The Quality is First: Tagging Nascent Aberrant Proteins for Destruction

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Abbreviations  
ROS: Oxygen Reactive Species; QC: Protein Quality Control; UPS: Ubiquitin Proteasome System; NMD: Nonsense-Mediated Decay; NGD: No-Go Decay; NSD: Non Stop-Decay; mRNA: Messenger RNA; RQC: Ribosome Quality Control; NAC: Nascent Polypeptide Associated Complex; RAC: Ribosome Associated Complex

Editorial  
For all the molecular processes that a cell needs to survive, divide or fulfill, billions of proteins have to be produced. To this end, millions of ribosomes are constantly working, with a synthesis rate of approximately six amino acids per second [1]. This makes the protein synthesis a process with a high probability of errors not only regarding the inefficient folding but also errors in translation and the presence of aberrant mRNAs. In addition, nascent proteins are more susceptible to environmental changes such as high temperatures or oxygen reactive species (ROS). Thus, eukaryotic cells have developed protein quality control (QC) pathways to prevent the accumulation of aberrant proteins that cells identify, label and direct to destruction mainly by the ubiquitin-proteasome system (UPS) [2,3]. Ubiquitin is a 76 amino acid molecule that is covalently attached to protein substrates by a sequential enzymatic cascade catalyzed by three classes of enzymes namely E1 ubiquitin-activating enzymes, E2 ubiquitin-conjugating enzymes, and E3 ubiquitin ligases. E1 enzymes activate ubiquitin in an ATP-dependent fashion, this enzyme transfer ubiquitin to the E2, which further will be used for ubiquitination of E3 bound substrates [4]. E3 therefore are the enzymes responsible to selectively recognize the proteins to be tagged for degradation.

During translation, ribosomes can stop during elongation and stall for different reasons, mainly due to mRNA defects. The ribosome stalling activates both protein and mRNA QC pathways. To date, there are three QC pathways recognized in eukaryotes: nonsense-mediated decay (NMD), no-go decay (NGD), and nonstop-decay (NSD). The NMD is activated by the translation of mRNAs containing premature termination codons, and the NGD pathway is activated with stable mRNA hairpin structures, rare codons, and positively charged pol lysine or polyarginine residues. Finally, the NSD pathway is activated by ‘‘non-stop codon” (those lacking of a stop codon) and damaged mRNAs [2,5].

QC pathways have been deeply studied in terms of mRNA turnover; however, the mechanisms for nascent polypeptide degradation or ribosome recycling have been matter of study only in the recent years. The most studied pathways are the NSD and NGD that are activated by a stalled ribosome. In yeasts, the Hbs1-Dom34 complex, two translation elongation factors (homologs of eFR3-eFR1, respectively), binds to the ribosome probably as a heterodimer. The Hbs1-Dom34 complex together with the ribosome recycling ATPase Rli1 promotes disassembly of ribosomes [6,7]. After ribosome splitting, the E3 ligase Ltn1, together with the ribosome quality control 1 (Rqc1), and the translation-associated element 2 (Tac2) are recruited and the aberrant nascent peptide is ubiquitinated [8]. The association of these three proteins and the ubiquitinated protein recruits the AAA-ATPase Cdc48, Ufd1, and Npl4 complex. Based on the known activity of Cdc48, it has been proposed that it might extract the polyubiquitinated peptides from the dissociated 60S and escort them to the proteasome for degradation (Figure 1A) [9,10].

Ltn1 was the first E3 ubiquitin ligase discovered in this RQC complex. Ltn1 mutation has been related to neurodegeneration in mice and has been found related to aberrant peptide synthetized from a non-stop mRNA [11]. Another E3 ubiquitin ligase related to these processes is Not4, a subunit of the Ccr4-Not deadenylase complex. This E3 ubiquitin ligase targets poly basic-containing nascent chains. In this case, Not4 seems to associate with 80S and polyribosomes ubiquitinating aberrant nascent polypeptides by the recruitment of the E2 enzymes Ubc4p and Ubc5p as well as the proteasome [12]. In addition to these E3 ligases, Hel2 has been also described in these QC pathways. Hel2 together with Asc1 participate in the polypeptide quality control in the RQC pathway, but unlike Ltn1, this E3 ligase may be participating in an upstream stage since in some cases (polybasic chains), the initial recognition step appears to require Asc1 and Hel2. These factors may mediate RNA cleavage [10]. Furthermore, other E3 ligases, Upf1 and Ubr1 have been identified. These are also related to this RQC pathway stimulating the degradation of stalled protein chains. However, the exact moment of their participation have not been elucidated [2,7,13].

Other translation errors that conduct to protein ubiquitination and degradation are nascent polypeptides not correctly folded due to mutations, errors during transcription and RNA processing among others [14]. In order to avoid this, cells uses different complexes such as the nascent polypeptide associated complex (NAC), the ribosome
Some pathways and their mechanisms within the cell are imperative, since these processes are not well regulated. Understanding these QC mechanisms is crucial to achieve substrate specificity. Which would be the consequences if we recognize a defective/misfolded nascent protein? How does the cell regulate the translation rate of certain proteins? Which are the signals to alert the cell on the synthesis or when those are being synthetized in a process named co-translational quality control. This mechanism has not been completely understood; however, some participating proteins have been identified. In yeasts, the nuclear import factor Srp1, and Sts1 couple ribosome-bound nascent polypeptides to proteasomes for degradation [15]. Also, it was found that the E3 ubiquitin ligase Hsl2 and Rkr1 work in parallel with other ubiquitin ligases such as Doa10, Hrd1, Hul5, and Hel2, are able to recognize the misfolded nascent polypeptide and tag them with ubiquitin. After ubiquitination, the Cdc48 complex is recruited to direct the tagged chain to the proteasome for its degradation.

In both cases, the molecular mechanisms for aberrant nascent polypeptides are not completely understood. Many questions remain unsolved. How many E3 ligases are working to recognize aberrant nascent proteins? All proteins could be damaged when are synthetized? All proteins could be damaged when are synthetized? Or are there proteins with more potential to be defective? It depends on the translation rate of certain proteins? Which are the signals to recognize a defective/misfolded nascent protein? How does the cell achieve substrate specificity? Which would be the consequences if these processes are not well regulated? Understanding these QC pathways and their mechanisms within the cell is imperative, since some diseases, mainly neurodegenerative, have been related to misfolded protein degradation impairment.

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