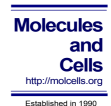
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N-Terminal Acetylation-Targeted N-End Rule Proteolytic System: The Ac/N-End Rule Pathway

Kang-Eun Lee^{1,2}, Ji-Eun Heo^{1,2}, Jeong-Mok Kim¹, and Cheol-Sang Hwang^{1,*}

Although No-terminal acetylation (Nt-acetylation) is a pervasive protein modification in eukaryotes, its general functions in a majority of proteins are poorly understood. In 2010, it was discovered that Nt-acetylation creates a specific protein degradation signal that is targeted by a new class of the N-end rule proteolytic system, called the Ac/N-end rule pathway. Here, we review recent advances in our understanding of the mechanism and biological functions of the Ac/N-end rule pathway, and its crosstalk with the Arg/N-end rule pathway (the classical N-end rule pathway).

INTRODUCTION

Conditional and constitutive proteolysis are essential life processes. Living cells exploit metabolic turnover to adjust intracellular protein levels, restrict unwanted or aberrant protein accumulation, and supply protein and peptide fragments or free amino acids for physiological needs (Varshavsky, 2011; Zattas and Hochstrasser, 2015). Malfunctions in regulated proteolysis lead to the accumulation of misfolded or abnormal proteins, causing many fatal diseases, including cancers, immunological or neurobiological disorders, as well as aging (Ciechanover and Kwon, 2015; Varshavsky, 2011). Detailed knowledge of intracellular proteolytic machineries and their modes of action will therefore provide new insight into drug interventions for the treatment of devastating maladies or general health maintenance.

The ubiquitin-proteasome system (UPS) is involved in the majority of regulated proteolysis within a cell. The first well-defined protein degradation signals (degrons) in the UPS were N-degrons, which comprise destabilizing Nt-residues. Their detailed examination has revealed the N-end rule, which enables an Nt-residue or its modification to regulate the in vivo half-life of a protein (Varshavsky, 2011). The original (classical) N-end rule pathway (now called the Arg/N-end rule pathway) targets unmodified destabilizing Nt-residues for degradation

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(Bachmair et al., 1986). In contrast, a new branch of the N-end rule pathway (called the Ac/N-end rule pathway) recognizes the Nt-acetyl group of cellular proteins for proteolysis (Hwang et al., 2010b). Recent studies on the Ac/N-end rule pathway have revealed a variety of cellular functions, including protein quality and subunit stoichiometry control (Hwang et al., 2010b; Kim and Hwang, 2014; Kim et al., 2014; Shemorry et al., 2013), blood pressure regulation via G-protein signaling (Park et al., 2015), and immune responses to pathogens in plants (Xu et al., 2015). Here, we review current knowledge related to the mechanism and biological functions of the Ac/N-end rule pathway.

NT-ACETYLATION, MACHINERY, AND FUNCTIONS

Nascent cellular proteins harbor Nt-methionine (Nt-Met), which is co-translationally removed by ribosome-bound Metaminopeptidases (MetAPs) if small residues (Ala, Gly, Ser, Cys, Thr, Pro, or Val) are positioned at the penultimate site. Nt-Met removal is evolutionarily conserved and occurs in approximately two-thirds of cellular proteins (Giglione et al., 2015). In eukaryotes, both retained Nt-Met and newly exposed Nt-residues (immediately after Nt-Met removal or their proteolytic cleavage) are very frequently Nα-terminally acetylated (Nt-acetylated); Nt-acetylation occurs in 80-90% and 50-70% of cytosolic human and yeast proteins, respectively (Van Damme et al., 2012). Unlike the reversible Nεacetylation/deacetylation, Nt-acetylation appears to be irreversible because no Nt-deacetylases have been identified to date (Starheim et al., 2012). Additionally, Nt-acetylation takes place both cotranslationally and posttranslationally in vivo (Gautschi et al., 2003).

Nt-acetylases catalyze the covalent attachment of an acetyl moiety from acetyl-CoA to the free $\alpha\textsc{-amino}$ group of a protein, which alters the steric or chemical properties of the N-terminus of a protein by neutralizing its positive charge and making the Nt-residue larger and more hydrophobic (Aksnes et al., 2015b). Nt-acetylases belong to the GCN5-related N-acetyltransferase (GCNT) family (Polevoda et al., 1999) and are classified as NatA, NatB, NatC, NatD, NatE, and NatF based on substrate specificity and subunit compositions. The substrate specificity of each Nt-acetylase is mostly based on the first two protein residues. Of the 6 Nt-acetylases (NatA to F), NatA, NatB, and NatC primarily act on protein N-termini (Aksnes et al., 2015b) (Fig. 1).

NatA consists of a catalytic subunit Naa10 (Ard1) and an auxiliary subunit Naa15 (Nat1) (Mullen et al., 1989; Park and Szostak, 1992), and acetylates Ser, Ala, Cys, Gly, Thr, and Val

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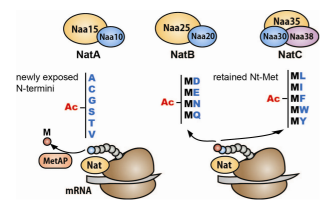


Fig. 1. Substrate specificity and subunit compositions of Nt-acetylases. Among 6 Nt-acetylases (NatA-F), NatA, NatB, and NatC mainly Nt-acetylate cellular proteins. NatA consists of a catalytic subunit, Naa10, and an auxiliary subunit, Naa15, and acetylates Ser (S), Ala (A), Cys (C), Gly (G), Thr (T), or Val (V) N-termini of its substrates after Nt-Met removal by methionine aminopeptidases (MetAPs). NatB consists of a catalytic subunit, Naa20, and an auxiliary subunit, Naa25, and Nt-acetylates Met-Asn (MN), Met-Asp (MD), Met-Gln (MQ), or Met-Glu (ME) on cellular proteins. NatC contains a catalytic subunit, Naa30, and auxiliary subunits, Naa35 and Naa38, and Nt-acetylates Met-Ile (MI), Met-Phe (MF), Met-Trp (MW), Met-Leu (ML), or Met-Try (MY) on its substrates.

N-termini (Aksnes et al., 2015b) (Fig. 1). NatA functions in cell proliferation, gene silencing, stress resistance, mating process, ribosome biogenesis, apoptosis, protein folding and disaggregation, photosynthesis, development, and stress responses (Aksnes et al., 2015b; Dorfel and Lyon, 2015). Loss of NatA causes a wide range of phenotypes depending on organisms, from slight growth defects to lethality (Aksnes et al., 2015b; Dorfel and Lyon, 2015). In humans, dysregulation of NatA results in various cancer types and neuronal diseases (Kalvik and Arnesen, 2013). In particular, missense or splicing mutations of Naa10, a catalytic subunit of NatA, cause X-linked human genetic disorders, including Ogden Syndrome (Rope et al., 2011) and Lenz microphthalmia syndrome (Esmailpour et al., 2014).

NatB contains a catalytic subunit, Naa20, and an auxiliary subunit, Naa25, and specifically targets N-terminal Met with Asn, Asp, Gln, or Glu at the second position (Fig. 1). NatB plays roles in cell growth in response to various stressors, mitochondrial inheritance, actin cable formation, cell wall maintenance, flowering regulation, plant development, etc. (Aksnes et al., 2015b; Ferrandez-Ayela et al., 2013; Lee et al., 2014).

NatC has a catalytic subunit, Naa30, and two accessory subunits, Naa35/Naa38, and primarily acetylates Nt-Met with bulky hydrophobic residues at position 2 (Fig. 1). It affects cell growth, protein targeting, viral particle maintenance, stress responses, chloroplast development, etc. (Aksnes et al., 2015b; Pesaresi et al., 2003).

NatD, NatE, and NatF include only catalytic subunits Naa40, Naa50, and Naa60, respectively. NatD acetylates solely the Nterminus of histone H2A or H4 and partially regulates histone modification and ribosomal DNA silencing (Polevoda et al., 2009). NatE and NatF display slightly overlapping substrate specificities with NatC. NatE is involved in chromosome segregation and microtubule growth. Interestingly, membrane-localized NatF is only found in multicellular organisms and mainly acetylates cytosolic-faced N-termini of transmembrane proteins, thus maintaining Golgi structural integrity (Aksnes et al., 2015b; 2015c; Van Damme et al., 2011).

The function of Nt-acetylation on a vast number of cellular proteins has remained a conundrum since its discovery (Narita, 1958). However, studies on individual proteins have revealed that Nt-acetylation contributes to protein stability, degradation, interactions, targeting, activity, etc. (Behnia et al., 2004; Forte et al., 2011; Hwang et al., 2010b; Jornvall, 1975; Scott et al., 2011; Setty et al., 2004). Schulman and colleagues have shown that Nt-acetylation of the E2 enzyme Ubc12 increases its binding affinity to the E3 ligase Dcn1 by approximately ~100-fold, thereby promoting E3 ligase activity (Scott et al., 2011). In addition, Nt-acetylation affects global protein folding, chaperone

expression, prion stability, and disaggregation of neurodegenerative proteins, such as β -amyloid, huntingtin, or α -synuclein (Arnesen et al., 2010; Holmes et al., 2014; Pezza et al., 2009). Nt-acetylation also precludes the targeting of cytosolic proteins to the endoplasmic reticulum (ER) (Forte et al., 2011) and transit polypeptide import into plastids (Bischof et al., 2011).

Nt-acetylation has been largely regarded as a metabolic stabilizer that protects proteins or peptides from destruction (Jornvall, 1975). For example, Nt-acetylated globins and lysozymes are more stable than their unacetylated counterparts (Hershko et al., 1984). Nt-acetylation also blocks the Nt-ubiquitylation-mediated degradation of several cellular proteins, in which a linearly attached Nt-Ub moiety serves as a specific degron (Ciechanover and Ben-Saadon, 2004). Furthermore, Nt-acetylation stabilizes Arg/N-end rule substrates by directly inhibiting N-degron recognition by the Arg/N-end rule pathway (Varshavsky, 2011) (see below). However, Nt-acetylation also creates a specific degron that is targeted by the Ac/N-end rule pathway (Hwang et al., 2010a; Varshavsky, 2011).

N-DEGRONS, THE N-END RULE, AND THE ARG/N-END RULE PATHWAY

Ubiquitin (Ub) is a small 76-amino-acid protein that is highly conserved across all eukaryotes. Ubiquitylation links the carboxyl terminus of Gly in Ub to the Nε-group of Lys or, relatively rarely, to the Nα-amino group of Met in cellular proteins. The covalent linkage of Ub to target substrates requires consecutive reactions of Ub-activating enzymes (E1s), Ub-conjugating enzymes (E2s), and Ub ligases (E3s). The ubiquitylation reaction repeats several times to generate a polyubiquitylated substrate that is selectively eliminated by the 26S proteasome. In the UPS, E3 ligase directly contacts target substrates for polyubiquitylation and subsequent proteasomal degradation (Varshavsky, 2011; Zattas and Hochstrasser, 2015). One fundamental question related to the UPS is exactly what structural or sequence features within a target protein trigger its Ubmediated destruction by the 26S proteasome; are there any specific degrons (Kim and Hwang, 2014; Ravid and Hochstrasser, 2008)?

Varshavsky and colleagues first defined the primary degrons in the UPS (called N-degrons) based on the unexpected observation that the half-lives of engineered β -galactosidases with varied Nt-residues are determined by an exposed Nt-residue in Saccharomyces cerevisiae (Bachmair et al., 1986). According to the stability of the resulting β -galactosidases, they classified Nt-amino acids as either stabilizing or destabilizing residues (Bachmair et al., 1986). N-degrons include primary destabilizing

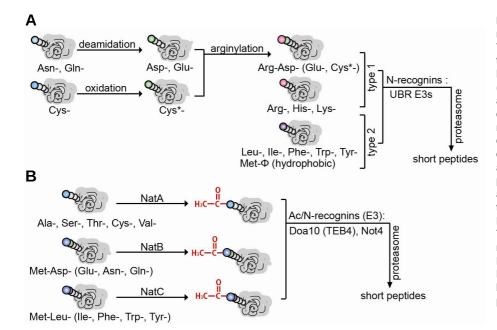


Fig. 2. Two branches of the N-end rule pathways in eukaryotes. (A) The Arg/N-end rule pathway, which targets unmodified Arg, His, Lys, Leu, Ile, Phe, Trp, Tyr, and Met-Φ (hydrophobic) Nt-residues. Nt-Gln and Asn are destabilizing after Ntdeamidation and subsequent arginylation. Nt-Cys also becomes destabilizing through preliminary oxidation and subsequent Ntarginylation. (B) The Ac/N-end rule pathway, which targets acetylated residues of cellular proteins for degradation. Doa10 and Not4 are yeast Ac/N-recognins and Teb4 is a mammalian Ac/Nrecognin. In addition to the NatA, NatB, and NatC substrates, other Nt-acetylated proteins are potentially targeted by the Ac/N-end rule pathway for degradation.

Nt-residues, internal Lys residue(s) for ubiquitylation, and flexible region(s) for the exposure of substrate Nt-residues. Extensive examination of N-degrons has revealed the N-end rule and the related proteolytic system, called the N-end rule pathway (Tasaki et al., 2012; Varshavsky, 2011).

The N-end rule pathway is generally grouped into the Arg/Nend rule pathway and the Ac/N-end rule pathway in eukaryotes (Fig. 2). The Arg/N-end rule pathway targets specific unmodified Nt-residues for polyubiquitin-mediated proteolysis by the 26S proteasome (Varshavsky, 2011) or, to a lesser extent, by autophagy (Cha-Molstad et al., 2015) (Fig. 2A). In eukaryotes, the Arg/N-end rule pathway employs specific UBR-type E3 ligases as N-recognins, which are recognition components of the N-end rule pathway. The UBR-type E3s bind directly to unmodified basic (Arg, Lys, His) and large hydrophobic (Leu, Phe, Tyr, Trp, Ile) destabilizing Nt-residues. Nt-Asn and Gln can act as destabilizing residues through their deamination via Ntamidases, resulting in Asp or Glu, and subsequent Ntarginylation via Arg-tRNA-protein transferases (ATEs) (Kwon et al., 1999; Varshavsky, 2011). Nt-Cys also becomes destabilizing through its oxidation by NO, oxygen, or cysteine oxidases, and entails Nt-arginylation by ATEs. Subsequently, Ntarginylated proteins are directly recognized by UBR-type Nrecognins for polyubiquitin-mediated degradation by the 26S proteasome (Gibbs, 2015; Tasaki et al., 2012; Varshavsky, 2011). In addition to primary destabilizing Nt-residues, the Arg/N-end rule pathway directly recognizes, for proteolysis, Nt-Met of cellular proteins with a hydrophobic residue at the 2nd position, termed MΦ-degrons (Kim et al., 2014) (Fig. 2A).

The functions of the Arg/N-end rule pathway include sensing small molecules (e.g., heme, di/tripeptides, and oxygen), eliminating abnormal proteins, regulating genome stability, apoptosis, DNA repair, G-protein signaling, autophagy, fungal pathogenesis, plant hormone responses, leaf senescence, cardiac signaling, and the viral life cycle (Cha-Molstad et al., 2015; Dougan et al., 2012; Gibbs et al., 2014; Hwang et al., 2010a; Sriram et al., 2011; Tasaki et al., 2012; Varshavsky, 2011). The Arg/N-end rule pathway also mediates the degradation of breast cancerrelated tumor suppressor 1 (BRCA1) (Xu et al., 2012) and the

Parkinson's disease-associated protein PTEN-induced putative kinase 1 (PINK1) (Yamano and Youle, 2013). Interestingly, loss of human UBR1, an E3 ligase of the Arg/N-end rule pathway, causes Johanson-Blizzard Syndrome, which is characterized by physical malformations and pancreatic exocrine insufficiency (Hwang et al., 2011; Zenker et al., 2005).

THE NT-ACETYLATION-TARGETED N-END RULE PATHWAY (THE AC/N-END RULE PATHWAY)

The Arg/N-end rule pathway classifies Met, Ser, Thr, Val, Cys, Ala, and Pro as stabilizing Nt-residues (Varshavsky, 2011). However, Nt-Cys is readily oxidized in mammalian and plant cells, and subsequently arginylated by ATEs following its targeting to UBR-type N-recognins (Gibbs et al., 2011; Hu et al., 2005; Lee et al., 2005; Weits et al., 2014). Interestingly, reporters with Nt-Cys are apparently long-lived in S. cerevisiae, which does not retain typical NO synthases (Hu et al., 2005), but produces significant amounts of NO or reactive oxygen species (Bhattacharjee et al., 2009). Thus, reporters with Nt-Cys were assumed to be destabilized only if the Nt-Cys could be oxidized, which depends on downstream sequence contexts. Indeed, Nt-Cys-reporters bearing non-basic residues at position 2 become destabilized. Additionally, otherwise identical reporters with Met, Ser, Thr, Ala, or Val N-termini are short-lived, in contrast to longlived reporters with Nt-Gly or Pro (Hwang et al., 2010b) (Fig. 2B). The first two Nt-residue-dependence of short-lived reporters implicates likely correlation with the substrate specificity of S. cerevisiae Nt-acetylases that act on Met, Ala, Val, Ser, Thr, or Cys N-termini, but very rarely or never on Nt-Gly or Pro, and Ntresidues with basic residues at the penultimate site (Aksnes et al., 2015b). More interestingly, short-lived reporters with nonbasic residues at position 2 do not involve Ubr1, the sole Nrecognin of the Arg/N-end rule pathway in S. cerevisiae. Instead, the reporters are dramatically stabilized in the absence of an ER-transmembrane Doa10 E3 ligase, which mediates the degradation of both cytosolic and nuclear or ER proteins in concert with Ubc6/Ubc7 E2s (Hwang et al., 2010b; Swanson et al., 2001) (Fig. 2B). Strikingly, Doa10 preferentially interacts with

acetylated Nt-Met, Ala, Ser, Cys, Thr, and Val, rather than their non-acetylated counterparts. Furthermore, Doa10 is required for the degradation of not only artificial model substrates, but also some Nt-acetylated native proteins, including MATα2 (a mating type regulator), Tbf1 (a telomerase regulator), Slk19 (a chromosome segregation regulator), His3 (a histidine biosynthetic enzyme), Hsp104 (a chaperone), Ubp6 (a deubiquitylating enzyme), Aro8 (an aromatic aminotransferase), and Ymr090w (an unknown protein) (Hwang et al., 2010b).

Khemlinskii and Knop showed that the degradation of a CL-mCherry-sfGFP fluorescent protein timer with Nt-Cys-Leu, a plausible Ac/N-degron, involves Doa10 E3 ligase and its cognate E2s Ubc6/Ubc7 (Khmelinskii and Knop, 2014). Fields and colleagues also demonstrated that chimeric fusions between Deg1 (1-67 residues of Mato2) and Leu2 are longer-lived when Asn at position 2 is replaced by a basic Lys or Arg (Kim et al., 2013), which suppresses Nt-acetylation in yeast (Arnesen et al., 2009). Additionally, Deg1-Leu2 becomes stabilized in an naa20Δ mutant that lacks a catalytic subunit of cognate NatB Nt-acetylase (Kim et al., 2013), similar to another Deg1-Ura3 reporter (Hwang et al., 2010a). However, Nt-acetylation slightly perturbs the proteolysis of different versions of Deg1-fusion and endogenous MATo2 (Zattas et al., 2013).

COMPLEMENTARITY BETWEEN THE ARG/N-END RULE AND THE AC/N-END RULE PATHWAYS

Features of Nt-acetylation, e.g., its prevalence, cotranslationality, and apparent irreversibility, indicate that most proteins should retain acetylated N-degrons (Ac/N-degrons) from their emergence from ribosomes to their destruction (Mogk and Bukau, 2010; Varshavsky, 2011). Nonetheless, many cellular proteins bear both acetylated and unacetylated Nt-residues owing to partial Nt-acetylation. Moreover, some cellular proteins are rarely or never Nt-acetylated (Aksnes et al., 2015b). The built-in property of Ac/N-degrons suggests that unacetylated Nt-residues also act as imprinted intrinsic degrons. Hence, we presumed that unacetylated stabilizing Nt-residues of the Arg/N-end rule pathway might be destabilizing depending upon their downstream sequence contexts, especially the 2^r residues. To test this possibility, we focused on Nt-Met because it is present almost every nascent polypeptide. We found that Nt-Met acts as specific protein degrons if it is followed by hydrophobic residues (Φ) (Leu, Phe, Ile, Trp, Tyr, Gly, or Ala), and these are termed Met-Φ degrons (Kim et al., 2014). Interestingly, yeast Ubr1 and mouse UBR1 and UBR2 E3 ligases of the Arg/N-end rule pathway bind specifically to Met-Φ/N-degrons. We also showed that the Arg/N-end rule pathway eliminates, through Met-Φ/N-degrons, quasi-randomly selected natural Met-Φ protein Msn4 (a transcription activator), Sry1 (a 3hydroxyaspartate dehydratase), Arl3 (a Golgi-bound GTPase), and Pre5 (a proteasome subunit) as well as misfolded Met-Φ proteins (Kim et al., 2014). The finding of Met-Φ/N-degrons enormously increases the number of Arg/N-end rule substrates because more than 15% of DNA-encoded proteins have Nt-Met-Φ sequences in both yeast and humans. Furthermore, detailed analyses of Met-Φ degrons have unraveled the complementary crosstalk between the Ac/N-end rule and the Arg/Nend rule pathways. For example, when Met-Φ proteins protrude from ribosomes, the Arg/N-end rule pathway immediately attacks them for degradation. Otherwise, if Nt-acetylated, the Ac/N-end rule pathway is activated and eliminates Met-Φ proteins by recognizing their Nt-acetyl moiety. Consequently, the complementary collaboration between the Arg/N-end rule and

Ac/N-end rule pathways makes it possible to efficiently eliminate Met-Φ proteins irrespective of their Nt-acetylation states for physiological needs (Kim and Hwang, 2014; Kim et al., 2014). Furthermore, Nt-acetylation not only precludes the targeting of Met-Φ proteins by the Arg/N-end rule pathway, but also converts Met-Φ/N-degrons into AcMet-Φ/N-degrons, making them susceptible to the Ac/N-end rule pathway (Kim and Hwang, 2014; Kim et al., 2014).

A combined analysis of bioinformatics and proteomic data has revealed that substantial fractions of proteins (~10%) are potentially destroyed by retained Nt-Met (Meinnel et al., 2005). In particular, the transient retention of Nt-Met destabilizes chloroplast D2 variants (Giglione et al., 2003), β -glucuronidase in plants (Sawant et al., 2001), and a GST variant in S. cerevisiae (Chen et al., 2002). Furthermore, Ubr1 may mediate the degradation of a previously identified Met-Φ-type extension of Ura3 (with Nt-MLDDKCRVTP) via its Nt-Met-Leu sequence (Ghislain et al., 1996). In contrast, treatment with the MetAP2 inhibitor TNP-470 dramatically stabilizes a Rab37 GTPase (with a Met-Thr Nt sequence) in murine pulmonary endothelial cells, suggesting that the retained Nt-Met of Rab37 also prevents its NatA-mediated Nt-acetylation and subsequent degradation by the Ac/N-end rule pathway (Sundberg et al., 2011). Together, the identification of Nt-Met as an N-degron indicates that virtually all 20 amino acids may serve as potential destabilizing Nt-residues according to their downstream sequence contexts and Nt-acetylation status. For example, fructose-1.6-bisphosphatase, a key regulatory enzyme of gluconeogenesis, is targeted by the GID (glucose-induced degradation-deficient) E3 ligase complex in S. cerevisiae. Interestingly, substitutions of the Nt-Pro of fructose-1.6bisphosphatase with other residues abrogate its degradation, suggesting the GID E3 ligase recognizes the Nt-Pro of the metabolic enzyme (Hammerle et al., 1998; Menssen et al., 2012). Additionally, the Drosophila apoptosis inhibitor DIAP1 and DIAP2 E3 ligases directly contact Nt-Ala of processed caspases for proteolysis as plausible specific Ala/N-degron and Ala/N-recognins (Ditzel et al., 2003). Furthermore, the replacement of the penultimate site (Pro-2) in thymidylate synthase with certain amino acids profoundly influences its half-life through Ub-independent proteasomal degradation (Pena et al., 2009), similar to bacteria that contain the Ublacking prokaryotic N-end rule pathway (Dougan et al., 2012). It is unclear which components directly recognize the Nt-Pro of thymidylate synthase for Ub-independent proteolvsis.

CONTROL OF SUBUNIT STOICHIOMETRY IN COMPLEXES BY THE AC/N-END RULE PATHWAY

Although most proteins obtain Ac/N-degrons during or immediately following their synthesis, the lack of individual Nt-acetylases does not significantly affect global protein expression levels, suggesting little correlation between Nt-acetylation and protein stability (Garrels et al., 1997; Helbig et al., 2010). Ac/N-degrons were, therefore, proposed to be conditionally active depending on the accessibility of Ac/N-recognins within a polypeptide or a multi-subunit complex (Fig. 3A). For example, if a protein is properly folded or assembled into a cognate complex, the Ac/N-degron may be protected from the attack by the Ac/N-end rule pathway. Otherwise, the Ac/N-degrons will remain exposed, leading to their elimination by the Ac/N-end rule pathway (Hwang et al., 2010a; Mogk and Bukau, 2010). The conditionality of built-in Ac/N-degrons plays a pivotal role in the control of protein quality and the adjustment

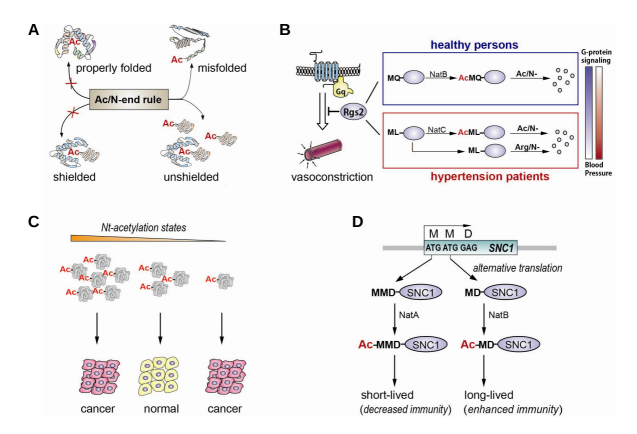


Fig. 3. Functions of the Ac/N-end rule pathway. (A) Control of protein quality and subunit stoichiometry. The Ac/N-end rule pathway targets misfolded proteins or unassembled subunit(s) of complex proteins with Ac/N-degrons. (B) Regulation of blood pressure via G-protein signaling. The Ac/N-end rule pathway degrades Nt-acetylated wild-type MQ-Rgs2. In contrast, the hypertension-related ML-Rgs2 with a Met-Φ degron is targeted by either the Ac/N-end rule pathway or the Arg/N-end rule pathway according to its Nt-acetylation status. Dual targeting of ML-Rgs2 by two branches of the N-end rule pathways dramatically decreases Rgs2 levels, and thereby increases blood pressure via augmented G-protein signaling and subsequent vasoconstriction. (C) Dysregulation of Nt-acetylation most likely affects protein stability, thus causing many cancers. (D) Control of pathogen immunity in plants. Plants contain two SNC1 (suppressors of nod-like protein receptor 1) variants with Met-Met-Asp (MMD) or Met-Asp (MD) N-termini owing to alternative translation. NatA Nt-acetylates MMD-SNC1 for degradation, whereas NatB Nt-acetylates MD-SNC1 for stabilization. Consequently, the steady-state levels of SNC1 are directly involved in plant immunity.

of subunit stoichiometries of protein complexes (Fig. 3A). For example. Nt-acetylated Cog1, a subunit of the conserved oligomeric Golgi (COG) complex is targeted for degradation by another Ac/N-recognin, Not4 E3 ligase, rather than Doa10. Interestingly, the short-lived Cog1 becomes longer-lived upon co-overexpression of the binding partner proteins Cog2, Cog3, and Cog4 by shielding its Ac/N-degron within the COG complex (Shemorry et al., 2013). The crystal structure of the Hcn1 and Cut9 subunits of Schizosaccharomyces pombe APC/C E3 ligase suggests that Hcn1 escapes from the Ac/N-end rule pathway by placing its acetylated Nt-Met in the cavity of Cut9 (Zhang et al., 2010). Indeed, unmasked Hcn1 is degraded via its Ac/N-degron when it is heterologously expressed in S. cerevisiae. Similar to the stoichiometry-mediated degradation control of the COG complex, co-expression of Cut9 represses the degradation of Hcn1 by shielding its Ac/N-degron (Shemorry et al., 2013). Additionally, Nt-acetylated Ser of the H4 peptide or Nt-acetylated Met of Dcn1 is specifically enclosed within the cavity of the double PHD1/2 finger DPF3b transcriptional protein or Ubc12 E3 enzyme, respectively (Scott et al., 2011; Zeng et al., 2010). It remains to be determined whether DPF3b and Ubc12 shield the Ac/N-degrons of H4 and Dcn1. The conditionality of the Ac/N-degron is further demonstrated in human wildtype Rgs2, a regulator of G-protein signaling. Rgs2 bearing an Ac/N-degron is strongly stabilized by the co-overexpression of one of its binding partners, the Gq protein (Park et al., 2015).

The conditional nature of Ac/N-degrons provides a new paradigm for how protein levels are sensed and balanced with respect to their interacting proteins. The conditionality of Ac/Ndegrons also holds true for other degrons, thus providing new insight into the regulation of protein quality and stoichiometric levels of individual proteins. For example, the steady-state levels of the decapping enzyme Dcp2 are modulated by competition between its degradation and assembly into decapping complexes, despite the apparent internal degrons of Dcp2 (Erickson et al., 2015). As another example, S. cerevisiae fatty acid synthase (FAS) complex subunits become short-lived in the absence of respective ligands. Interestingly, unassembled Fas2 is eliminated by the Ubr1-mediated degradation pathway, possibly via its internal degron(s), similar to subunit stoichiometric control by the Ac/N-end rule pathway (Scazzari et al., 2015). In addition, frequent aneuploidy in cancer cells and trisomy 21 in Down syndrome may perturb input subunit stoichiometries of protein complexes due to altered specific gene

dosages and a subsequent increase in unassembled or misfolded proteins (Hwang et al., 2010b). Accordingly, intracellular proteolytic systems are likely to regulate the subunit stoichiometry of complexes by targeting aberrant or unassembled proteins for the maintenance of protein quality and homeostasis.

THE AC/N-END RULE PATHWAY IN MAMMALS

The Arg/N-end rule pathway is conserved across eukaryotic species, from fungi to mammals and plants, and has recently been identified in S. cerevisiae (Hwang et al., 2010b). We demonstrated the existence of the Ac/N-end rule pathway in mammals and identified wild-type MQ-Rgs2 (with Nt-Met-Gln), a regulator of G-protein signaling, as the first physiological substrate of the mammalian Ac/N-end rule pathway (Park et al., 2015) (Fig. 3B). Additionally, two hypertension-associated 2ndposition missense human mutants, ML-Rgs2 (with Nt-Met-Leu) and MR-Rgs2 (with Nt-Met-Arg), are physiological substrates of the human Ac/N-end rule pathway (Bodenstein et al., 2007: Park et al., 2015). Both ML-Rgs2 and MR-Rgs2 are significantly shorter-lived than MQ-Rgs2, partially explaining the association between these mutants and hypertension in humans; reduced Rgs2 levels upregulate vasoconstrictors and have a weaker inactivation effect on Gq proteins, leading to high blood pressure (Aksnes et al., 2015a; Bodenstein et al., 2007; Park et al., 2015) (Fig. 3B). The ER membrane-embedded human Teb4 (March6) E3 ligase, an ortholog of yeast Doa10, (Hassink et al., 2005) was identified as the first mammalian Ac/N-recognin. Teb4 targets wild-type MQ-Rgs2 and hypertensive ML-Rgs2 and MR-Rgs2 by directly recognizing their N-terminal acetyl groups (Park et al., 2015). While human MQ-Rgs2 and its mutant MR-Ras2 are targeted for degradation solely by the Ac/Nend rule pathway, the other hypertension-associated (and the shortest-lived) ML-Rgs2 mutant is targeted by both the Ac/Nend rule pathway (as Nt-acetylated ML-Rgs2) and the Arg/Nend rule pathway (as Nt-unacetylated ML-Rgs2), consistent with the dual targeting mechanism of Met-Φ-proteins by two branches of the N-end rule pathways in yeast (see below). Consequently, the dual degradation of the hypertensive ML-Rgs2 strongly decreases ML-Rgs2 levels, leading to high blood pressure by promoting Gq-protein signaling (Aksnes et al., 2015a; Park et al., 2015) (Fig. 3B).

Out of ~30 RGS family proteins in humans, Nt-Cys of Rgs4, Rgs5, and Rgs16 are subject to oxidation, Nt-arginylation, and subsequent degradation by the Arg/N-end rule pathway (Gibbs, 2015; Hu et al., 2005; Lee et al., 2005; 2015). The identification of Rgs2 as an Ac/N-end rule substrate suggests that other RGS proteins, including Rgs1, Rgs6, and Rgs7, may also be substrates of either the Ac/N-end rule or Arg/N-end rule pathways depending on their Nt-sequence contexts and Nt-acetylation states (Park et al., 2015).

Similar to Rgs2 variants in hypertension patients, 2nd-position mutations are observed in many proteins, particularly in cancer cell exomes (Kandoth et al., 2013). Some cancer-related proteins may become longer- or shorter-lived as a result of 2ndposition mutations, which cause their targeting to either the Arg/N-end rule pathway or the Ac/N-end rule pathway (Table 1) (Fig. 3C). For example, the tumor suppressor Dab2 (Disabled homolog 2). 80-90% of which is lost in ovarian and breast cancer cells (Bagadi et al., 2007), and ENSA (α-endosulfine), of which overexpression suppresses hepatic tumor growth (Chen et al., 2013), are expected to be short-lived due to their 2ndposition mutations in cancer cells. In contrast, Reg3A (regenerating islet-derived protein 3α), which accelerates pancreatic cancer cell growth (Liu et al., 2013), may be long-lived due to its 2nd-position mutation. Accordingly, it is unclear whether particular 2nd-position mutations contribute to the development of extant malignant phenotypes by faster or delayed degradation of specific proteins (Table 1).

A handful of cellular proteins are shown to be putative substrates of the Ac/N-end rule pathway. For example, a short-lived endogenous p21 $^{\text{Cip1}}$, a CDK inhibitor in mammalian cells, is acetylated at its Nt-Ser, indicating that the resulting Nt-acetyl Ser may control the half-life of p21 $^{\text{Cip1}}$ (Lu and Hunter, 2010). In addition, *C. elegans* RHY-1, an ER acyltransferase-like protein, appears to promote the degradation of CYSL-1, a putative sulfhydrylase/cysteine synthase, potentially by increasing the Nt-acetylation of CYSL-1 (Ma et al., 2012). Furthermore, Nt-Ala of NaV1.5, α subunit of the cardiac voltage-gated sodium channel, is acetylated at the end of heart failure. The resulting

Table 1. Cancer-related proteins with mutations at position 2

Cancer-related proteins	Original sequence	Mutations at position 2	Carcinoma
FARSB (Phe-tRNA ligase β subunit)	MPTVSV	MLTVSV	Head/neck squamous cell
POLR2C (RNA polymerase II subunit)	MPYANQ	MLYANQ	Bladder urothelial
eIF1AX (eukaryotic translation initiation factor 1A)	MPKNKG	M S KNKG	Uterine corpus Endometrial
Dab2 (disabled homolog 2)	MSNEVE	MFNEVE	Breast invasive
ENSA (alpha-endosulfine)	M S QKQE	MYQKQE	Breast invasive
HIST2H2AB (histone H2A type 2-B)	M S GRGK	MLGRGK	Head/neck squamous cell
Rgs2 (regulator of G-protein signaling 2)	MQSAMF	MLSAMF	Breast invasive
Cdk19 (cyclin-dependent kinase 19)	M D YDFK	MYYDFK	Kidney renal clear cell
Reg3A (regenerating islet-derived protein 3- α)	MLPPMA	M P PPMA	Lung adeno
eIF5 (eukaryotic translation initiation factor 5)	MSVNVN	MPVNVN	Lung squamous cell

^{*} Similar to Rgs2 variants in hypertension patients, many cancer cell exomes encode cellular proteins with mutations at position 2 (Kandoth et al., 2013), which are likely to make cancer-related proteins shorter-lived or longer-lived by affecting two branches of the N-end rule pathways. 2nd-position mutations listed above have mutation assessor scores above 2.0, indicating significant impacts on protein function. All information about cancer-related mutations was acquired and analyzed from datasets of The Cancer Genome Atlas (https://tcga-data.nci.nih.gov/tcga).

acetylated Nt-Ala of NaV1.5 is predicted to have an Ac/N-degron, because its steady-state level is significantly lower in typical heart failure than in normal conditions (Beltran-Alvarez et al., 2014). However, additional studies are needed to prove that the Ac/N-end rule pathway mediates the degradation of p21^{Cip1}, CYSL-1, and NaV1.5.

The Ac/N-end rule pathway may alleviate ER stresses because Nt-acetylation prevents cellular protein targeting to the ER, leading to the elimination of retained Nt-acetylated proteins in the cytosol (Forte et al., 2011). Additionally, Teb4 may mediate the degradation of some previously identified substrates, such as the thyroid hormone-activating type 2 iodothyronine deiodinase (D2) (Zavacki et al., 2009), the cholesterol biosynthetic enzyme squalene monooxygenase (SM) (Foresti et al., 2013; Zelcer et al., 2014), and the salt export pump BSEP (Wang et al., 2008), at least in part via their acetylated Ntresidues before or after the targeting to the ER under some stress conditions.

THE AC/N-END RULE PATHWAY IN PLANTS

Nt-acetylation is also an abundant modification in plant cells, which contain a unique chloroplast-localized Nt-acetylase, AtNaa70, as well as 6 Nt-acetylases, NatA-F (Dinh et al., 2015). Loss of the Nt-acetylases NatA. NatB. or NatC in plants results in lethal or pleiotropic deleterious phenotypes (Gibbs, 2015). Despite the presence of SUD2 (suppressor of dry2 defects 1) E3 ligase (Doblas et al., 2013), a sequelog of yeast Doa10 and human TEB4, the Ac/N-end rule pathway has not yet been established in plants. Interestingly, the plant immune receptor SNC1 (suppressor of nod-like protein receptor 1, constitutive 1) has an Ac/N-degron. The SNC1 contains two distinct Nt-variants starting with Met-Met-Asp and Met-Asp Nt-sequences, likely owing to alternative translation initiation: they are differentially regulated by NatA and NatB Nt-acetylases, respectively (Xu et al., 2015) (Fig. 3D). More interestingly, NatA acetylates the Met-Met-Asp Nterminus of SNC1 and triggers its degradation by creating an Ac/N-degron. In contrast, NatB antagonistically stabilizes SNC1 by acetylating the Met-Asp N-terminus. These results uncover the first Ac/N-degron in plants and show the dual roles of Ntacetylation as a destabilizer or a stabilizer for newly synthesized proteins to reach appropriate amounts in a timely manner within a cell (Gibbs. 2015: Xu et al., 2015) (Fig. 3D).

Many plastid precursor proteins are Nt-acetylated and accumulate outside of plastids in the absence of Toc159, a main receptor for photosynthetic proteins. Consequently, Nt-acetylated plastid precursors are most likely to be eliminated by the Ac/N-end rule pathway to avoid the accumulation of unfolded proteins in the cytosol (Grimmer et al., 2014). Furthermore, populations of Nt-acetylated proteins are likely to be modulated in response to external stress in plants. For example, drought and abscisic acid-associated stresses drastically reduce Nt-acetylated proteins through the depletion of NAA10 mRNA and a subsequent decrease in the Naa10 expression levels, leading to impaired growth and development (Linster et al., 2015). It is not clear whether the dramatic changes in Naa10 levels influence the degradation patterns of the global proteome by the Ac/N-end rule pathway.

REGULATION OF THE AC/N-END RULE PATHWAY BY NT-ACETYLASES AND ACETYL-COA

Despite the abundance and irreversibility of Nt-acetylation, the transcription levels of Nt-acetylase subunits change significantly

during development or under stress conditions (Graveley et al., 2011; Linster et al., 2015; Silva and Martinho, 2015). In addition, specific Nt-acetylases are modified by phosphorylation or acetylation for different cellular responses (Malen et al., 2009; Seo et al., 2010). Therefore, altered amounts or activities of specific Nt-acetylases are likely to affect protein expression patterns during development or stress conditions by changing the production of Ac/N-degrons. Intracellular levels of a key metabolic intermediate, acetyl-CoA, also change according to the metabolic states of cells, thus affecting gene expression, cell proliferation, cancer metabolism, autophagy, etc. (Pietrocola et al., 2015). Because acetyl-CoA acts as a cosubstrate for the reaction, its levels are also likely to affect the Nt-acetylation of cellular proteins. For example, reduced acetyl-CoA levels decrease the acetylation of apoptosis effectors, thus linking metabolic conditions with cell death (Yi et al., 2011). The prevalence of Ac/N-degrons in a vast majority of proteins also suggests that changing acetyl-CoA levels may have effects on the global proteome by modulating the activities of Ntacetylases and Ac/N-end rule components.

N-FORMYL DEGRON AND ITS RELEVANT N-END RULE PATHWAY IN BACTERIA

The discovery of Ac/N-degrons led to the hypothesis that the transient Nt-formyl Met (fMet) of nascent proteins acts as an fMet-based N-degron owing to the similarity between acetyl and formyl groups on Nt-Met and their identical localization at Ntermini (Hwang et al., 2010b). The fMet residue occurs in nascent proteins of bacteria and eukaryotic organelles, such as mitochondria and chloroplasts. The inhibition or loss of deformylase activity decreases the levels of chloroplast D2 in Chlamydomonas reinhardtii (Giglione et al., 2003) as well as D2derived reporters and many high-molecular-weight proteins in E. coli by accelerating their degradation via the retained formyl group (Piatkov et al., 2015). The effects are also abrogated by mutations that prevent formylation. These and related findings strongly suggest that fMet acts as a new kind of N-degron by a bacterial fMet/N-end rule pathway (Piatkov et al., 2015). The ATP-dependent protease FtsH may be a fMet/N-recognin that mediates the proteolysis of the E. coli membrane protein YfgM by targeting its fMet (Dohmen, 2015; Piatkov et al., 2015). The fMet/N-end rule pathway is also thought to be involved in protein quality control by the preferential and largely cotranslational degradation of Nt-formylated misfolded or unassembled proteins (Dohmen, 2015; Piatkov et al., 2015).

CONCLUSION

Most cellular proteins are Nt-acetylated, but the universal functions of this modification are largely unknown. The discovery of Ac/N-degrons and the Ac/N-end rule pathway represents a remarkable breakthrough in our understanding of the cellular functions of Nt-acetylation. Recent studies of the Ac/N-end rule pathway have unraveled the complementary crosstalk between two branches of the N-end rule pathway, fMet/N-degrons in bacteria, and a wide range of cellular roles, including the control of protein quality and subunit stoichiometry, the regulation of blood pressure via G-protein signaling, and plant stress and immune responses. Nonetheless, the discovery of Ac/N-degrons and the Ac/N-end rule pathway is still the beginning of an entirely new and broad research field. Because Ac/N-degrons are present in nearly all proteins and can be conditionally modulated, a detailed understanding of the Ac/N-end rule

pathway will set the stage for new therapeutic approaches based on inhibitors or activators of MetAPs, specific Nt-acetylases, and other components of the Ac/N-end rule pathway, including Ac/N-recognins (Hwang et al., 2010a; Lee et al., 2015; Park et al., 2015).

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