential for neuronal function if DNA damage can be caused by physiological brain activities [89]. According to the new hypothesis, neuronal vulnerability lies in the inability of efficient repair synthesis due to the dysfunction of APP. APP dysfunction may also impair stem cell differentiation which depends on prior DNA synthesis [90,91]. Although it has not been tested, the function of APP in the AppBp1 pathway may involve APP cleavage into monomeric A β peptides since these peptides promote DNA synthesis-linked neural stem cell differentiation [90]. The function of APP may also involve neddylation of APP in the C-terminus [92]. In order to gain a greater understanding of APP's function as proposed, many questions remain to be answered. What factors cause the gradual decrease of APP's activity during the development of AD? How does APP activate AppBp1? Further confirmation of the molecular pathway from APP may provide novel drug targets and new therapeutic strategies for AD treatment and prevention.

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