N-degron and C-degron pathways of protein degradation

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This perspective is partly review and partly proposal. N-degrons and C-degrons are degradation signals whose main determinants are, respectively, the N-terminal and C-terminal residues of cellular proteins. N-degrons and C-degrons include, to varying extents, adjoining sequence motifs, and also internal lysine residues that function as polyubiquitylation sites. Discovered in 1986, N-degrons were the first degradation signals in short-lived proteins. A particularly large set of C-degrons was discovered in 2018. We describe multifunctional proteolytic systems that target N-degrons and C-degrons. We also propose to denote these systems as “N-degron pathways” and “C-degron pathways.” The former notation replaces the earlier name “N-end rule pathways.” The term “N-end rule” was introduced 33 years ago, when only some N-terminal residues were thought to be destabilizing. However, studies over the last three decades have shown that all 20 amino acids of the genetic code can act, in cognate sequence contexts, as destabilizing N-terminal residues. Advantages of the proposed terms include their brevity and semantic uniformity for N-degrons and C-degrons. In addition to being topologically analogous, N-degrons and C-degrons are related functionally. A proteolytic cleavage of a subunit in a multisubunit complex can create, at the same time, an N-degron (in a C-terminal fragment) and a spatially adjacent C-degron (in an N-terminal fragment). Consequently, both fragments of a subunit can be selectively destroyed through attacks by the N-degron and C-degron pathways.

debrin | proteolysis | ubiquitin | proteasome | N-end rule

The lifespans of protein molecules in a cell range from less than a minute to many days. Regulated protein degradation protects cells from misfolded, aggregated, or otherwise abnormal proteins, and also controls the levels of proteins that evolved to be short-lived in vivo. Some proteolytic pathways can selectively destroy a specific subunit of a protein complex. Such pathways can act as protein-remodeling devices (1). They can either activate or inactivate a protein machine, change its enzymatic specificity, alter its subunit composition, or repair an oligomeric complex, for example, by destroying fragments of a cleaved subunit that are still embedded in the complex. This would allow a replacement of the cleaved subunit by its intact counterpart. Many biological transitions involve remodeling of protein complexes through subunit-selective degradation, in settings that range from cell-division cycles and circadian circuits to cell differentiation and responses to stresses.

One function of protein degradation is the quality control of nascent and newly formed proteins. Selective proteolysis eliminates those proteins (including mutant ones) that fold too slowly, misfold, or do not satisfy other requirements of quality control. Most proteins function as multisubunit complexes, which often assemble cotranslationally. Quality-control systems destroy subunits that are either overproduced relative to other subunits of a complex or do not become incorporated into a complex rapidly enough. The intracellular protein degradation is mediated largely by the ubiquitin (Ub)-proteasome system (UPS) and by autophagy-lysosome pathways, with molecular chaperones being a part of both systems (1–14).

The UPS comprises a set of pathways that have in common two classes of enzymes: E3-E2 Ub ligases and deubiquitylases (DUBs). A Ub ligase recognizes a substrate protein through its degradation signal (degron) (15) and conjugates Ub, a 9-kDa protein (usually in the form of a poly-Ub chain), to an amino acid residue (usually an internal lysine) of the targeted substrate (SI Appendix, Fig. S1A). DUBs deubiquitylate Ub-conjugated proteins and

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edit poly-Ub chains. In addition, DUBs produce free Ub through cleavages of Ub precursors encoded by Ub genes (1–9).

Most UPS pathways also involve a multisubunit, ATP-dependent protease called the 26S proteasome (SI Appendix, Fig. S1A). This protease binds to a ubiquitylated protein substrate through a substrate-linked poly-Ub chain, unfolds the protein using proteasome’s ATPases (often with involvement of the Clcβ48/p97 unfoldase), and processively destroys the protein to ~10-residue peptides (16–22).

Some UPS pathways have nonproteolytic functions as well. A mammalian genome encodes more than 800 E3 Ub ligases, which target, in general, different degrons. The multitude and diversity of Ub ligases underlie the immense functional reach of UPS. Its pathways participate in just about every physiological process in all eukaryotes, play major roles in aging, and are involved in causation of many diseases, from impairments of immunity to cancer and neurodegeneration.

**Terminology for Proteolytic Pathways That Target N-Terminal and C-Terminal**

Degradation signals are features of proteins that make them short-lived in vivo. Such signals determine, in particular, the specificity of UPS. The problem of degradation signals preceded the onset of Ub studies in the 1980s, and remained a mystery until 1986, when the first degradation signals, later termed degrons (15), were discovered at the N-termini of short-lived proteins, an advance made possible by the invention of the Ub fusion technique (SI Appendix, Fig. S1B) (23–26). A set of these N-terminal (Nt) signals, later called N-degrons (15), was referred to by the term “N-end rule,” which related the in vivo half-life of a protein to the identity of its Nt-residue (23–27). Studies over the next three decades identified proteolytic systems that recognize distinct classes of N-degrons and destroy, often conditionally, specific proteins or their natural fragments that bear N-degrons (Figs. 1 and 2 and SI Appendix, Fig. S2A) (22–41). Other studies, in the 1990s and afterward, have also identified many internal degrons, defined as degradation signals whose functionally essential elements do not include either Nt-residues or C-terminal (Ct) residues.

The first example of physiologically relevant Ct-degradation signals, called C-degrons below, was identified in 1996. A specific RNA (SrA) can terminate, in trans, a stalled translation of a bacterial protein while tagging the released protein with the C-tetrasidue ANDENVALAA. This segment acts as a C-degron, targeting a protein for degradation by the proteasome-like bacterial protease ClpXP (42). In 2018, the laboratories of Elledge and Yen discovered a remarkably large set of diverse natural C-degrons in human proteins (SI Appendix, Fig. S3) (43–45). While differing from N-degrons mechanistically and location-wise, C-degrons are topologically analogous to N-degrons. Specific C-degrons and N-degrons can also be associated functionally through their coformation upon a proteolytic cut, as described below.

In 1986, only some Nt-residues were thought to be destabilizing (23). However, later studies by our laboratory showed that every one of the 20 amino acids in the genetic code can act, in cognate sequence contexts, as a destabilizing Nt-residue of an N-degron (Figs. 1 and 2 and SI Appendix, Fig. S2A) (10, 23–26, 35, 38, 39, 46–48). The term N-end rule and its definition, cited above, are not commensurate with involvement of the entire gamut of Nt-residues in protein degradation. This understanding, as well as benefits of accurate notations, is the reason for renaming N-end rule pathways as “N-degron pathways.” Advantages of this terminology include a recall of both the N-tetrasidue ("N") and degradation ("degron"), and the ease of extending this notation from N-degrons to C-degrons.

In sum, proteolytic systems that target N-degrons are proposed to be called the "Arg/N-degron pathway", the "Pro/N-degron pathway", and the "Ac/N-degron pathway" in eukaryotes, the "Imet/N-degron pathway" in eukaryotes and bacteria, and the "Leu/N-degron pathway" in bacteria (Fig. 1 and SI Appendix, Fig. S2A). The prefixes Arg, Pro, Ac, Imet, and Leu specify each pathway by highlighting their unique features, for example, the step of N-arginylation as a part of the Arg/N-degron pathway (Fig. 1G and SI Appendix, Fig. S2A).

A Ub ligase of an N-degron pathway can contain several degron-recognition sites. Such a ligase can bind not only to N-degrons but also to internal degradation signals in proteins that lack an N-degron (26). In the proposed terminology, a substrate of, for example, the Arg/N-degron pathway (Fig. 1G and SI Appendix, Fig. S2A) can be called an Arg/N-degron substrate or an Arg/N-d substrate. Another protein, recognized by the same Ub ligase through a protein’s internal degron, can be denoted as an Arg/N-id substrate: that is, a substrate bearing an internal degron of the Arg/N-degron pathway.

**N-Degron Pathways of Protein Degradation**

The N-degron pathways (formerly “N-end rule pathways”) comprise a set of proteolytic systems whose unifying feature is their ability to recognize proteins containing N-degrons, thereby causing the degradation of these proteins by the 26S proteasome or autophagy in eukaryotes and by the proteasome-like ClpAP protease in bacteria (Fig. 1 and SI Appendix, Figs. S2A and S4) (13, 23, 24, 26, 28, 30–38, 40, 46, 49–54). The main determinant of an N-degron is a destabilizing Nt-residue of a protein. In eukaryotes, an N-degron includes an internal lysine (or lysines) of a substrate protein that acts as the site of polyubiquitylation.

Initially, most N-degrons are pro-N-degrons. They are converted to N-degrons either constitutively (e.g., during the emergence of a protein from a ribosome) or conditionally, via regulated steps. Among the routes to N-degrons are cleavages of proteins by proteases that can expose a destabilizing Nt-residue (29, 55–57). An exopeptidase, for example the mammalian Dpp9 aminopeptidase (it removes dipeptides from N-tetraside), can convert a pro-N-degron at the N-terminus of a specific protein, such as the Syk kinase, to an N-degron (58). The Dpp9 aminopeptidase, Met-aminopeptidases (that remove Met-Met from some nascent proteins) (SI Appendix, Fig. S1D), and endoproteases that include caspases, separases, calpains, and cathepsins, have all been shown to generate N-degrons in vivo through their cleavages of intracellular proteins (26, 29, 55–59). Operationally, these proteases are components of N-degron pathways.

A different and mutually nonexclusive route to N-degrons is through enzymatic Nt-modifications of proteins, including Nt-acetylation, Nt-deamidation, Nt-arginylation, Nt-leucylation, and Nt-formylation of the ε-amino groups of Nt-residues (Fig. 1 and SI Appendix, Fig. S2A). Recognition components of N-degron pathways are called N-recogins. They are either specific E3 Ub ligases or other proteins that can target N-degrons (Figs. 1A and SI Appendix, Figs. S2A and S4). All 20 amino acids of the genetic code can act, in cognate sequence contexts, as destabilizing Nt-residues (Fig. 1A). Consequently, many proteins in a cell are conditionally short-lived N-degron substrates, either as full-length proteins or as protease-generated Ct-frgments (29, 35, 47, 55–57).

Selective degradation of proteins or their natural fragments by N-degron pathways has been shown to regulate a multitude of processes, including: the sensing of oxygen, nitric oxide (NO), heme, and short peptides; the control of subunit stoichiometries in protein complexes; the elimination of misfolded or otherwise abnormal proteins; the degradation of proteins that are retrotranslocated to the cytosol from other compartments, such as mitochondria; the regulation of apoptosis and repression of neurodegeneration; the regulation of DNA repair, transcription, replication, and chromosome cohesion/ segregation; the regulation of G proteins, cytoskeletal proteins, autophagy, gluconeogenesis, peptide transport, meiosis, immunity, circadian rhythms, fat metabolism, cell migration, cardiovascular development, spermatogenesis, and neurogenesis; and the regulation of leaf and shoot development, oxygen/NO sensing, and many other...
processes in plants (see refs. 26, 30–35, 48, 56, 57, and 59 and references therein).

The field of N-degrons and C-degrons is too large for a comprehensive review in a Perspective-size article. Instead of describing all pathways equally briefly, the Arg/N-degron pathway is discussed below in relative detail, followed by much shorter accounts of other pathways.

The Arg/N-Degron Pathway

This eukaryotic pathway targets unacetylated Nt-residues (Figs. 1G and 2B and SI Appendix, Figs. S2A and S4) (23, 55, 59). Nt-Arg, -Lys, -His, -Leu, -Phe, -Tyr, -Trp, -Ile, and -Met (if Nt-Met is followed by a bulky hydrophobic residue) are directly recognized by Arg/N-recognins. Examples of Arg/N-recognins include: the Saccharomyces cerevisiae Ubr1 E3; the mammalian Ubr1, Ubr2, Ubr4, and Ubr5 E3s; the Prt1 and Prt6 E3s of plants; and the mammalian non-E3 autophagy regulator p62/Sqstm1 (Fig. 1G and SI Appendix, Figs. S2A and S3) (13, 26, 32, 33, 47, 49, 50). The Nt-Asn, -Gln, -Glu, and -Asp residues (as well as Nt-Cys, under some conditions) are destabilizing because of enzymatic deamidation of Nt-Asn and -Gln, and Nt-arginylation of Nt-Asp, -Glu, and (oxidized) -Cys (Fig. 1G and SI Appendix, Fig. S2A) (40, 60–62).

Double-E3 Design of the Arg/N-Degron Pathway. Ubr1 is the sole Arg/N-recognin in S. cerevisiae, but the pathway’s targeting complex contains two E3s: the 225-kDa RING-type Ubr1 and the 168-kDa HECT-type Ufd4, in association with their respective E2 enzymes Rad6 and Ubc4/Ubc5 (63) (Fig. 1G). The Ubr1-bound Ufd4 increases the processivity of polyubiquitylation (63). In contrast to Ubr1, Ufd4 is not an Arg/N-recognin. Specifically, Ufd4 does not, by itself, recognize Arg/N-degrons. However, Ufd4 can bind to substrate proteins such as Mgt1, Cup9, and Chk1, through their internal degrons that are also recognized by Ubr1 (63, 64). Exactly how the recognition of an internal degron by both Ubr1 and Ufd4 is achieved within the Ubr1–Ufd4 complex (do Ubr1 and Ufd4 compete for the same elements of a degron, or did...
these E3s evolve to recognize a cognate degron simultaneously?) remains to be understood.

**Substrates of the Arg/N-Degron Pathway and Their Protection by Chaperones.** A molecule of a protein, including a newly formed protein, would be longer-lived if that molecule succeeds, rapidly enough, to become a subunit of a “protective” complex, often a cognate complex in which that subunit normally functions. Stabilization of the subunit would be caused by steric shielding of its degrons within the complex (10). A protection can also be attained through the binding of a vulnerable protein to a molecular chaperone, particularly the Hsp90 system, which comprises Hsp90 and more than 10 of its cochaperones. Hsp90 reversibly binds to at least 20% of cellular proteins, called Hsp90 clients, including most kinases and transcriptional regulators. Hsp90 assists its clients, often repeatedly, in maintaining their active conformations (14).

Oh et al. (64) showed that a weakening of the *S. cerevisiae* Hsp90 system (i.e., an increase in the fraction of Hsp90 clients that are not bound to Hsp90), causes many otherwise long-lived proteins to become short-lived, because of their rapid degradation by the Arg/N-degron pathway. Diverse Hsp90 clients, including Chk1, Kar4, Tup1, Gpd1, Ste11, and also, remarkably, Hsp82 [i.e., the Hsp90 chaperone itself (suggesting that Hsp90 is its own client)], become short-lived substrates of the Arg/N-degron pathway under conditions of hypactive Hsp90 (64). The cited proteins are targeted by Ubr1/Ufd4 through their internal degrons (64). The Arg/N-degron pathway has also been shown to destroy a variety of misfolded proteins (reviewed in refs. 26, 31, and 34).

**Mammalian Ubr1, Ubr2, Ubr4, and Ubr5.** In contrast to *S. cerevisiae*, in which the Ubr1 E3 is the sole Arg/N-recognition (Figs. 1G and 2B), a mammalian genome encodes at least four E3s that can recognize Arg/N-degrons: the 200-kDa Ubr1 and Ubr2, the 570-kDa Ubr4 (p600; Big), and the 300-kDa Ubr5 (Edd1; Hyd) (31) (SI Appendix, Fig. S2A). Ubr1 and Ubr2 are highly sequelogous (similar in sequence) (65) to each other and to *S. cerevisiae* Ubr1 (26, 31). In contrast, the sequelogy (sequence similarity) (65) between, for example, Ubr1 and either Ubr4 or Ubr5, is largely confined to their ∼80-residue UBR domains. [“Sequelog” denotes a sequence that is similar, to a specified extent, to another sequence (65). Derivatives of sequelog include “sequelogy” (similar in sequence) and “sequelology” (sequence similarity). The usefulness and appeal of sequelog and derivative terms stem from the rigor of their evolutionary neutrality. In contrast, the terms “homolog,” “ortholog,” and “paralog” which invoke, respectively, common descent and functional similarity or dissimilarity, are interpretation-laden and often less than precise notations. Homolog, ortholog, and paralog are compatible with the sequelog terminology, and can be used to convey understanding about common descent and biological functions, if this additional information (it is distinct from sequence similarities per se) is actually present (65).] Ubr4, a huge (570 kDa) Arg/N-recognition, functions, in particular: in neurogenesis; in cell migration; in the biogenesis of endosomes; in cardiovascular development and autophagy; in the degradation of misfolded proteins (reviewed in refs. 26, 31, and 34).

Fig. 2. Structural basis of N-degron recognition. The upper diagrams schematically depict the substrate-binding sites of different N-recognins, with corresponding space-filling images indicating electrostatic potential (red, negative; blue, positive) below the diagrams. (A) The substrate-binding site of the human Gid4 Pro/N-recognin (35–37). (B) One of substrate-binding sites (the UBR box, which recognizes basic Nt-residues) of the *S. cerevisiae* Ubr1 Arg/N-recognin. (C) The substrate-binding site of the *E. coli* ClpS Leu/N-recognin, which recognizes bulky hydrophobic Nt-residues (30, 52, 53). Modified with permission from ref. 35.

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podocin, a protein that maintains the renal filtration barrier; and in auxin transport in plants (refs. 66 and references therein). The functions of the Ubr5 Arg/N-recognin include: regulation of Wnt/β-catenin; the degradation of huntingtin, hPXR, Gkn1, and many other proteins; and also specific roles as either an oncoprotein or a tumor suppressor (ref. 67 and references therein). Connections between the functions of Ubr4/Ubr5 and their ability to recognize Arg/N-degrons remain to be understood.

**Johanson-Blizzard Syndrome.** Johanson-Blizzard Syndrome (JBS) patients lack Ubr1 but retain other Arg/N-recognins, including Ubr2, a sequolog (65), and functional analog of Ubr1 (SI Appendix, Fig. S2A). Symptoms of JBS include an exocrine pancreatic insufficiency and inflammation, multiple malformations (e.g., a near-absence of nasal wings), as well as mental retardation and deafness (refs. 26 and 68 and references therein). Ubr1−/− mice exhibit JBS symptoms in a milder form. Mice lacking Ubr2 have other defects, including infertility in males (because of apoptosis of spermatozoa) and genomic instability (31). In contrast to viability of Ubr1−/− and Ubr2−/− mouse strains, double-mutant mice, lacking both Ubr1 and Ubr2, die as midgestation embryos, with defects in neurogenesis and cardiovascular development (refs. 26 and 31 and references therein).

**Structure and Targeting of Arg/N-Degrons.** The main determinant of an Arg/N-degron is a substrate’s specific Nt-residue (1, 22, 26, 29–31). Once an Arg/N-recognin (as a part of a targeting complex) binds to a destabilizing Nt-residue of a substrate, a race against time begins, given the transience of the bound state and the necessity for a successful targeting, to produce a substrate-linked polyUb chain. The synthesis of a proteasome-binding polyUb chain is initiated at an internal lysine of a substrate. This lysine is the second determinant of an Arg/N-degron (SI Appendix, Fig. S5) (1, 22, 24, 26, 31). The third determinant of an Arg/N-degron is a universegmented segment that the substrate-bound proteasome uses to initiate proteolysis (21, 22, 24, 26).

**Subunit Selectivity of Protein Degradation.** The Arg/N-degron pathway can destroy a subunit of a complex while sparing the rest of the complex (1, 26). Subunit selectivity, discovered in 1990, involved in addition the discovery of trans-targeting (SI Appendix, Fig. S5) (1). In this process, an Arg/N-recognin binds to a destabilizing Nt-residue of a subunit that lacks an efficacious second-determinant lysine. It was found that the subunit-bound Arg/N-recognin could polyubiquitylate in trans another subunit of the same complex (if it contained a “suitable” second-determinant lysine), and thereby would target for degradation specifically that subunit rather than the initially bound one. In sum, the multideterminant organization of an Arg/N-degron allows it to be “split” between subunits of a complex, leading to a targeting in trans (SI Appendix, Fig. S5) (1, 26).

**Physiological Arg/N-Degron Substrates.** The list of physiological Arg/N-degron substrates is already large and continues to grow (SI Appendix, Figs. S6–S8). An example of Arg/N-degron substrates that are not cited in SI Appendix, Figs. S6–S8 is Phe-Pink1, a Ct-fragment of the Pink1 kinase. Pink1 is imported into mitochondria and is conditionally cleaved there. The Youle laboratory (see ref. 69 for review) showed that the Phe-Pink1 Ct-fragment is retrotranslocated to the cytosol and is destroyed by the Arg/N-degron pathway. Pink1 phosphorylates, in particular, the E3 Ub ligase parkin and Ub itself. Null mutations in both copies of human PINK1 result in early-onset Parkinson disease, a neurodegeneration syndrome. The uncleaved Pink1 accumulates in the outer mitochondrial membrane (OMM) and recruits parkin to OMM, a step that can lead to an autophagosomemediated engulfment of damaged mitochondria and their destruction in lysosomes. Generation and degradation of the Phe-Pink1 Ct-fragment are a part of circuits that regulate the levels of OMM-bound uncleaved Pink1 and mitochondrial quality control (ref. 69 and references therein).

**Roq1 as a Substrate and Regulator of the Arg/N-Degron Pathway.** A natural Ct-fragment of S. cerevisiae Roq1 acts as both a substrate and regulator of the Arg/N-degron pathway (41). Tunicamycin, a drug that causes protein misfolding in the endoplasmic reticulum (ER), increases the level of ROQ1 mRNA. An artificial increase of ROQ1 mRNA can accelerate the degradation, by the Arg/N-degron pathway, of an ER membrane-embedded reporter protein. The Ynm3 endoprotease can cleave Roq1, generating the Arg-Roq1 Ct-fragment. This cleavage of Roq1 is required for the accelerated degradation of the above reporter. Arg-Roq1 is destroyed, in part, by the Arg/N-degron pathway (41). Remarkably, interactions between Arg-Roq1 and Ubr1 can alter the targeting efficacy of Ubr1 toward its other substrates, such as, for example, Cup9, which bears an internal degron. One possibility is that Arg-Roq1 modulates the specificity and efficacy of Ubr1 under conditions of stress (41).

**Accelerators of Apoptosis as Arg/N-Degron Substrates.** During apoptosis, caspases cleave more than 1,000 different proteins in a mammalian cell. Caspase-mediated cleavages of cellular proteins can generate proapoptotic Ct-fragments, defined as those that increase the probability of apoptosis. Such Ct-fragments often bear destabilizing Nt-residues (SI Appendix, Fig. S6). It was found that the natural proapoptotic Ct-fragments Cys-Ripk1, Cys-Traf1, Asp-Brc1, Leu-Likk1, Tyr-Nedd9, Arg-Bid, Asp-Bcl-xs, Arg-Bimxs, Asp-Epha4, and Tyr-Met were short-lived substrates of the Arg/N-degron pathway (SI Appendix, Fig. S6) (refs. 34 and 55 and references therein). In agreement with these results, even a partial ablation of the Arg/N-degron pathway sensitizes cells to apoptosis (55). In sum, the Arg/N-degron pathway is a regulator of apoptosis, acting largely but not necessarily exclusively (34) as an antiapoptotic circuit (55). By destroying proapoptotic Ct-fragments, the Arg/N-degron pathway contributes to thresholds that prevent a transient or otherwise weak proapoptotic signal from reaching the point of commitment to apoptosis.

**Ubr1 Binds to Caspases.** Weaver et al. (70) showed that the Ubr1Arg/N-recognin of the nematode Caenorhabditis elegans binds to both the procaspase Ced3 and its proteolytically activated form. One substrate of Ced3 is Lin28, a regulator of cell differentiation. Activated Ced3 cleaves Lin28, generating its Nt-Asn-bearing Ct-fragment that is rapidly destroyed by the Arg/N-degron pathway. In ubr1Δ worms the level of Lin28 was increased (as would be expected), but Lin28 was also at most weakly cleaved by Ced3 (70). The latter finding suggested that Ubr1 not only mediates the degradation of the caspase-generated Asn31-Lin28, but may also activate the Ced3 procaspase. If so, the Arg/N-degron pathway might be a previously unknown route for activation of caspases, a most interesting possibility.

**Regulation of Peptide Transport by the Arg/N-Degron Pathway.** In the absence of extracellular di/tripeptides, the S. cerevisiae transcriptional repressor Cup9 shuts off (nearly but not entirely) the PTR2 gene, which encodes the transmembrane peptide importer. This makes cells nearly (but not entirely) incapable of importing di/tripeptides (SI Appendix, Fig. S9). The type 1 and type 2 binding sites of Ubr1 recognize Arg/N-degrons through their binding, respectively, to basic and bulky hydrophobic Nt-residues in either proteins or short peptides (26, 63, 71).

If a cell finds itself in the presence of extracellular di/tripeptides, they are imported inefficiently at first, because of low initial levels of the Ptr2 transporter. However, imported di/tripeptides
that bear destabilizing Nt-residues can bind to the type 1/2 sites of Ubr1. These interactions activate, allosterically, a separate (third) binding site of Ubr1, the one that recognizes an internal degron of the Cup9 repressor (SI Appendix, Fig. S9) (26, 71). The resulting “activated” form of Ubr1 targets Cup9 for degradation, reducing its half-life to ~1 minute and its levels to negligible. As a result, PTT2 is derepressed and the Prt2 transporter is overproduced, greatly increasing the capacity of cells to import di/tripeptides (SI Appendix, Fig. S9) (26, 71). This positive-feedback circuit enables both the budding yeast S. cerevisiae and the fission yeast Schizosaccharomyces pombe to detect the presence of extracellular di/tripeptides and to react by accelerating their uptake (71, 72).

Deamidation of Nt-Asn and -Gln. In S. cerevisiae, Nt-deamidation is mediated by the 52-kDa Nta1 Nt-amidase (Fig. 1G) (40, 73). Remarkably, the bulk of Nta1 is located in the inner mitochondrial matrix (https://yeastgfp.yeastgenome.org/). Nevertheless, a low cytosolic (and presumably nuclear) level of Nta1 suffices to mediate the Arg/N-degron pathway (40, 73). Physiological substrates of yeast Nta1 remain to be discovered. Mitochondrial Nta1 might be a component of a distinct (still to be identified) N-degron pathway in the mitochondrial matrix (26).

In animals and plants, Nt-deamidation is mediated by the Ntn1-encoded, Nt-Asn-specific Nt8-amidase and the Ntaq1-encoded, Nt-Gln-specific Nt8-amidase (SI Appendix, Fig. S2A) (74). Ntn1 and Ntaq1 are present in the cytosol/nucleus, in contrast to the largely mitochondrial S. cerevisiae Nta1 (ref. 74 and references therein). In the fly Drosophila melanogaster, the cleavage, by a caspase, of the antiapoptotic Ub ligase Diap1 generates the short-lived Asn21-Diap1 Ct-fragment that is much less efficacious than full-length Diap1 in repressing apoptosis. Degradation of Asn21-Diap1 requires Nta1 (SI Appendix, Fig. S2A) (75). A virus would benefit from a delay of apoptosis, as this would facilitate the completion of viral replication in an infected cell. Remarkably, a picomolar-like RNA virus induces, through an unknown mechanism, the proteasome-dependent degradation of the Ntn1 Nt8-amidase in infected insect cells, resulting in a partial stabilization of Asn21-Diap1 (76). This way, a viral infection can down-regulate apoptosis, benefiting the virus (76).

Nt-Acetylation. The 60-kDa Ate1 R-transferase catalyzes the conjugation of Arg (provided by Arg-tRNA) to the e-amino group of a specific Nt-residue of a protein. The resulting Nt-Arg can be bound by Arg/N-recogins (Fig. 1G and SI Appendix, Figs. S2 and S4). In mammals, there are six isoforms of R-transferase, produced through alternative splicing of the Ate1 pre-mRNA (SI Appendix, Fig. S2B and C) (ref. 26 and references therein). A number of natural Ct-fragments, including nearly full-length proteins, are either confirmed or putative substrates of the Ate1 R-transferase and the rest of the Arg/N-degron pathway (SI Appendix, Figs. S6–S8).

Arginylation and the Sensing of Oxygen and NO. In 2005, it was discovered that the mammalian Arg/N-degron pathway is a new kind of oxygen (O2) and NO sensor. The NO/O2-dependent oxidation of Nt-Cys converts it to Nt-Cys-sulfinate or Nt-Cys-sulfonate, which can be Nt-arginylated, in contrast to unmodified Nt-Cys (SI Appendix, Fig. S2A) (60, 61). The NO/O2-dependent proteolysis by the Arg/N-degron pathway controls the levels of a subset of proteins that bear Nt-Cys, including Rgs4, Rgs5, and Rgs16 (60, 61). These conditionally short-lived proteins are regulators of specific G proteins.

The Arg/N-degron pathway is also the main sensor of NO/O2 in plants, through the NO/O2-dependent oxidation of Nt-Cys in conditionally short-lived transcription factors that include Rap2.12, Rap2.2, Rap2.3, Hre1, and Hre2 (33, 77, 78). In plants, and possibly in other multicellular eukaryotes as well, the NO/O2-dependent oxidation of Nt-Cys is catalyzed by Cys-oxidases, in addition to a nonenzymatic oxidation of Nt-Cys. In vivo levels of the above transcription factors and the expression of regulons controlled by them underlie adaptations to a broad range of stresses experienced by plants (refs. 32, 33, 77, and 78 and references therein).

The Arg/N-Degron Pathway as a Sensor of Heme. Both mammalian and yeast Ate1 R-transferases are inhibited by low micromolar levels of heme (Fe2+/heme) (79). Hemin also accelerates, in vivo, the degradation of mouse Ate1, thereby acting as both a “stoichiometric” and “catalytic” down-regulator of Nt-arginylation. Thus, in addition to being a sensor of NO, O2, and short peptides, the Arg/N-degron pathway is also a sensor of heme (SI Appendix, Fig. S2A) (79).

Arginylation, Autophagy, and the Arg/N-Degron Pathway. Kwok and colleagues (13, 49, 50) discovered that p62/Sqstm1 (called p62 below), a component of the autophagy-lysosome system, is also a non-E3 Arg/N-recogin that binds to cytosolic proteins that bear either Nt-Ac or specific hydrophilic Nt-residues. p62 mediates the capture of these proteins by autophagy and their subsequent degradation by autophagy-lysosomes (refs. 49 and 50 and references therein). Either a proteasome inhibitor or natural stresses can up-regulate the p62/autophagy branch of the Arg/N-degron pathway, termed the Arg/N-degronp62 pathway (SI Appendix, Fig. S4) (12, 13, 49, 50).

BiP (one of Hsp70 chaperones), calreticulin (another ER chaperone), and protein disulfide isomerase are among ER resident proteins that bear Nt-arginylatable Nt-residues, such as Nt-Asp or Nt-Glu. Upon stresses, including heat shock and unfolded protein response, a fraction of these ER proteins is transferred to the cytosol, followed by their Nt-arginylation. The resulting Nt-Arg-bearing proteins are captured either by the p62 Arg/N-recogin (the p62 Arg/N-recogins, and are degraded by the autophagy-lysosome system the Arg/N-degronp62 pathway) or by one or proteasome (SI Appendix, Fig. S4) (49, 50). In sum, the Arg/N-degron pathway is a major functional link between UPS and autophagy (refs. 12, 13, 49, and 50 and references therein).

The Ac/N-Degron Pathway. About 60% and more than 80% of, respectively, S. cerevisiae and human proteins are irreversibly N*-terminally acetylated (Nt-acetylated) by Nt-acetylases (80). The 2010 discovery of Ac/N-degrons (46) identified a major function of Nt-acetylation, a universally present modification whose significance was, until then, largely obscure. The Ac/N-degron pathway targets proteins for degradation by recognizing their Nt-acetylated Nt-residues (Fig. 1F) (10, 46–48). The E3 Ub ligases (Ac/N-recogins) of this pathway are the ER membrane-embedded yeast Doa10 and its mammalian counterpart Teb4, and also Not4, the E3 subunit of Ccr4-Not, a multifunctional cytosolic/nuclear complex (10, 46, 48).

Schulman and coworkers (81) showed that the Nt-Ac group of a subunit in a protein complex usually increases thermodynamic stability of the complex. The affinity-enhancing effect of Nt-acetylation provides an explanation for at least intermittently long half-lives of many Nt-acetylated proteins. Specifically, natural Ac/N-degrons tend to be conditional, because of their rapid sequestration within cognate protein complexes (10). The functions of the Ac/N-degron pathway (Fig. 1F) include quality control and the regulation of input protein stoichiometries in vivo. For example, S. cerevisiae Nt-Ac-Cog1, a short-lived Ac/N-degron substrate, can be made long-lived by coexpressing Cog2 or Cog3, the Cog1-binding subunits of the Golgi-associated COG complex (10). Analogously, S. pombe Nt-Ac-Hcn1, a short-lived Ac/N-degron substrate, can be stabilized by coexpressing Cut9, a cognate ligand of Hcn1 in the APC/C Ub ligase (10).

Varshavsky
The Pro/N-Degron Pathway
When glucose is low or absent, cells synthesize it through gluco-
neogenesis. In yeast, the main gluconeogenesis-specific cytosolic
enzymes are the Fbp1 fructose-1,6-biphosphatase, the Icl1 isocra-
tic lyase, the Mdh2 malate dehydrogenase, and the Pck1 phospho-
enolpyruvate carboxykinase. When S. cerevisiae grows on a
nonfermentable carbon source such as, for example, ethanol,
the gluconeogenic enzymes are expressed and long-lived. Transition
to a medium containing glucose inhibits the synthesis of these en-
zeymes and induces their degradation, mediated by the multisubunit
GID Ub ligase and the proteasome (ref. 35 and references therein).

We discovered that Gid4, a subunit of GID, is the N-recognin
of a proteolytic system termed the Pro/N-degron pathway (Figs.
1E and 2A) (35). Gid4 recognizes a substrate through its Nt-Pro
residue or a Pro at position 2, in the presence of distinct (but nonunique)
adjacently following sequence motifs. The gluconeogenic en-
zeymes Fbp1, Icl1, Mdh2, and Pck1 bear either Nt-Pro or a Pro
at position 2, and are conditionally short-lived substrates of the
Gid4-dependent Pro/N-degron pathway (Fig. 1E) (35–37). The
structure of Gid4 comprises an antiparallel β-barrel that contains
a deep and narrow substrate-binding cleft (Fig. 2A) (36, 37).

The Eukaryotic fMet/N-Degron Pathway
Nascent proteins bear Nt-Met, encoded by the AUG initiation codon.
In bacteria and in eukaryotic organelles, mitochondria, and chloro-
plasts, formyltransferases Nt-formylate the Met moiety of initiator
Met-tRNAs. Consequently, nascent bacterial proteins start with Nt-
Met. In contrast, proteins synthesized by the cytosolic ribosomes of
eukaryotes bear unformylated Nt-Met, which is often cotranslationally
Nt-acetylated, resulting in Ac/N-degrons (Fig. 1D) (10, 46–48).

In 2015, it was found that Nt-Met residues of nascent bacterial
proteins can act as bacterial N-degrons, termed Imet/N-degrons (Fig.
1C) (38). Remarkably, it was recently discovered that Nt-formylation
of proteins, previously thought to be confined to bacteria and bacteria-
derived eukaryotic organelles, can also occur at the start of translation
by the cytosolic ribosomes of an eukaryote, such as S. cerevisiae (Fig.
1B) (39). Nt-formylation of yeast cytosolic proteins is mediated by the
nuclear DNA-encoded Fmt1 formyltransferase, whose translocation
from the cytosol to the inner matrix of mitochondria was found to be
not as efficacious, even under normal conditions, as had previously
been assumed, and is strongly impaired under conditions of sta-
tionary phase and other stresses (39). The cytosolic retention of Fmt1,
and the resulting upsurge in the levels of Nt-formylated cytosolic
proteins in nutritionally stressed cells, require Gcn2, a protein kinase
(39). It was also discovered that Nt-formylated cytosolic proteins are
selected for targeted degradation by the Psh1 E3 Ub ligase, which
acts as the Imet/N-recognin of the previously unknown eukaryotic
fMet/N-degron pathway (Fig. 1B) (39).

The Bacterial Leu/N-Degron Pathway
The bacterial Leu/N-degron pathway, which does not involve ubiqui-
itylation, was discovered in 1991 (28) and characterized in Gram-
negative bacteria (Fig. 1D) (refs. 26, 30, and 51–54 and references therein).
This pathway comprises the following components: (i) CtpAP,
a proteasome-like, ATP-dependent protease; (ii) CtpS, the 12-kDa
Leu/N-recognin that binds to Nt-Leu, -Phe, -Trp, or -Tyr and delivers
bound substrates to the CtpAP protease; (iii) Aat, an L/F-transferase
that employs Leu-tRNA or Phe-tRNA as a cosubstrate to conjugate
largely Leu (and occasionally Phe) to the N-termini of proteins bearing
Nt-Lys or Nt-Arg (Fig. 1D); and (iv) Bpt, an L-transferase that employs
Leu-tRNA to conjugate Leu to Nt-Asp, -Glu, and (possibly) oxidized
-Cys (Fig. 1D). Vibrio vulnificus, a human pathogen, contains both Aat
and Bpt, while Escherichia coli contains only Aat (51). Physiological
substrates of the E. coli Leu/N-degron pathway include Dps, an
18-kDa DNA-binding protein that compacts the E. coli nucleoid in
staining cells, and the YgiG putrescine-aminotransferase (PATase) (refs.
30 and 52 and references therein). Although E. coli ClpS is nearly 20-
fold smaller than yeast or human Ubr1, there is a significant sequelloing
between the substrate-binding region of ClpS and a functionally
analogous region of Ubr1, suggesting a common descent of bacterial
eukaryotic N-recognins (26, 30).

Studies by Groisman and coworkers indicated that ClpS can target
not only N-degrons (Fig. 1D), but also N-terminus–proximal internal
degrons in bacterial proteins, such as PhoP (ref. 34 and references therein).
In a pathway that regulates PhoP, the MgtC protein competes
with ClpS for the binding to PhoP, and thereby protects PhoP from
degradation. In addition, PhoP, a conditionally short-lived substrate
of ClpS, is a transcriptional repressor of ClpS expression. The resulting
circuits differentially regulate the rates of degradation of specific ClpS
substrates under conditions of low intracellular Mg2+ (54). Because
bacterial ClpS is a sequelog (65) of much bigger eukaryotic Arg/N-
recognins E3s, such as Ubr1 (26, 31), the largely unexplored regula-
tion of the Arg/N-degron pathway in yeast and multicellular eukaryotes
may prove to be at least as functionally rich as the already revealed
regulation of ClpS and the bacterial Leu/N-degron pathway (ref. 54
and references therein).

Eukaryotic C-Degron Pathways
Because of its free carboxyl group, the Ct-residue of a poly-
peptide is stereochemically unique, analogously to the Nt-residue
and its α-amino group. In 2018, the laboratories of Elledge and
Yen discovered a remarkably large set of Ct-degradation signals
in human proteins (SI Appendix, Fig. S3) (43–45). They also
showed that specific E3 Ub ligases of the cullin-RING (CRL) family,
and other E3s as well, can recognize these degrons (43–45).

The authors’ terms for Ct-degradation signals and pathways
that recognize them were, respectively, “C-end degrons” and
“DesCEND” (destruction via C-end degron) (43, 44). For reasons
discussed at the beginning of this paper, we propose to denote
“C-end degrons” as “C-degrons,” and “DesCEND pathways” as “C-
degron pathways” (SI Appendix, Fig. S3). In addition to their suc-
cinctness as well as semantic uniformity vis-à-vis N-degrons, it is easy
to adapt these terms to specific settings. For example, a pathway
mediated by the C-degron–recognizing Kldhc3 subunit of the Crl2
Ub ligase (43, 44) can be called the C-degronKldhc3
pathway.

Functional Aspects of C-degrons
C-degrons can be present in full-length proteins, in truncated proteins
that result from premature termination of translation, and in protein fragments
that form upon proteolytic cuts (SI Appendix, Fig. S3) (43, 44). All such proteins
would be afforded, in vivo, a transient stochastic opportunity to fold or
associate in ways that would shield their C-degrons. A C-degron-
containing polypeptide that fails to shield its C-degron rapidly
enough would face the rising probability of destruction by a cognate
C-degron pathway. This temporal pattern is universal among
C-degrons and other degradation signals, in that it is relevant
to any protein whose degron-based susceptibility to a proteolytic attack
changes as a function of time, with the clock beginning to tick at
the time of protein’s emergence from the ribosomal tunnel.

Corcreation of C-Degrons and N-Degrons upon a Proteolytic
Cut
Usp1 is a mammalian DUB (82). Usp1 forms a heterodimer with
Uaf1, a non-DUB protein (SI Appendix, Fig. S10). Usp1 can autocleave
immediately after its internal Gly-Gly sequence (82). The resulting Ct-
fragment, Gln-Usp1Ct, bears a deamidation/arginylation-dependent
Arg/N-degron (SI Appendix, Fig. S10) (56). Nevertheless, the DUB
activity of autocleaved Usp1 can be transiently maintained, inasmuch
as Usp1Nt, the Nt-fragment of autocleaved Usp1, can remain bound

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to the Gin-Usp1Ct-Ct-fragment within the cleaved Usp1-Uaf1 heterodimer. The Ct-sequence Gly-Gly of the Usp1Nt-Nt-fragment can act as a C-degron, which is recognized by the Kldch2 adaptor subunit of the Crl2 Ub ligase (44). In the resulting mechanism (not yet analyzed in detail), Uaf1 would hold together two Usp1 fragments, allowing them to function, temporarily, as a DUB enzyme, until successful attacks on both fragments by the N-degron and C-degron pathways (refs. 56 and 82 and references therein). Usp1 is the first experimentally addressed setting in which an N-degron and a C-degron can be cocreated upon a cleavage (self-cleavage, in this case) of a full-length protein.

Concluding Remarks

In 1984–1990, studies by our laboratory described the discovery of the first degradation signals (N-degrons) in short-lived proteins; the singular biological significance of UPS; the first physiological functions of ubiquitination, in the cell cycle, DNA repair, protein synthesis, transcriptional regulation, and stress responses; the Arg/N-degron pathway as the first specific UPS pathway; the subunit selectivity of Ub-dependent proteolysis; the first specific poly-Ub chains and their necessity for protein degradation; the Matz2 repressor as the first physiological substrate of UPS; the first nonproteolytic function of Ub (its role as a cotranslational chaperone); and initiated the molecular genetic understanding of UPS, including the cloning of the first E3 Ub ligase (Ubr1), the first DUBs (Ubp1–Ubp3), and the first precursors of free Ub (Ubi1–Ubi4) (refs. 3 and 4 and references therein). Just how broad and elaborate Ub functions are was understood systematically over the next three decades through studies by many laboratories that entered the field in the 1990s and afterward, an expansion that continues to this day.

Studies of N-degron pathways remained a fount of new genetic and biochemical methods for more than three decades, giving rise to the Ub fusion technique, the Ub reference technique, the Ub translocation technique, the split-Ub technique, the Ub sandwich technique, the heat-inducible N-degron (refs. 3, 4, and 25 and references therein), and other methods by other laboratories.

UPS is of major relevance to medicine. Pharmaceutical companies and academic laboratories are developing compounds that target specific UPS components. The fruits of their labors have already become—or will soon become—clinically useful drugs. Work in this arena is producing not only “conventional” inhibitors or activators of specific enzymes, but also drugs that can direct a Ub ligase to target, destroy, and thereby down-regulate any specific protein. Given the broad functional range of N-degron and C-degron pathways, they will be a part of these advances.

Acknowledgments

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41 Szoradi T, et al. (2018) SHRED is a regulatory cascade that reprograms Ubr1 substrate specificity for enhanced protein quality control during stress. Mol Cell 70:1025–1037.e5
Supplementary Information (SI) for

The N-degron and C-degron pathways of protein degradation

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This PDF includes:

Figures S1 to S10.
References for SI reference citations.
**Fig. S1.** The ubiquitin (Ub)-proteasome system (UPS), the Ub fusion technique, the N-terminal processing of newly formed proteins, and specificity of Met-aminopeptidases.

(A) The UPS. Conjugation of Ub to other proteins involves a preliminary ATP-dependent step in which the last residue of Ub (Gly76) is joined, via a thioester bond, to a Cys residue of the E1 (Ub-activating) enzyme. The “activated” Ub moiety is then conjugated to a Cys residue in one of Ub-conjugating (E2) enzymes, and from there, through an isopeptide bond, to a Lys residue of an ultimate acceptor, denoted as “protein”. E2s function as subunits of E2-E3 Ub ligase complexes that can produce substrate-linked poly-Ub chains. These chains have specific Ub-Ub topologies, depending on the identity of a specific Lys residue of Ub that forms an isopeptide bond with C-terminal Gly76 of the adjacent Ub moiety in a poly-Ub chain. Specific poly-Ub chains can confer a processive degradation of a substrate by the 26S proteasome, or other metabolic fates. One role of E3 is the recognition of a substrate’s degradation signal (degron). An individual mammalian genome encodes more than 800 distinct E3 Ub ligases. See also the main text.

(B) The Ub fusion technique. In eukaryotes, linear fusions of Ub to other proteins are cotranslationally cleaved by deubiquitylases (DUBs) at the last residue of Ub, making it possible to produce, in vivo, different residues at the N-termini of otherwise identical proteins (1, 2).

(C) N-terminal processing of nascent eukaryotic proteins by Nα-terminal acetylases (Nt-acetylases) and Met-aminopeptidases (MetAPs) (3). “Ac” denotes the Nα-terminal acetyl moiety. M, Met. X and Z, single-letter abbreviations for any amino acid residue. Yellow ovals denote the rest of a protein.

(D) Met-aminopeptidases (MetAPs) cotranslationally cleave off the N-terminal Met residue of a nascent protein if a residue at position 2 belongs to the set of indicated residues (4, 5). Gly and Pro at position 2 are depicted in a different color because these residues, in contrast to other (indicated) small residues, are very rarely Nt-acetylated after the removal of N-terminal Met (3).
Fig. S2. The mammalian Arg/N-degron pathway and isoforms of the mouse Ate1 R-transferase. Amino acid residues are denoted by single-letter abbreviations. A yellow oval denotes the rest of a protein substrate.

(A) The mammalian Arg/N-end rule pathway. It targets proteins for degradation through their specific unacetylated N-terminal (Nt) residues. “Primary”, “secondary”, and “tertiary” refer to mechanistically distinct classes of destabilizing Nt-residues. Ntan1 and Ntaq1 are N-terminal amidases (Nt-amidases) that convert, respectively, the tertiary destabilizing Nt-residues Asn and Gln to Asp and Glu. The Ate1 R-transferase (arginyltransferase or Arg-tRNA-protein transferase) conjugates Arg, a primary destabilizing residue, to N-terminal Asp, Glu and (oxidized) Cys. “Type 1” and “type 2” refer, respectively, to two sets of primary destabilizing N-terminal residues, basic (Arg, Lys, His) and bulky hydrophobic (Leu, Phe, Trp, Tyr, Ile, and also Met, if the latter is followed by a bulky hydrophobic residue (Φ)). These sets of N-terminal residues are recognized by two distinct substrate-binding sites of N-recognins, the pathway’s E3 ubiquitin ligases Ubr1, Ubr2, Ubr4, and Ubr5. Also indicated is the ability of Arg/N-recognin E3s to recognize specific internal (non-N-terminal) degrons in proteins that lack Arg/N-degrons. See the main text for references and further details. Yet another
branch of this proteolytic system, denoted as the Arg/N-degron\textsuperscript{p62} pathway and discovered by Kwon and colleagues, involves p62/Sqstm1, a non-E3 Arg/N-recognin and a component of autophagy-lysosome pathways (see the main text and Fig. S4) (6-15) and refs. therein).

(B) The exons, including alternative exons (1A/1B and 7A/7B) of the mouse \textit{Ate1} gene, which encodes alternative splicing-derived isoforms of R-transferase. The deduced lengths (in amino acid residues) of the encoded \textit{Ate1} exons are indicated on top.

(D) Mouse R-transferase isoforms that are produced through alternative splicing of \textit{Ate1} pre-mRNA. The terminology of these isoforms (\textit{Ate1}\textsuperscript{1A7A}, \textit{Ate1}\textsuperscript{1A7B}, \textit{Ate1}\textsuperscript{1B7A}, \textit{Ate1}\textsuperscript{1B7B}, \textit{Ate1}\textsuperscript{1A7AB}, \textit{Ate1}\textsuperscript{1B7AB}) is based on the presence or absence of the alternative exons 1A/1B and 7A/7B. \textit{Ate1}\textsuperscript{1A7AB} and \textit{Ate1}\textsuperscript{1B7AB} are minor \textit{Ate1} isoforms that result from the retention of both variants of exon 7 (16, 17).
**Fig. S3.** C-degrons and C-degron pathways in human cells. This diagram is a simplified summary of the 2018 discovery, by the laboratories of Elledge and Yen, of a large set of C-degrons in human proteins (18-20). Amino acid residues are denoted by single-letter abbreviations. A yellow oval denotes a protein substrate upstream of its C-terminus. The indicated C-terminal (Ct) sequences and individual Ct-residues, referred to as C-degrons, are targeted, in conjunction with internal Lys residues of individual C-degron substrates, by a broad range of Ub ligases, largely but not solely of the CRL class (18-20). See the main text for a brief discussion of C-degron pathways and the term “C-degron” for denoting this class of degradation signals.
Fig. S4. Autophagy-mediated versus proteasome-mediated degradation of Arg/N-degron substrates. Amino acid residues are denoted by single-letter abbreviations. A yellow oval denotes the rest of a protein substrate.

Kwon and colleagues discovered that p62/Sqstm1 (called p62 below), a component of the autophagy-lysosome system, is also a non-E3 Arg/N-recognin that binds to cytosolic proteins bearing Nt-Arg or specific hydrophobic Nt-residues. p62 mediates the capture of these proteins by autophagy and their subsequent destruction in lysosomes (6-15). Either a proteasome inhibitor or natural stresses can up-regulate the p62/autophagy branch of the Arg/N-degron pathway (6-15). BiP (one of Hsp70 chaperones), calreticulin (CRT, another ER chaperone), and protein disulfide isomerase (PDI) are among ER-resident proteins that bear Nt-arginylatable Nt-residues such as Nt-Asp or Nt-Glu. Upon stresses, including heat stress and unfolded protein response (UPR), a fraction of these ER proteins is transferred to the cytosol, followed by their Nt-arginylation. The resulting Nt-Arg-bearing proteins are captured either by the p62 Arg/N-recognin and/or by E3 Arg/N-recognins, and are destroyed, respectively, by the autophagy-lysosome system and/or by the 26S proteasome (Fig. S2A) (6-15). In sum, the Arg/N-degron pathway is a major functional link between UPS and autophagy.

The affinity of p62 for Nt-Arg (K_d of ~44 nM) is at least 50-fold higher than that of Ubr1/Ubr2 (12). If so, how can intracellular Ubr1, Ubr2, Ubr4 and Ubr5 E3s (Fig. S2A) compete, in vivo, with the relatively abundant p62 for their binding to proteins bearing Nt-Arg? One possibility is that the affinity of p62 for Nt-Arg may be regulated in living cells. Both basic and aromatic Nt-residues of polypeptides that interact with p62 bind to its ~50-residue zinc finger-like ZZ domain, which is sequelogous (21) to the ~80-residue UBR domain of Ubr1/Ubr2 (6).
Fig. S5. Organization and cis-trans targeting of eukaryotic N-degrons. Amino acid residues are denoted by single-letter abbreviations. A yellow oval denotes the rest of a protein substrate.

(A) Determinants of a eukaryotic N-degron. \textit{d}, a destabilizing Nt-residue. \textit{K}, a “ubiquitylatable” internal Lys residue. See also the main text and refs. (17, 22, 23).

(B and C) Cis versus \textit{trans} polyubiquitylation of an oligomeric Arg/N-degron substrate that can result in the degradation of a subunit that becomes linked to a poly-Ub chain (24).

(D) \textit{Trans}-degradation, in which a specific subunit of an oligomeric protein is polyubiquitylated but is not degraded by the 26S proteasome, for example, because that subunit lacks an unstructured region that is required for the initiation of degradation. Instead, a subunit-selective degradation another, nonubiquitylated subunit takes place. This mode of degradation was discovered by Matouschek and colleagues for oligomeric substrates of the Ub-fusion-degradation (UFD) pathway (25, 26). It remains to be determined whether an analogous \textit{trans}-degradation of an oligomeric Arg/N-degron substrate can also occur.
**Fig. S6.** Natural proapoptotic protein C-terminal (Ct) fragments that are experimentally confirmed substrates of the Arg/N-degron pathway, and some proapoptotic Ct-fragments that remain to be verified as such substrates. The list of either confirmed or putative Arg/N-degron substrates in this and other figures (Figs. S7 and S8) is but a subset of the known list of putative Arg/N-degron substrates, most of which remain to be experimentally confirmed.

<table>
<thead>
<tr>
<th>Protein</th>
<th>Cleavage site</th>
<th>Nt-residue</th>
<th>C-terminal fragment</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Proapoptotic protein fragments that are experimentally confirmed Arg/N-degron substrates</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>Mm</em> Ripk1</td>
<td>321 SLOHDCVPL...S</td>
<td>656</td>
<td>Proapoptotic fragment, generated by caspase-8, of the RIPK1 kinase. Cys-RIPK1 is an N-end rule substrate.</td>
</tr>
<tr>
<td><em>Mm</em> Traf1</td>
<td>152 DLEVDVRALL...A</td>
<td>409</td>
<td>Proapoptotic fragment, generated by caspase-8, of TRAF1, a regulator of apoptosis. Cys-TRAF1 is an N-end rule substrate.</td>
</tr>
<tr>
<td><em>Mm</em> Brca1</td>
<td>1118 DDDLDVWEI...D</td>
<td>1812</td>
<td>Proapoptotic fragment, generated by caspase-3, of the tumor suppressor BRCA1. Asp-BRCA1 is an N-end rule substrate.</td>
</tr>
<tr>
<td><em>Hs</em> Limk1</td>
<td>236 LDEIDLQQ...D</td>
<td>647</td>
<td>Proapoptotic fragment, generated by caspase-3, of LIMK1, a Ser/Thr protein kinase. Leu-LIMK1 is an N-end rule substrate.</td>
</tr>
<tr>
<td><em>Hs</em> Nedd9</td>
<td>626 MDDYDVVLH...F</td>
<td>834</td>
<td>Proapoptotic fragment, generated by caspase-3, of NEDD9, a regulator of cell adhesion. Tyr-NEDD9 is an N-end rule substrate.</td>
</tr>
<tr>
<td><em>Hs</em> Bid</td>
<td>66 HSRLGRIEA...D</td>
<td>195</td>
<td>Proapoptotic fragment, generated by calpains, of BID, a regulator of apoptosis. Arg-BID is an N-end rule substrate.</td>
</tr>
<tr>
<td><em>Mm</em> Bcl-xl</td>
<td>56 SWHLADSPA...K</td>
<td>233</td>
<td>Proapoptotic fragment, generated by calpains, of the antiapoptotic regulator BCL-XL. Asp-BCL-XL is an N-end rule substrate.</td>
</tr>
<tr>
<td><em>Mm</em> Blm</td>
<td>9 ECDRREGG...H</td>
<td>106</td>
<td>Proapoptotic fragment, generated by caspase-3, of the apoptosis regulator BLM. Arg-BLM is an N-end rule substrate.</td>
</tr>
<tr>
<td><em>Mm</em> Epha4</td>
<td>RVLEDPDEA...V</td>
<td>986</td>
<td>Proapoptotic fragment, generated by caspase-3, of the dependence receptor EPHA4. Cys-EPHA4 is an N-end rule substrate.</td>
</tr>
<tr>
<td><em>Mm</em> Met</td>
<td>996 NESYDRAT...S</td>
<td>1379</td>
<td>Proapoptotic fragment, generated by caspase-3, of the dependence receptor MET. Tyr-MET is an N-end rule substrate.</td>
</tr>
</tbody>
</table>

| **Proapoptotic protein fragments that remain to be verified as Arg/N-degron substrates** |
| *Hs* PkCold | 325 EDMQDNSGT...D | 676 | Proapoptotic fragment, generated by caspase-3, of the antiapoptotic protein kinase C5. Asn-PKCo5 is a likely N-end rule substrate. |
| *Hs* PkCold | 350 LDEVDKLMCH...S | 706 | Proapoptotic fragment, generated by caspase-3, of the antiapoptotic protein kinase C8. Lys-PKCo8 is a likely N-end rule substrate. |
| *Hs* Etk | 236 EDYPDDGWGV...H | 675 | Proapoptotic fragment, generated by caspase-3, of the antiapoptotic ETK/BMX tyrosine kinase. Trp-ETK is a likely N-end rule substrate. |
| *Mm* Slk | 432 PDTQDRQQTV...S | 1202 | Proapoptotic fragment, generated by caspase-3, of the SLK kinase, a regulator of actin. Gln-SLK is a likely N-end rule substrate. |
| *Hs* Hpk1 | 381 YDDVDPITP...L | 833 | Proapoptotic fragment, generated by caspase-3, of the Ser/Thr kinase HPK1. Ile-HPK1 is a likely N-end rule substrate. |
| *Hs* Mlh1 | 414 EDKTDISSC...C | 756 | Proapoptotic fragment, generated by caspase-3, of the mismatch repair protein MLH1. Ile-MLH1 is a likely N-end rule substrate. |
verified as such. The name of a protein on the left is preceded by the acronym of a species (Hs, *Homo sapiens*; Mm, *Mus musculus*). Amino acid residues are indicated by single-letter abbreviations. Arrowheads and enlarged residues, in red, indicate P1′ residues of cleavage sites, i.e., the Nt-residues of Ct-fragments of the indicated cleaved full-length proteins. Primary, secondary and tertiary destabilizing N-terminal residues are denoted as Nd0, Nd1 and Nd2, respectively (17, 27). If a depicted cleavage site is the one recognized by a caspase (the exceptions, in the present list, are the calpain cleavage sites that yield Asp-BclXL and Arg-Bid), the site is underlined. The residue number on the left is of the first shown residue of uncleaved protein, numbered as in a full-length protein. The number on the right is of the last residue of a full-length protein.

The first section of this figure describes 10 previously identified natural proapoptotic Ct-fragments, specifically Cys-Ripk1, Cys-Traf1, Asp-Brca1, Leu-Limk1, Tyr-Nedd9, Arg-Bid, Asp-BclXL, Arg-BimEL, Asp-Epha4, and Tyr-Met. They were found to be short-lived substrates of the Arg/N-end rule pathway (28). The figure’s second section describes six other previously identified proapoptotic Ct-fragments (all of them are produced by caspases) bearing destabilizing Nt-residues that can be recognized by the Arg/N-degron pathway. These fragments remain to be verified as Arg/N-degron substrates.

Asn-PkCδ is the Ct-fragment of the protein kinase Cδ (PkCδ) that can be generated by (in particular) caspase-3. This fragment bears Nt-Asn and is proapoptotic, in contrast to the full-length PkCδ kinase (29-33). Lys-PkCθ is the Ct-fragment of the protein kinase Cθ (PkCθ). This fragment can be generated by (in particular) caspase-3, bears Nt-Lys, and is proapoptotic, in contrast to the full-length PkCθ kinase (34). Trp-Etk is the Ct-fragment of the Etk/Bmc tyrosine kinase, a member of the Btk/Tek family of kinases, at least some of which regulate apoptosis. The Trp-Etk fragment can be generated by (in particular) caspase-3, bears Nt-Trp, and is proapoptotic, in contrast to the full-length Etk kinase (35).

Gln-Slk is the Ct-fragment of Slk, a Ste20-related protein kinase that plays a role in regulation of actin fibers. The Gln-Slk fragment can be generated by (in particular) caspase-3, bears Nt-Gln, and is proapoptotic. The concomitantly produced Nt-fragment of Slk is also proapoptotic (36). Ile-Hhp1 is the Ct-fragment of the hematopoietic progenitor kinase 1 (Hpk1), a Ste20-related protein kinase whose functions include stimulation of the stress-activated protein kinases SAPKs/JNKs and the NF-κB transcriptional regulon. The Ile-Hhp1 fragment can be generated by (in particular) caspase-3, bears Nt-Ile, and is proapoptotic, in contrast to the full-length Hk1 kinase (37). Ile-Mlh1 is the Ct-fragment of the mismatch repair Mlh1 protein that can be generated by (in particular) caspase-3, bears Nt-Ile, resides in the cytosol (unlike the full-length nuclear MLH1) and is proapoptotic, in contrast to full-length MLH1 (38).
Calpain-generated C-terminal (Ct) protein fragments that are either identified or predicted substrates of the Arg/N-degron pathway.

<table>
<thead>
<tr>
<th>Ct fragment &amp; description of its full-length precursor</th>
<th>Calpain cleavage site</th>
<th>Ct fragment &amp; description of its full-length precursor</th>
<th>Calpain cleavage site</th>
</tr>
</thead>
<tbody>
<tr>
<td>Glu-Bak; proapoptotic regulator of apoptosis</td>
<td>11 GVGCD E</td>
<td>Arg-Bid; proapoptotic regulator of apoptosis</td>
<td>2 SFNQG R</td>
</tr>
<tr>
<td>Asp-BclXL; antiapoptotic regulator of apoptosis</td>
<td>56 SWHLSA D</td>
<td>Arg-c-Fos; leucine-zipper transcription factor</td>
<td>4 APSQT R</td>
</tr>
<tr>
<td>Glu-IκBα; inhibitor of the NFκB transcription factor</td>
<td>46 VKELER D</td>
<td>Arg-Igfbp2; insulin-like growth factor-binding prot.</td>
<td>6 VNEQHR R</td>
</tr>
<tr>
<td>Asp-Capns1; calpain, regulatory subunit</td>
<td>138 DFGFI D</td>
<td>Arg-Atp2b2; PMCA, pl. membr. Ca^{2+} ATPase</td>
<td>8 FRGLNR R</td>
</tr>
<tr>
<td>Leu-Capn1; calpain, catalytic subunit</td>
<td>23 DKLFG L</td>
<td>Arg-Ankrd2; regulator of muscle differentiation</td>
<td>9 SLDLRL R</td>
</tr>
<tr>
<td>Tyr-Gm1; subunit of metabotropic Glu receptor</td>
<td>11 PLTKS Y</td>
<td>Lys-ica512; member of receptor phosphatase fam.</td>
<td>10 SLRQDR K</td>
</tr>
<tr>
<td>Phe-GluN2a; sub. of NMDA Glu receptor, Ca^{2+} channel</td>
<td>1274 NNLQP F</td>
<td>Asn-DSCR1 (RCAN1); inhibitor of calcineurin</td>
<td>12 HFRNF N</td>
</tr>
<tr>
<td>Arg-GlyT1A; transmembrane Glycine transporter</td>
<td>15 DQNLKT R</td>
<td>Asn-Ca^{2+} 1.1; transmembrane voltage-gated Ca^{2+} channel</td>
<td>16 NTNNA N</td>
</tr>
<tr>
<td>Lys-cortactin; actin-binding cytoskeletal protein</td>
<td>17 FENLAK K</td>
<td>Asn-Bfl-1; antiapoptotic member of Bcl-2 family</td>
<td>18 ARIIF N Q</td>
</tr>
<tr>
<td>Arg-dystrophin; cytoskeletal protein</td>
<td>19 TMVTL R</td>
<td>Glu-Ryr1; Ca^{2+} channel (ryanodine receptor)</td>
<td>20 VAMMT Q P</td>
</tr>
<tr>
<td>Arg-Mef2d; enhancer factor (transcription)</td>
<td>21 EPHES R</td>
<td>Glu-talin; cytoskeletal protein</td>
<td>22 STVLQY Q</td>
</tr>
<tr>
<td>Arg-p39; activator of Cdk5; cyclin-dependent kinase</td>
<td>23 LLRKGP R</td>
<td>Gln-Egrf; transmembrane receptor for EGF</td>
<td>24 LLRLQP E</td>
</tr>
<tr>
<td>Leu-β-catenin; cytoskeletal and transcription factor</td>
<td>25 QQQSY L</td>
<td>Leu-Nf2 (merlin); tumor suppressor, cytoskeleton</td>
<td>26 KLIQK L C</td>
</tr>
<tr>
<td>Arg-caspase-9; caspase, inactivated by calpain</td>
<td>11 RPEIK R</td>
<td>Leu-tropinin T2; cytoskeletal protein</td>
<td>28 PKPSRL L</td>
</tr>
<tr>
<td>Lys-PKCα; protein kinase C-alpha</td>
<td>29 PSEDRK K</td>
<td>Leu-RAD21; subunit of chromosomal cohesion</td>
<td>30 ASNLRL L E</td>
</tr>
<tr>
<td>Phe-PKCγ; protein kinase C-gamma</td>
<td>31 SKRCF F</td>
<td>Leu-STEP33; striatiol-enriched tyrosine phosphatase</td>
<td>32 GSNVS L T</td>
</tr>
<tr>
<td>Leu-CAMK-IV; Ca^{++}/calmodulin kinase IV</td>
<td>19 ASTEN L</td>
<td>Leu-vimentin; intermediate filament protein</td>
<td>34 VSKPD L T</td>
</tr>
</tbody>
</table>

Fig. S7. Calpain-generated C-terminal (Ct) fragments of mammalian proteins that are either identified or predicted substrates of the Arg/N-degron pathway. The entries whose numbers are colored in green are the experimentally identified substrates of the Arg/N-degron pathway (39). The entries whose numbers are colored in black are predicted Arg/N-degron substrates. Each entry cites a calpain-generated Ct-fragment of a protein and
the fragment’s Nt-residue (in red, using three-letter abbreviations for amino acids), followed by a brief description of uncleaved (full-length) precursor protein. A calpain cleavage site, denoted by an arrowhead, is shown using single-letter abbreviations for amino acids. An enlarged P1’ residue (in red) becomes N-terminal upon the cleavage. The indicated residue numbers are the number of the first shown residue of a full-length protein and the number of its last residue, respectively. All entries are mouse proteins, with the exception of #14 and #27, which are human proteins.

#1. Glu-Bak is the proapoptotic Ct-fragment of the apoptotic regulator BAK. Glu-BAK is generated by calpain-1 in vitro and is apparently formed in vivo as well (40).

#2. Arg-Bid. Bid is a 22 kDa member of the BCL-2 family of apoptosis regulators (41-43). Although full-length Bid is already a proapoptotic protein, its Ct-fragments, which can be naturally produced by activated caspas, calpains or granzyme B, can be even more active than intact Bid as proapoptotic protein fragments (44). The cleavage of Bid by calpains produces the 14 kDa Arg-31-Bid fragment (44-47) that has been shown to be a short-lived substrate of the Arg/N-degron pathway (48). #3. Asp-BclXL. BclXL is a 26 kDa antiapoptotic regulatory protein (41, 49). Under conditions that include glucose and oxygen deprivation, BclXL can be cleaved by activated calpain-1, resulting in the 21 kDa Asp61-BclXL Ct-fragment. In contrast to full-length BclXL, Asp61-BclXL has proapoptotic activity (50), and has been shown to be a short-lived substrate of the Arg/N-degron pathway (48).

#4. Arg-c-Fos is the Ct-fragment of the c-Fos transcriptional regulator. c-Fos is targeted for conditional degradation through more than one degron, including the path that includes the cleavage by calpains (51). #5. Glu-IkBα is the Ct-fragment of the IκBα subunit of the NFκB transcriptional regulator. c-Fos is cleaved by activated calpains, resulting in the 21 kDa Asp61-IκBα Ct-fragment of the IκBα subunit of the NFκB-IκBα complex in which the NFκB transcriptional regulator is inhibited by IκBα. The IκBα subunit is targeted for degradation either through a conditional phosphodegron or through a specific calpain-mediated cleavage (52).

#6. Arg-Igfbp2 is the calpain-generated Ct-fragment of the insulin-like growth factor binding protein-2 (53).

#7. Asp-Capns1 is the Ct-fragment of the calpain regulatory subunit that is cleaved by activated calpains (54, 55).

#8. Arg-Atp2b2 is the Ct-fragment of the transmembrane Atp2b2 plasma membrane Ca2+ pump (PMCA) that everts Ca2+ from cells. This pump is activated either by the binding of Ca2+/calmodulin or by the calpain-mediated truncation of Atp2b2 that generates the Arg-Atp2b2 fragment and thereby activates the pump (56).

#9. Leu-Capn1 is the auto-generated, catalytically active Ct-fragment of calpain-1 (57, 58).

#10. Arg-Ankrd2. Ankrd2 (Marp2, Arpp), a member of the MARP (muscle ankyrin repeat protein) family, functions as a negative regulator of muscle differentiation (59). Calpains can produce the 30 kDa Arg103-Ankrd2 Ct-fragment (60).

#11. Tyr-Grm1. Grm1 is the Ct-fragment of the mGluR1α transmembrane metabotropic glutamate receptor (61). Receptors containing the calpain-truncated mGluR1α Ct-fragment can elevate cytosolic Ca2+ but cannot activate Plk-Akt signaling pathways, in contrast to uncleaved receptors (61, 62).

#12. Lys-1ca512. Ica512 (Ptpn) is a member of the transmembrane receptor protein phosphatase family (63). The 43 kDa calpain-generated mouse Lys609-Ica512 Ct-fragment enters the nucleus and acts as a transcriptional regulator (63, 64).

#13. Phe-GluN2a. GluN2a (NMDA-R2a) is a subunit of the N-methyl-D-aspartate receptor (NMDAR), a glutamate receptor that can function as a ligand-gated Ca2+ channel (65, 66). The GluN2b subunit of NMDAR can also be cleaved by calpains (67). Ct-fragments of NR2A and NR2B contain domains required for the association of these subunits with other synaptic proteins. NMDAR receptors lacking a Ct-region of GluN2a could function as glutamate-gated Ca2+ channels but the intracellular traffic of cleaved receptors and their electrophysiological properties were altered (68). #14. Asn-DSCR1
(Rcan1) is the calpain-generated Ct-fragment of the Down syndrome critical region 1 protein Dscr1, which binds to Raf1, inhibits the phosphatase activity of calcineurin, and enhances its degradation. The Asn-DSCR1 fragment does not bind to the Raf1 kinase (69). #15. Arg-Glyt1a is the Ct-fragment of the transmembrane Glyt1a glycine transporter (70). Another Gly transporter, Glyt1b, is also cleaved by calpains, yielding the Arg-Glyt1b fragment (70). These Ct fragments are still active as transporters but are impaired in their ability to remove Gly (an inhibitory neurotransmitter) from synaptic clefts (70). #16. Asn-Ca, 1.1 is the Ct fragment of the voltage-gated transmembrane Ca\(^2^+\) channel. This (apparently) calpain-generated fragment is noncovalently associated with the rest of the channel and can inhibit its activity. Upon dissociation from the channel, the Asn-Ca, 1.1 fragment enters the nucleus and functions as a transcriptional regulator (71-74). #17. Lys-cortactin is the Ct-fragment of cortactin, a protein that regulates actin polymerization (75). #18. Asn-Bfl-1. Bfl-1 is antiapoptotic regulatory protein whose cleavage by calpain-1 generates the Asn72-Bfl-1 proapoptotic Ct-fragment (76). #19. Arg-dystrophin is the calpain-generated Ct-fragment of a major cytoskeletal protein in the skeletal muscle (77). #20. Gln-Ryr1 is the Ct-fragment of the Ryr1 ryanodine receptor, a Ca\(^2^+\) channel in the ER (78) that mediates the efflux of Ca\(^2^+\) from the ER into the cytosol. Calpain-mediated cleavage of Ryr1 increases Ca\(^2^+\) efflux (79). #21. Arg-Mef2d is the Ct-fragment of the Mef2d myocyte enhancer factor 2d, a transcriptional regulator that contributes to neuronal survival, development, and synaptic plasticity (80). #22. Gln-talin is the calpain-generated Ct-fragment of talin, an adaptor protein that interacts with the integrin family of cell adhesion transmembrane proteins (56, 81, 82). #23. Arg-p39 is the calpain-generated Ct-fragment of the p39 activator of the Cdk5 protein kinase (83). The indicated cleavage site is located immediately downstream of two other closely spaced (and strongly conserved) calpain cleavage sites in p39. A cleavage at any one of these sites yields a predicted Arg/N-degron substrate. #24. Gln-Egfr is one of calpain-generated Ct-fragments of the transmembrane epidermal growth factor (EGF) receptor protein kinase (84). Remarkably, all 7 calpain cleavage sites in the cytosol-exposed domain of the 170-kDa EGFR contain P1’ residues that are recognized as destabilizing by the Arg/N-degron pathway (84). #25. Leu-β-catenin is the calpain-generated Ct-fragment of β-catenin, a conditionally short-lived cytoskeletal protein and transcriptional regulator. The Leu-β-catenin fragment is a nuclear protein that activates specific genes in conjunction with other transcription factors (85). #26. Leu-NF2 is the calpain-generated Ct-fragment of NF2 (merlin), a tumor suppressor and cytoskeletal protein. Loss of function NF2 mutants result in autosomal-dominant neurofibromatosis, a predisposition to specific kinds of brain tumors (86). #27. Arg-caspase-9 is the Ct-fragment of caspase-9, which can be inactivated by calpains (87), followed by the (predicted) degradation of the Arg-caspase-9 Ct-fragment by the Arg/N-degron pathway. #28. Leu-troponin T2 is the Ct-fragment of the cardiac troponin T that is produced by calpain-1 in the troponin-containing cardiac myofibril complex (88). #29. Lys-PkCaα is the calpain-generated Ct-fragment of PkCaα, a broadly expressed Ser/Thr kinase of the PKC family (89). Being catalytically active but no longer controlled by the regulatory Nt-domain of the full-length PkCaα, the Lys-PkCaα fragment can be toxic, for example, upon its formation in an ischemic heart (90). #30. Leu-Rad21 is the calpain-generated Ct-fragment of the Sec1/Rad21 subunit of the chromosome-associated cohesin complex (91). The calpain-mediated generation of Leu-Rad21 contributes to the control of chromosome cohesion/segregation, together with processes that include the separatease-mediated cleavage of the same Rad21 subunit of cohesin (91-94). #31. Phe-PkCγ is the calpain-generated Ct-fragment of PkCγ, a Ser/Thr kinase of the PKC family (91-94).
family (89). The *Phe*-PkCγ fragment is constitutively active as a kinase, because it lacks the regulatory Nt-domain of the full-length PkCγ kinase (89). #32. *Leu*-STEP33 is the Ct-fragment of the striatal-enriched STEP61 phosphatase, a brain-specific Tyr-phosphatase whose substrates include the MAPK-family kinases Erk1/2 and p38. The calpain-generated *Leu*-STEP33 fragment lacks phosphatase activity (95). #33. *Leu*-Camk-IV is the calpain-generated Ct-fragment of the Ca$^{2+}$/calmodulin-dependent kinase-IV. This fragment lacks kinase activity (96). #34. *Leu*-vimentin is the calpain-generated Ct-fragment of vimentin, a component of intermediate filaments (97).
Fig. S8. Retention of destabilizing activity (but not necessarily of the identity) of P1’ residues in calpain cleavage sites during evolution of vertebrates. Arrowheads indicate calpain cleavage sites. P1’ residues, which become N-terminal upon the cleavage, are larger and colored. Domain organization and approximate location of a calpain cleavage site are indicated for each protein. The diagrams and indicated residue numbers are of mouse (*Mus musculus* (*Mm*)) proteins. (A) Bak. (B) Grm1. (C) IκBα. (D) GluN2a. (E) Ankrd2. (F) Capsn1. Note the retention, during evolution, of destabilizing activity (but not necessarily of the identity) of P1’ residues in these proteins. See also Fig. S7.
Fig. S9. Regulation of peptide import by the Arg/N-degron pathway in *S. cerevisiae*. (A) The “primed” cascade of Ubr1 phosphorylation in which the Yck1/Yck2-mediated phosphorylation on Ser\textsuperscript{300} of Ubr1 is essential for the normal regulation of peptide import (98). (B) Ubr1-mediated regulation of peptide import, and the involvement of the SPS pathway (99-103). Cup9 is a transcriptional repressor of the regulon that includes *PTR2*, which encodes the major importer of di/tripeptides. In the absence of Ubr1 (in *ubr1Δ* cells), Cup9 becomes relatively long-lived, accumulates to higher levels, and extinguishes expression of *Ptr2*. Therefore *ubr1Δ* cells cannot import di/tripeptides. In wild-type (*UBR1*) cells growing in the absence of extracellular di/tripeptides, a low but non-zero number of Ubr1 molecules have their third substrate-binding site “open” (not autoinhibited) and therefore can target Cup9 for degradation (t\textsubscript{1/2} \sim 5 min) via its internal degron, resulting in a weak but non-zero expression of the *Ptr2* transporter. In wild-type (*UBR1*) cells growing in the presence of extracellular di/tripeptides (some of which bear type-1 and type-2 destabilizing Nt-residues), the imported di/tripeptides interact with the type-1 and type-2 binding sites of Ubr1. This binding allosterically increases the fraction of Ubr1 molecules whose third (Cup9-specific) site is “open” (active). The result is a decrease in the t\textsubscript{1/2} of Cup9 from \sim 5 min to 1 min or less, leading to negligible steady-state levels of Cup9, and consequently to a strong induction of the *Ptr2* transporter (101-103). Also shown is the amino acid-sensing SPS pathway which can influence the import of peptides at least in part through the Yck1/Yck2-mediated phosphorylation of Ubr1 on Ser\textsuperscript{300}. This phosphorylation is required (through a mechanism that remains to be understood) for normal levels of Ubr1 activity in the *Ptr2*-Cup9-Ubr1 circuit (98).
**Fig. S10.** Co-formation of C-degron and N-degron upon Usp1 autocleavage (18, 19, 28).

(A) Sequence alignments, among vertebrates, of the catalytic domains of Usp1 and its autocleavage site, with amino acid residues numbered as in human Usp1. Key residues of the catalytic triad are in red, and are also marked by black circles. The Ub-like domain (Ubl, in green), with its last Gly-Gly 2-residue sequence (in green and a larger font size) abuts the autocleavage site of Usp1. The Gln (Q) P1’ residue of the autocleavage site is in red and a larger font size. This residue, #672 in human Usp1 and #670 in mouse Usp1, becomes N-terminal after the autocleavage of Usp1.

(B) The enzymatic cascade that includes the autocleavage of Usp1 and the Arg/N-degron pathway, which destroys the Gln-Usp1Ct fragment (28).

As described in the main text, the Usp1 deubiquitylase (DUB) forms a heterodimer with Uaf1, a non-DUB protein. Usp1 can autocleave immediately after its internal Gly-Gly sequence (104). Nevertheless, the DUB activity of autocleaved Usp1 can be transiently maintained, inasmuch as Usp1Nt, the Nt-fragment of autocleaved Usp1, can remain bound to the Gln-Usp1Ct Ct-fragment within the cleaved Usp1-Uaf1 heterodimer. The Ct-sequence Gly-Gly of the Usp1Nt Nt-fragment can act as a C-degron, which is recognized by the Kldch2 adaptor subunit of the Clr2 Ub ligase. In the resulting mechanism (not yet analyzed in detail), Uaf1 can hold together two Usp1 fragments, allowing them to function, temporarily, as a DUB enzyme, until successful attacks, on both fragments, by the N-degron and C-degron pathways. Usp1 is the first experimentally addressed setting in which an N-degron and a C-degron can be co-created upon a cleavage (self-cleavage, in this case) of a full-length protein((18, 19, 28, 104) and refs. therein)).
SI References


