

Rapid Proteasomal Degradation of Posttranscriptional Regulators of the TIS11/Tristetraprolin Family Is Induced by an Intrinsically Unstructured Region Independently of Ubiquitination

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The TIS11/tristetraprolin (TTP) CCCH tandem zinc finger proteins are major effectors in the destabilization of mRNAs bearing AU-rich elements (ARE) in their 3' untranslated regions. In this report, we demonstrate that the *Drosophila melanogaster* dTIS11 protein is short-lived due to its rapid ubiquitin-independent degradation by the proteasome. Our data indicate that this mechanism is tightly associated with the intrinsically unstructured, disordered N- and C-terminal domains of the protein. Furthermore, we show that TTP, the mammalian TIS11/TTP protein prototype, shares the same three-dimensional characteristics and is degraded by the same proteolytic pathway as dTIS11, thereby indicating that this mechanism has been conserved across evolution. Finally, we observed a phosphorylation-dependent inhibition of dTIS11 and TTP degradation by the proteasome *in vitro*, raising the possibility that such modifications directly affect proteasomal recognition for these proteins. As a group, RNA-binding proteins (RNA-BPs) have been described as enriched in intrinsically disordered regions, thus raising the possibility that the mechanism that we uncovered for TIS11/TTP turnover is widespread among other RNA-BPs.

Posttranscriptional regulation plays a central role in the control of gene expression in eukaryotic cells. *cis* regulatory elements, often located in the 3' untranslated region (UTR), influence the localization, translation, and degradation status of mRNAs (1, 2) by binding *trans*-acting microRNAs or RNA-binding proteins (see references 3 and 4 for reviews). AU-rich elements (AREs) are the most common *cis*-acting determinants found in metazoan mRNAs (5, 6). A large variety of AU-rich element-binding proteins (ARE-BPs) affect either positively or negatively the expression of their target ARE-containing mRNAs (reviewed in reference 7). For example, proteins of the TIS11/tristetraprolin (TTP) and AUF-1 families most frequently promote the degradation of their target mRNA (reviewed in references 8 and 9); TIA-1/R proteins inhibit mRNA translation (10), while HuR binding to AREs generally induces stabilization of the RNA messengers (11).

Redundant, additive, or antagonist effects for ARE-BPs have been reported when these factors are expressed in the same cells. Therefore, the outcome of ARE-mediated gene regulation is largely dependent on the functional availability of each ARE-BP (12–14). In this context, the degradation rate for each ARE-BP is an important parameter for determining the output of ARE control.

The proteasome is an essential proteolytic complex whose main function is to degrade damaged or unnecessary proteins. Proteasome substrates are generally marked by covalent attachment of a Lys-48-type polyubiquitin chain to one of the target's internal lysines. The process requires the combined activity of three classes of enzymes: ubiquitin-activating enzymes (E1), ubiquitin-conjugating enzymes (E2), and ubiquitin ligases that ensure specific targeting for degradation (E3). The ubiquitin-proteasome system (UPS) has been described as essential for the degradation of several ARE-BPs and the regulation of ARE mRNAs. A positive correlation was established between AUF1 degradation by the proteasome and ARE mRNA turnover (15). More recent data have also demonstrated that ubiquitination of AUF1 by the E3 ligase b-Trcp leads to its destruction by the proteasome and to the stabilization of its target mRNAs (16). HuR ubiquitination and subsequent proteasomal degradation upon heat shock has also been described (17). Upon proteasome inhibition, HuR is stabilized, which in turn prevents degradation of the TRAIL receptor DR5 mRNA in an ARE-dependent manner (18). Interestingly, HuR is also subject to nondegradative ubiquitination by the p97-UBXD8 complex, which in turn induces the remodeling of mRNAs that contain ARE-bound HuR protein (19).

The mechanism controlling the degradation of the TTP/TIS11 protein family is not fully understood. In mammals, this family is composed of three members, TTP (ZFP36), BRF1 (ZFP36L1), and BRF2 (ZFP36L2), which all participate in ARE mRNA decay (reviewed in reference 8). The stability of TIS11/TTP proteins is known to be defined by their phosphorylation status, which in turn determines their degradation rate (20-22). Deregulation of this control can have dramatic effects on cell fate, leading to defects in the inflammatory response (23), increased radiation-induced lung toxicity (24), and tumorigenesis (25). Previous studies established the proteasome as the proteolytic complex responsible for the degradation of several TIS11/TTP proteins, as its inhibition stabilizes TTP (20) and BRF1 (21). PEST domains, which are susceptible to target proteins to the proteasome, are also predicted in human and mouse TTP, but mutation of these domains did not alter the level of the corresponding proteins (26). An association

Received 9 May 2014 Returned for modification 1 June 2014 Accepted 17 September 2014 Published ahead of print 22 September 2014 Address correspondence to Cyril Gueydan, cgueydan@ulb.ac.be. V.K. and C.G. made equal contributions to this work. Copyright © 2014, American Society for Microbiology. All Rights Reserved. doi:10.1128/MCB.00643-14 of TIS11/TTP proteins with the ubiquitination machinery was also described, as the Cul4B ubiquitin ligase was identified in TTPcontaining messenger ribonucleoprotein particles (mRNPs). However, this enzyme did not target TTP for degradation (27). In addition, nondegradative Lys-63-type polyubiquitination of TTP was reported in response to tumor necrosis factor stimulation (in mouse embryonic fibroblast and human umbilical vein endothelial cell lines), thereby reassigning the biological function of TTP from ARE mRNA degradation to modulation of JNK signaling (28).

The molecular mechanism responsible for TIS11/TTP protein decay has remained elusive, in part due to the fact that proteasome inhibitors can affect the expression of this protein family at several levels. The transcriptional activation of the *Zfp36* gene encoding TTP depends on NF- κ B signaling, whose activity is regulated by the proteasome (20, 29). Moreover, proteasome inhibitors suppress the AKT pathway (30), which plays a central role in regulating BRF1 activity (31, 32). Proper marking of TIS11/TTP proteins as proteasomal substrates by Lys-48-type polyubiquitination has not been described, raising the question of how these proteins are targeted for proteolysis.

In order to get a better insight into the molecular mechanisms controlling the degradation of the TIS11/TTP family of ARE-BPs, we have investigated the mechanisms responsible for the degradation of the unique member of this family found in Drosophila melanogaster, dTIS11. We and others have shown that dTIS11 is able to induce the destabilization of ARE mRNAs (6, 33, 34). Our previous data also implicated the proteasome in the regulation of dTIS11, as pharmacological inhibition of this degradative complex resulted in the accumulation of dTIS11. We report here that dTIS11 is an unstable protein, constitutively produced but rapidly degraded by the proteasome. Recognition of dTIS11 by the proteosomal complex does not require prior ubiquitination, as a nonubiquitinable mutant is efficiently processed by the proteasome. In addition, a proteasome deprived of its ubiquitin-binding domains directly degrades in vitro-translated dTIS11 protein. These results establish dTIS11 as a substrate for the proteasomedependent ubiquitin-independent proteolysis named "degradation by default." Many proteins targeted to this degradative pathway possess unstructured, disordered regions that are thought to be important for proteasomal recognition (35, 36). dTIS11 may be an intrinsically disordered protein (IDP), as predicted by bioinformatics analysis, as we found that dTIS11 shares the main characteristics of IDPs, namely, high sensibility to proteolysis and heat resistance. Finally, we report that the mammalian family founding member TTP undergoes the same degradation process, establishing a novel and conserved regulatory mechanism for TIS11/TTP proteins.

MATERIALS AND METHODS

Reagents. Clasto-lactacystin beta-lactone, epoxomicin, thapsigargin, bafilomycin A1, and NH₄Cl were purchased from Sigma-Aldrich. Okadaic acid was purchased from LC Laboratories. Puromycin was purchased from Invivogen. MG132 was purchased from UBPbio.

Plasmids. Expression vectors for dTIS11_{V5/HIS}, dDREDD_{V5/HIS}, and GFP-DCP1_{V5/HIS} were obtained via reverse transcription-PCR (RT-PCR; for dTIS11 and dDREDD) or amplification from the previously described green fluorescent protein (GFP)-DCP1 construct (37) (kindly provided by E. Izaurralde, Max Planck Institute for Developmental Biology) and then insertion into the pMT/V5-His vector (Life Technologies). GFP-DCP1-CdTIS11_{V5/HIS} was generated by fusion of the coding sequence for

GFP-DCP1 with that of the dTIS11 C terminus (amino acids 214 to 436) and insertion into the pMT/V5-His vector. Expression vectors for hemagglutinin (HA)-tagged ubiquitin (Ub) and FLAG-tagged dTIS11 were obtained by replacement of the start codon by an HA or FLAG tag via RT-PCR and insertion into the pAc5.1/V5-His vector (Life Technologies) for HA-ubiquitin and into the pMT/V5-His vector for FLAG-dTIS11. A lysine-less dTIS11KR mutant was designed by replacement of all 14 lysines with arginines. The corresponding DNA sequence was synthesized by Genscript and subcloned into the pMT/V5-His vector (Life Technologies). The Ub^{G76V}-GFP_{V5/HIS} coding sequence was generated by PCR from the previously described pcDNA3-UbG76V-eGFP-V5His_20aa construct (38) (Addgene) and inserted into the pMT/V5-His vector. The expression vector for PARP12_{HIS} was kindly provided by D. Hutin (Laboratoire d'Immunobiologie, ULB). The lysine-less TTP mutant form was obtained by replacement of all lysines with arginines, synthesized by Genscript, and subcloned into the pcDNA3.1/Myc-His vector (Life Technologies). All constructs were verified by sequencing. The primers used for the cloning procedures are available on request.

Cell culture, treatments, RNA interference (RNAi), and transfection. *Drosophila* S2 cells (Invitrogen) were maintained in serum-free medium (Express Five; Life Technologies) at 24°C. Cells were transfected with FuGENE HD (Promega) according to the manufacturer's instructions. For pMT constructs, transcription driven by the metallothionein promoter was induced by overnight incubation with CuSO₄ (0.5 mM). RAW 264.7 and HEK293T cell lines were maintained in Dulbecco's modified Eagle's medium supplemented with 10% fetal bovine serum. RAW 264.7 cells were activated with lipopolysaccharide (100 ng/ml; Sigma-Aldrich) for 4 h. HEK293T cells were transfected with Lipofectamine 2000 (Life Technologies), following the manufacturer's instructions.

For RNA interference, double-stranded RNA (dsRNA) was produced with a MEGAscript RNAi kit (Life Technologies), following the manufacturer's instructions. S2 cells were incubated with 10 μ g/ml dsRNA targeting β 5 for 4 days or *gfp* or *dtis11* for 7 days.

For mRNA half-life measurements, transcription was blocked by using actinomycin D (5 μ g/ml; Sigma-Aldrich), and the cells were harvested after 30 min, 1 h, 2 h, and 3 h. Proteasome inhibitors were used in 4-h treatments at final concentrations of 5 μ M for clasto-lactacystin-beta-lactone, 40 μ M for MG132, and 10 μ M for epoxomicin. Lysosome function inhibitors were used in 16-h treatments at final concentrations of 10 mM for NH₄Cl, 1 μ M for thapsigargin, and 150 nM for bafilomycin A1. Puromycin was used at 50 μ g/ml.

For pulse-chase analyses, S2 cells maintained in serum-free medium (ESF 921; Expression Systems) were washed in labeling medium lacking methionine for 1 h, then labeled for 2 h in medium supplemented with 0.1 mCi/ml [³⁵S]methionine (43.48 TBq/mmol; PerkinElmer).

Western blotting and antibodies. Cells were lysed in EBC buffer (120 mM NaCl, 0.5% NP-40, 50 mM Tris-HCl [pH 8]) supplemented with phosphatase inhibitors NaF (100 mM), Na₃VO₄ (0.2 mM), and beta-glycerophosphate (10 mM), and also Complete EDTA-free protease inhibitor cocktail (Roche Applied Science). For ubiquitination detection experiments, cells were lysed in denaturing buffer (6 M urea, 10% glycerol, 20 mM Tris-HCl [pH 8], 1 M NaCl, 1% Triton X-100) supplemented with 15 mM *N*-ethylmaleimide (Sigma-Aldrich) and 25 μ M clasto-lactacystin-beta-lactone. Protein extracts were then incubated on ice for 10 min and cleared by centrifugation. For nickel affinity chromatography purification, protein extracts were incubated with equilibrated Ni-nitrilotriacetic acid (NTA) Superflow beads (Qiagen) at 4°C for 1 h in the corresponding lysis buffer supplemented with 10 mM imidazole, washed in lysis buffer with 25 μ M imidazole, and eluted with 150 μ M imidazole lysis buffer.

Western blotting was performed with the following primary antibodies: anti-V5, anti-FLAG, anti-GFP, anti-Myc, and antiactin (Sigma-Aldrich) and anti-glyceraldehyde 3-phosphate dehydrogenase (anti-GAPDH; Santa Cruz Biotechnology). The detection of dTIS11 protein by Western blotting was performed with a previously described anti-dTIS11 antibody (39). The antiubiquitin specific antibody (FK2 clone; Enzo Life science) was used as previously described (40). The anti-TTP (Carp-3) and anti-PARP12 antibodies were kindly provided by W. Rigby (Dartmouth Hitchcock Medical Center) and D. Hutin, respectively. Quantifications were performed with the ImageJ software.

Northern blot assays and quantification. Total RNA was purified using Tri reagent (MBI Fermentas) and analyzed as described previously (33). Quantification of the radioactive signals was performed with a PhosphorImager (GE Healthcare).

In vitro translation and 20S proteasome degradation assay. *In vitro* translation was performed in rabbit reticulocyte lysate (Promega) in the presence of [³⁵S]methionine (0.4 mCi/ml; 43.48 TBq/mmol) according to the manufacturer's instructions. *In vitro*-translated or cell lysate purified proteins were incubated with 1 µg or 0.5 µg purified human 20S proteasome (Enzo Life Sciences), respectively, in 20S buffer (20 mM Tris-HCl [pH 7.2], 1 mM EDTA, 1 mM dithiothreitol [DTT]) at 37°C for the indicated times. ³⁵S-labeled proteins were revealed by SDS-PAGE followed by sodium salicylate gel staining and fluorography.

FastPP assay. The fast pulse proteolysis (FastPP) assay was performed as described previously (41). Briefly, protein extracts were incubated with 1-mg/liter or 10-mg/liter thermolysin from *Bacillus thermoproteolyticus rokko* (Sigma-Aldrich) in TL buffer (10 mM CaCl₂, 20 mM Na₃PO₄ [pH 7.2], 150 mM NaCl, 5 mM DTT) for 1 min at the indicated temperatures in a thermocycler. After 1 min, proteolysis was stopped by addition of 17 mM EDTA.

Heat resistance assay. Protein extracts were heated at 95°C for 10 min and ultracentrifuged for 10 min at 100,000 \times g. The supernatant was retrieved, and the pellet was resuspended in 1× Laemmli buffer.

Bioinformatics analysis. Disorder prediction was performed using default settings with DISOPRED2 (42), IUpred (43), Pondr-fit (44), and DynaMine (45). Protein sequence alignment was performed with Clust-alW (46).

RESULTS

dTIS11 is rapidly degraded in Drosophila S2 cells. Protein halflives are highly variable, ranging from a few minutes to several days in eukaryotes (47, 48). Protein decay is thus a key regulatory step, as its balance with protein production establishes a dynamic equilibrium that defines the steady-state level of expression. As a central effector in AU-rich element-mediated mRNA decay (33), the availability and activity of dTIS11 can affect the expression of many target messengers (6, 34). We sought to assess the stability of dTIS11 by measuring its half-life. Drosophila S2 cells were treated with the translation inhibitor puromycin for 0 to 6 h, and resulting levels of dTIS11 were evaluated by Western blotting. Specific detection of dTIS11 by Western blotting was verified upon RNAi inhibition of the corresponding gene (Fig. 1C). Figure 1A shows that dTIS11 was rapidly degraded, with a half-life of approximately 1.5 h ($t_{1/2} = 92 \pm 26$ min [mean \pm standard deviation, or SD]). Slower-migrating bands may correspond to phosphorylated dTIS11, as in vitro dephosphorylation by alkaline phosphatase results in comigration as a single 55-kDa band (data not shown). In parallel, we assessed the stability of a fusion dTIS11 protein, doubly tagged with a V5 epitope and a $6 \times$ His motif (dTIS11_{V5/HIS}), in S2 cells. Figure 1B shows that dTIS11_{V5/HIS} has a half-life similar to that of endogenous dTIS11 protein ($t_{1/2} = 118 \pm 9$ min). In contrast, the control GFP-DCP1 $_{\rm V5/HIS}$ produced from the same plasmid backbone and the endogenous actin protein were stable, with half-lives longer than 6 h. Decay curves calculated from four independent experiments confirmed that endogenous and V5/Histagged dTIS11 proteins are degraded with similar kinetics (Fig. 1D), which indicates that $dTIS11_{V5/HIS}$ may serve as a model for the analysis of the dTIS11 degradation process.

Half-life measurement using translation inhibitors such as puromycin is indirect and can be affected by side effects of the drug. We therefore sought to confirm the dTIS11 half-life measurement by pulse-chase analysis. In the absence of immunoprecipitaion (IP)-grade antibodies for dTIS11, the experiment was performed on the tagged protein dTIS11_{V5/HIS}. S2 cells expressing dTIS11_{V5/HIS} were cultured in medium containing [³⁵S]methionine and then chased with label-free medium and harvested at several time points. Total protein extracts were submitted to Ni²⁺-affinity chromatography, and purified dTIS11_{V5/HIS} was analyzed by fluorography after SDS-PAGE separation. The stability of dTIS11_{V5/HIS} directly measured by pulse-chase analysis ($t_{1/2} = 108 \pm 17 \text{ min}$) (Fig. 1E, lanes 1 to 6) was similar to that obtained in translation inhibition assays. Both experiments established dTIS11 as an unstable short-lived protein.

dTIS11 degradation requires proteasome activity. In eukaryotic cells, two pathways are responsible for most proteolytic events: slow selective digestion in the lysosome and targeted degradation by the proteasome. As dTIS11 appears to be efficiently and specifically degraded, we investigated the implications for the proteasome in this process. We previously showed that the proteasome inhibitor clasto-lactacystin-beta-lactone (Clasto) increases the level of dTIS11 protein detectable in S2 cells (33). Pulse-chase analysis of dTIS11_{V5/HIS} degradation performed in S2 cells treated with Clasto suggested that this protein is indeed degraded in a proteasome-dependent manner, as its half-life increased from 108 min to more than 4 h in the presence of Clasto (Fig. 1E, lanes 7 to 10). To confirm this result and to assess the potential role of lysosomal function in endogenous dTIS11 proteolysis, S2 cells were incubated in the presence of inhibitors of both pathways. Treatment with MG132, Clasto, or epoxomicin, proteasome inhibitors of three different classes (49), induced a marked accumulation of dTIS11 in S2 cells (Fig. 2A, compare lanes 1 and 2 with lanes 3 to 5). In contrast, dTIS11 did not accumulate in response to treatment with bafilomycin and NH₄Cl (Fig. 2A, compare lanes 1 and 2 with lanes 6 and 8) at doses where these inhibitors of lysosomal function block the accumulation of Lysotracker in S2 cells (data not shown). Inhibition of the lysosomal function by thapsigargin treatment provided similar results (Fig. 2A).

As observed for endogenous dTIS11, the three classes of proteasome inhibitors increased the accumulation of dTIS11_{V5/HIS} in transfected S2 cells (Fig. 2B, upper lane). In contrast, the accumulation levels of the GFP-DCP1_{V5/HIS} control and of the endogenous actin protein were not modified by these treatments (Fig. 2B, lower lanes). Moreover, efficient inhibition of proteasome subunit β 5 by RNA interference led to endogenous dTIS11 accumulation (Fig. 2C). Together, our results thus indicate that dTIS11 is highly unstable in *Drosophila* S2 cells because of its continuous degradation by the proteasome.

Ubiquitin-independent degradation of dTIS11 by the pro-teasome. Because proteasomal activity is required for dTIS11 degradation, we investigated the implication of the ubiquitination system in dTIS11 processing. First, we sought to directly detect dTIS11 ubiquitination. This can be achieved by observing coprecipitation of ubiquitin with the protein of interest. Polyubiquitinated proteins were immunoprecipitated from S2 cell extracts by using an antiubiquitin specific antibody as previously described (40). Under these conditions, we failed to detect the endogenous dTIS11 protein in the immunoprecipitated fraction (data not shown). To assess the implication of the ubiquitin system in the

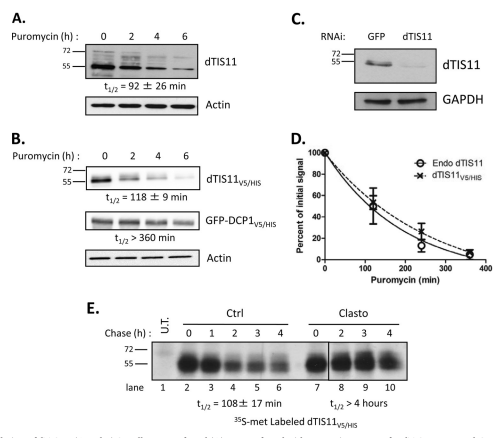


FIG 1 Rapid degradation of dTIS11. (A and B) S2 cells untransfected (A) or transfected with expression vectors for dTIS11_{V5/HIS} and GFP-DCP1_{V5/HIS} (B) were treated with puromycin for the indicated times. Endogenous dTIS11 was detected by Western blotting with a dTIS11-specific antibody (see Materials and Methods). dTIS11_{V5/HIS} and GFP-DCP1_{V5/HIS} were detected by Western blotting with an anti-V5 antibody. (C) Specificity of the anti-dTIS11 antibody. S2 cells were treated with 10 μ g/ml double-stranded RNA targeting GFP or dTIS11 for 7 days. Western blott analysis was performed with anti-dTIS11 and anti-GAPDH antibodies. (D) Quantification of dTIS11 and dTIS11_{V5/HIS} half-lives. Results of four independent experiments performed as described for panels A and B were quantified with the ImageJ software, and the ratios of dTIS11 or dTIS11_{V5/HIS} and actin signals were plotted. The amount of dTIS11 or dTIS11_{V5/HIS} at time zero of puromycin treatment was set to 100% in each experiment and plotted on a linear graph. Half-lives were determined by exponential regression. Error bars represent SD for each experimental condition. (E) Pulse-chase analysis of dTIS11_{V5/HIS} was precipitated on proteasome inhibition. Untransfected (U.T.) or dTIS11_{V5/HIS}-expressing S2 cells were labeled with [³⁵S]methionine-containing medium for 90 min, then chased with unlabeled medium supplemented with Clasto or dimethyl sulfoxide (DMSO) as a control, for the indicated times. dTIS11_{V5/HIS} was precipitated on nickel-coupled agarose beads, and precipitates were resolved by SDS-PAGE. Acrylamide gels were analyzed by fluorography. Half-lives were determined by exponential regression after quantification of results of three independent experiments. Vertical bars indicate cuts between separated parts of the same blot.

degradation of endogenous dTIS11, S2 cells were treated with the E1 ubiquitin-activating enzyme inhibitor PYR-41 (50). Although this compound is active in S2 cells (51), we did not detect any increase of dTIS11 accumulation when S2 cells were treated with sufficient levels of PYR-41 (Fig. 3A).

In order to potentiate the ubiquitination process and facilitate its detection, S2 cells were cotransfected with expression vectors coding for dTIS11_{V5/HIS} and for a HA-tagged ubiquitin (HA-ubiquitin). As a positive control, we also transfected cells with an expression vector for Ub^{G76V}-GFP_{V5/HIS}, since this fusion is known to undergo efficient polyubiquitination and subsequent proteasomal degradation (38). The cells were then treated with MG132 to prevent degradation of the ubiquitinated moieties. Proteins were extracted in highly denaturing urea buffer supplemented with the deubiquitinase inhibitor *N*-ethylmaleimide to prevent hydrolysis of the ubiquitin-substrate bonds in the extract. Affinity chromatography was then performed on Ni²⁺-coupled beads for isolation of dTIS11_{V5/HIS} or Ub^{G76V}-GFP_{V5/HIS}, followed by separation via SDS-PAGE. Western blotting with an anti-V5 antibody showed that precipitation of both proteins occurred with similar efficiency (Fig. 3B, left panel). Interestingly, Ub^{G76V}-GFP_{V5/HIS} was seen as two bands that were separated by a distance that matched the attachment of a single ubiquitin particle (8 kDa). Blotting of Ni²⁺-purified extracts with anti-HA antibodies showed the expected ladder/smear, which corresponded to polyubiquitinated Ub^{G76V}-GFP_{V5/HIS}, but no HA-ubiquitin signal associated with dTIS11_{V5/HIS} (Fig. 3B, right panel), which suggests that dTIS11_{V5/HIS} is not predominantly ubiquitinated under these conditions.

Detection of polyubiquitination may be limited by the low sensitivity of the assay, rapid deubiquitination in the protein extracts, and/or specific degradation of the modified moieties. Therefore, we could not exclude that dTIS11 ubiquitination might still occur in S2 cells. We thus sought to determine whether cryptic ubiquitination is necessary for dTIS11 recognition by the proteasome. Because degradative ubiquitination is generally initiated by the covalent binding of a ubiquitin molecule to an internal lysine of the substrate (52), we analyzed the behavior of dTIS11 when mu-

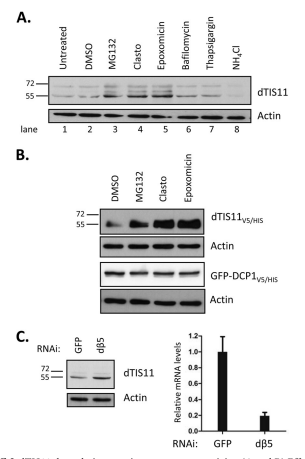


FIG 2 dTIS11 degradation requires proteasome activity. (A and B) Effects of proteasome and lysosomal activity inhibitors on dTIS11 accumulation. S2 cells untransfected (A) or transfected with an expression vector encoding dTIS11_{V5/HIS} or GFP-DCP1_{V5/HIS} (B) were treated with the indicated inhibitors for 4 h (proteasome inhibitors) or 16 h (lysosomal activity inhibitions) before harvesting and analysis by Western blotting. (C) RNAi inhibition of the proteasome leads to dTIS11 accumulation. S2 cells were treated with double-stranded RNA for GFP or d β 5. (Left panel) Endogeneous dTIS11 and actin levels were assessed by Western blotting as described for Fig. 1. (Right panel) d β 5 mRNA levels were measured by quantitative RT-PCR and normalized with RPL32 mRNA levels.

tated for all 14 lysines conservatively replaced with equally positively charged arginine residues (dTIS11^{KR}_{V5/HIS}). S2 cells were transfected with expression vectors encoding dTIS11_{V5/HIS} or the dTIS11KR with puromycin for 4 h. Western blot analysis showed that simultaneous mutation of all ubiquitin-accepting sites did not stabilize dTIS11 (Fig. 3C), while the mutant remained a substrate for the proteasome, as shown by the dTIS11KR_{V5/HIS} accumulation upon MG132, Clasto, or epoxomicin treatment (Fig. 3D). Importantly, simultaneous replacement of as many as 14 residues may extensively alter the protein structure, leading to the production of a polypeptide recognized as aberrant and targeted for degradation, regardless of its natural degradative process. We thus verified dTIS11^{KR}_{V5/HIS} functionality in promoting ARE mRNA decay. To do this, we compared the abilities of dTIS11_{V5/HIS} and dTIS11^{KR}_{V5/HIS} to destabilize a reporter mRNA construct (Luc-ARE; described in a previous study [33]). S2 cells expressing the reporter mRNA were cotransfected with expression vectors for dTIS11_{V5/HIS},

dTIS11^{KR}_{V5/HIS}, or GFP_{V5/HIS} as a control. We observed by Northern blotting and subsequent quantification of the Luc-ARE mRNA half-life that, in contrast to GFP, wild-type and lysine-less dTIS11were able to destabilize the target reporter mRNA to a similar extent (Fig. 3E), arguing in favor of the production of dTIS11^{KR}_{V5/HIS} as a properly folded and active protein.

While the vast majority of ubiquitination processes occur on internal lysines of the substrate, examples of alternative ubiquitination have been reported in which the N-terminal amino acid is used as the ubiquitination accepting site (see reference 53 for a review). We thus verified that modifying the environment of dTIS11 at its N terminus by addition of either a FLAG or a GFP coding sequence upstream of the *dtis11* start codon AUG (FLAG-dTIS11 and GFP-dTIS11) did not alter its degradation pattern. As for the lysine-less mutant, this modification did not stabilize the protein (Fig. 3F).

Taken together, our results suggest that while dTIS11 is a proteasome substrate in *Drosophila* S2 cells, its degradation does not require prior marking by the ubiquitination machinery.

Direct degradation of dTIS11 by the 20S proteasome in the absence of ubiquitin and ATP. Ubiquitin-independent proteasome-dependent degradation has been reported for a limited but growing number of proteins. It has been described to involve different forms of the proteasome: the 20S proteasome, the 26S proteasome, which has been most extensively studied, or other proteasome variants (see reference 36 for a recent review). The 20S core is formed by four heptameric rings that are stacked as a barrel-shaped complex, defining a narrow 53-Å degradation chamber shut by the N termini of its outer subunits. As such, and while bearing all three proteolytic activities of the proteasome, it alone stands gated, unable to recognize ubiquitin or to process large folded proteins. In contrast, association of the 20S core with 19S caps into the 26S proteasome allows for the degradation of ubiquitinated proteins, as the 19S provides opening of the 20S gated entrance, ubiquitin recognition, and ATP-dependent substrate unfolding (reviewed in reference 54).

As we established dTIS11 as a bona fide substrate for the proteasome with no requirement for prior ubiquitination, we investigated the ability of the 20S proteasome to directly recognize and process dTIS11. First, we tested whether purified 20S could degrade dTIS11 extracted from Drosophila S2 cells. For this, S2 cells were transfected with expression vectors for dTIS11_{V5/HIS}, GFP-dTIS11^{KR}_{V5/HIS}, or several negative controls (GFP_{V5/HIS}, GFP-DCP1_{V5/HIS}, Ub^{G76V}-GFP_{V5/HIS}, and dDREDD_{V5/HIS}). The corresponding proteins were purified by Ni²⁺-affinity chromatography, incubated with commercially available purified 20S proteasome in the absence of ATP, and submitted to Western blot analysis. Figure 4A shows that the 20S degradation assay for dTIS11_{V5/HIS} and dTIS11^{KR}_{V5/HIS} resulted in thorough degradation of these proteins, while all negative controls, including ubiquitinated Ub^{G76V}-GFP_{V5/HIS}, remained unaffected. Coincubation with clasto-lactacystin-beta-lactone reverted dTIS11_{V5/HIS} degradation, confirming the dependence on proteasomal activity in this assay. These results showed that dTIS11 can be directly degraded by the 20S proteasome, in the absence of the ATP necessary for the unfolding processes. To test whether dTIS11 was also sensitive to degradation by the 20S proteasome when produced in a heterologous, acellular system, in vitro translation of dTIS11 mRNA was programmed in reticulocyte lysate and the translation products were subsequently ex-

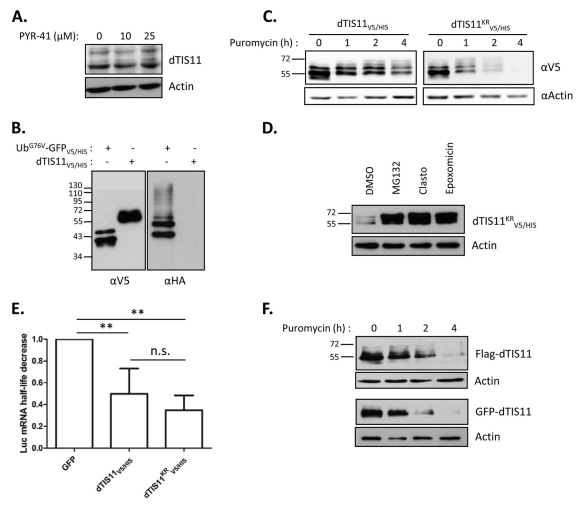


FIG 3 Ubiquitin-independent degradation of dTIS11 by the proteasome. (A) dTIS11 accumulation is insensitive to PYR-41 treatment. Cells were treated with PYR-41 at 10 μ M for 16 h or at 25 μ M for 2 h. Protein extracts were analyzed by Western blotting as described in the text. (B) Ubiquitination of dTIS11_{V5/HIS} was not detectable. Protein extracts from *Drosophila* S2 cells coexpressing HA-ubiquitin with either Ub^{G76V}GPF_{V5/HIS} or dTIS11_{V5/HIS} were subjected to IMAC. Western blotting of Ni²⁺ -NTA-retained proteins with anti-V5 and anti-HA antibodies revealed the presence of V5-tagged proteins and HA-ubiquitin in the His-enriched extract. (C and D) The lysine-less dTIS11^{KR}_{V5/HIS} mutant is unstable and degraded by the proteasome. S2 cells transfected with expression vectors encoding wild-type dTIS11_{V5/HIS} or lysine-less dTIS11^{KR}_{V5/HIS} were treated with puromycin for the indicated times (C) or with the proteasome inhibitors MG132, Clasto, epoxomicin, or dimethyl sulfoxide (DMSO) as a control for 4 h (D). Protein extracts were analyzed by Western blotting by using the indicated antibodies. (E) dTIS11^{KR}_{V5/HIS} was functional and destabilized ARE-bearing mRNA. S2 cells expressing an ARE-fused luciferase reporter with GFP_{V5/HIS}, dTIS11_{V5/HIS} or dTIS11^{KR}_{V5/HIS}, were treated with actinomycin D for up to 3 h. Luciferase mRNA levels were quantified by Northern blotting, and the associated half-lives were determined by exponential regression. The luciferase half-life decrease was calculated by reporting the half-life obtained under each condition and comparing it to that calculated for the GFP control. A Mann-Whitney U test was used for the statistical analysis. n.s., nonsignificant; *, P < 0.05; **, P < 0/01. (F) The context of the N-terminal amino acid does not control dTIS11 degradation. Flag-dTIS11-expressing cells were treated with puromycin for the indicated times. Protein extracts were analyzed by Western blotting using anti-FLAG, anti-FLAG, anti-GFP, or antiactin antibodies.

posed to purified 20S proteasome. Figure 4B shows efficient and 20S-dependent degradation of *in vitro*-translated dTIS11, while the negative-control GFP-DCP1 remained stable. Altogether, these results demonstrate that dTIS11 can be directly recognized and processed by the 20S proteasome in the absence of ATP and ubiquitin.

dTIS11 behaves as an intrinsically disordered protein. The ubiquitin-independent proteasome-dependent degradation, also known as "degradation by default" (55), is of increasing prevalence as the number of proteins described to undergo such degradation grows. Strikingly, many of these proteins present large disordered domains that lack a stable tertiary structure; hence, their recent classification as IDP (56). Moreover, protein disorder nor-

malized for protein length was shown as the major feature in determining a short half-life for proteins (57). As dTIS11 is involved in this "by default" pathway, we examined the possibility that dTIS11 protein would present one or more IDRs. We first performed bioinformatic analysis on the dTIS11 amino acid sequence, looking for predicted IDRs. Figure 5A shows the results returned by DISOPRED2, an algorithm trained on high-resolution X-ray crystal data to calculate a probability of disorder for each amino acid of the query protein. The predicted disorder for dTIS11 appeared very high throughout the whole sequence, with the notable exceptions of the tandem zinc finger domains (amino acids 135 to 163 and 173 to 201), as well as short stretches at the N and C termini. In contrast, the disorder profiles for GFP and *Dro*-

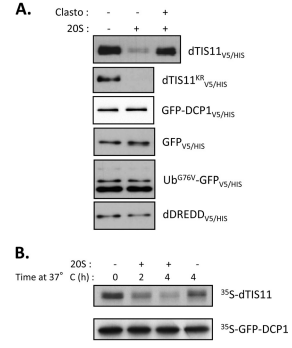


FIG 4 Direct recognition of dTIS11 by the 20S proteasome. (A) ATP-independent degradation of dTIS11 by purified 20S proteasome. The indicated proteins were purified using IMAC from S2 cells transfected with corresponding expression vectors and incubated for 1 h in the presence of 0.5 μ g of purified 20S proteasome with or without Clasto. Western blot analysis was performed with anti-V5 antibodies. (B) 20S proteasome-mediated degradation of dTIS11 produced by *in vitro* translation. dTIS11 and GFP-DCP1 were translated *in vitro* in the presence of [³⁵S]methionine and then incubated with 1 μ g 20S proteasome for the indicated times. Labeled proteins were revealed by SDS-PAGE followed by sodium salicylate gel staining and fluorography.

sophila DREDD were much flatter, which correlated with the inability of the 20S proteasome to process these proteins *in vitro* (Fig. 4A). Similar results were obtained when using Pondr-fit, IUpred, and DynaMine, three additional disorder prediction algorithms (data not shown).

Because dTIS11 is predicted to have extended disordered regions and because it complies with the main hallmark of IDPs, namely, susceptibility to degradation by the 20S proteasome, we next sought to determine whether dTIS11 also shared biochemical properties with IDPs. FastPP is a pulse proteolysis assay that measures a protein's degree of structuration in relation to temperature (41). This assay takes advantage of the specificity for unfolded states of the heat-resistant protease thermolysin (TL), which cleaves nearby exposed hydrophobic residues Phe, Val, Ile, and Leu. Incubation with TL thus results in degradation of the preferred disordered substrates, sparing folded proteins. In parallel, the incubation temperature can be increased to favor unfolding and thus susceptibility to TL. The amount of TL and the temperature needed for the TL to digest a protein give a measure of its structuration state. Figure 5B shows the results for FastPP analysis for dTIS11. Protein extracts from S2 cells untransfected (panels for dTIS11 and actin), or expressing dTIS11_{V5/HIS} or mRFP_{V5/HIS} as a highly structured control (Fig. 5A, right panel), were incubated with 1 or 10 mg/liter of TL at 4, 22, 50, or 80°C and then analyzed by Western blotting. Endogenous as well as overexpressed dTIS11 were both efficiently degraded, starting at 22°C.

Degradation of dTIS11 also required low TL concentrations (1 mg/liter) that proved insufficient for the degradation of the actin and mRFP_{V5/HIS} controls, even at the highest temperature. In fact, actin degradation required a 10-fold-higher concentration of TL and higher temperatures, of 50 to 60° C (Fig. 5B, bottom panel). As we would expect for IDPs, we found that dTIS11 was highly susceptible to this degradation process. Our results suggest that the dTIS11 unfolded state is already achieved at S2 cell culture temperatures of 24°C.

Another feature common in IDPs is their resistance to high temperatures. Because most IDPs have a low hydrophobic content, they do not present a hydrophobic core and are less likely to lose solubility upon heating (see reference 58 for a review). We thus tested dTIS11 heat resistance. Protein extracts from S2 cells expressing dTIS11_{V5/HIS}, mRFP_{V5/HIS}, or GFP-DCP1_{V5/HIS} were boiled at 95°C for 10 min and then ultracentrifuged at 100,000 × g to separate soluble from insoluble fractions. Western blot analysis showed that dTIS11_{V5/HIS} was mainly soluble, in contrast with the insoluble control proteins mRFP_{V5/HIS}, GFP-DCP1_{V5/HIS}, and endogenous GAPDH (Fig. 5C). Together, our data indicate that dTIS11, a protein predicted to be mainly disordered, also behaves like an IDP in several aspects.

As dTIS11 appears to be disordered and because disordered stretches have been shown to signal for proteasomal digestion, we next tested the capacity of regions of dTIS11 that were predicted to be disordered to change the fate of a stable protein. For this, we constructed expression vectors for GFP-DCP1-CdTIS11_{V5/HIS} and GFP-DCP1-NdTIS11_{V5/HIS}, built from the fusion of GFP-DCP1 with dTIS11 C-terminal region (CdTIS11; amino acids 214 to 436) and N-terminal region (NdTIS11; amino acids 1 to 110), respectively. The stability of these fusions was analyzed in S2 cells after translation inhibition. We observed that addition of CdTIS11 or NdTIS11 induced destabilization of the otherwisestable GFP-DCP1_{V5/HIS}, as shown by Western blotting using an anti-V5 antibody (Fig. 5D). Detection of the fusion protein by the N-terminal GFP moiety revealed that the degradation was complete, thus establishing dTIS11 C- and N-terminal domains as functional signals for degradation. While both domains are sufficient for destabilization, neither is necessary for dTIS11 turnover, as versions of dTIS11_{V5/HIS} lacking the N terminus (dTIS11- $\Delta \text{Nter}_{\text{V5/HIS}}$; amino acids 111 to 436) or most of the C-terminal domain (dTIS11- Δ Cter_{V5/HIS}, amino acids 1 to 253) were efficiently degraded by the 20S proteasome (Fig. 5E).

dTIS11 ubiquitin-independent proteasome-dependent degradation is conserved from Drosophila to mammals. The TIS11 family of proteins is present in eukaryotes from yeast to mammals. However, while the identification of proteins in this family is based on homology of their highly conserved tandem zinc finger domains, conservation for the rest of their sequence is much less conserved. This is easily observed when looking at a sequence alignment between Drosophila dTIS11 and the three mouse homologs, TTP, BRF1, and BRF2 (Fig. 6A). Yet, the implication of the proteasome in the degradation of TIS11/TTP proteins has been broadly established (20, 21). We thus addressed the hypothesis that the mammalian TIS11/TTP proteins would undergo the same degradation process as dTIS11. First, we analyzed the three mouse sequences by using DISOPRED2. We observed that despite sharing low sequence homology, their disorder profiles showed striking similarity with dTIS11, with an overall highly disordered degree, aside from the zinc finger domains and short stretches at

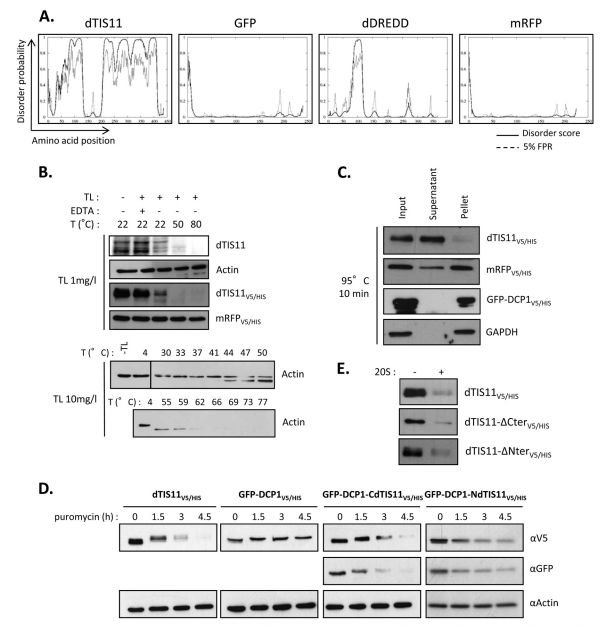


FIG 5 dTIS11 is a potential intrinsically disordered protein. (A) dTIS11 is predicted to be highly disordered. Disorder profiles were calculated using DISOPRED2 and are shown for the indicated proteins. Plain lines, disorder score per residue; dashed lines, prediction threshold at a 5% false-positive rate. (B) dTIS11 is a preferred substrate for proteolysis by thermolysin. Whole-protein extracts from untransfected S2 cells (dTIS11 and actin panels) or dTIS11_{V5/HIS}- or mRFP_{V5/HIS}-expressing cells were incubated with 1 or 10 mg/liter TL for 1 min at the indicated temperatures. Western blot analysis was performed with anti-dTIS11, antiactin, or anti-V5 antibodies. Vertical bars indicate cuts between separated parts of the same blot. (C) Heat resistance of dTIS11. Whole-protein extracts from dTIS11_{V5/HIS}-, mRFP_{V5/HIS}-, or GFP-DCP1_{V5/HIS}-expressing cells were boiled at 95°C for 10 min and then centrifuged at 100,000 × g for 10 min. Western blot analysis was performed on samples before centrifugation (input), and the soluble fraction was analyzed after centrifugation (supernatant) or the resuspended insoluble fraction (pellet). (D) dTIS11 N- and C-terminal domains act as signals for degradation. S2 cells expressing dTIS11_{V5/HIS}, GFP-DCP1_{V5/HIS}, GFP-DCP1-CdTIS11_{V5/HIS}, or GFP-DCP1-NdTIS11_{V5/HIS} were treated with puromycin for the indicated times. Western blot analysis was performed using the indicated antibodies. (E) The dTIS11 N terminus and C terminus are dispensable for 20S recognition. The indicated proteins were purified using tMAC from S2 cells transfected with are spondies. (E) The dTIS11 N terminus and C terminus are dispensable for 20S proteasome. Western blot analysis was performed with anti-V5 antibodies.

their N and C termini (Fig. 6B). We next sought to test our hypothesis experimentally on TTP, the founding member of the family. The corresponding Zfp36 gene is induced upon immunological challenge in macrophages and is then rapidly downregulated. We thus looked at the stability of the TTP protein in LPS-activated mouse macrophage cells, RAW 264.7. Translation

inhibition experiments showed that the TTP half-life is very short and is comparable to that of dTIS11 ($t_{1/2} = 114 \pm 35$ min). In addition, proteasome inhibition by clasto-lactacystin-beta-lactone prevented TTP degradation (Fig. 6C), implicating, as expected, the proteasome in this process. We also verified that TTP degradation did not require lysosomal activity in RAW 264.7 cells

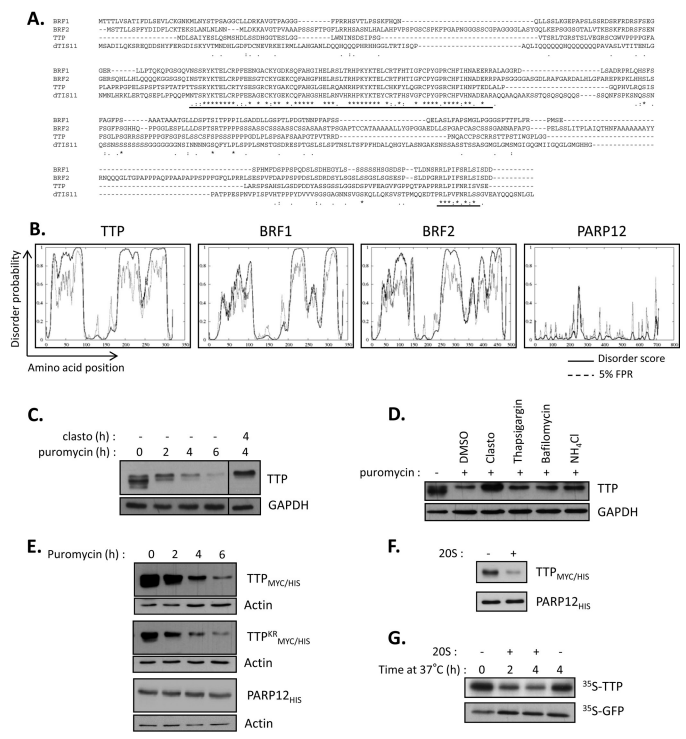


FIG 6 Evolutionary conservation of ubiquitin-independent proteasome-dependent degradation for TIS11/TTP proteins. (A) Sequence alignment based on ClustalW analysis of *Drosophila* dTIS11 and three mouse TIS11/TTP family proteins. Single letters, amino acids; asterisk, identical amino acid; colon, conserved substitution; period, semiconserved substitution. Highly conserved tandem zinc finger domains and the C-terminal stretch are underlined. (B) Disorder prediction calculated by using DISOPRED2 for the indicated proteins. Plain lines, disorder score per residue; dashed lines, prediction threshold at a 5% false-positive rate. (C) TTP is unstable and degraded by the proteasome. RAW 264.7 cells were stimulated with lipopolysaccharide for 4 h and treated with Clasto and puromycin for the indicated times. TTP was revealed by Western blotting with an anti-TTP antibody. Vertical bars indicate cuts between separated parts of the same blot. (D) Proteasome inhibition leads to TTP accumulation. RAW 264.7 cells were left untreated or cortreated with puromycin and the indicated times. TTP MYC/HIS and TTP^{KR}_{MYC/HIS} and analysis by Western blotting. (E) Ubiquitination is not required for TTP degradation. HEK293T cells were transfected with expression vectors for wild-type TTP_{MYC/HIS}, lysine-less mutant TTP^{KR}_{MYC/HIS}, or PARP12_{HIS} and treated with puromycin for the indicated times. TTP_{MYC/HIS} and TTP^{KR}_{MYC/HIS} by purified 20S proteasome. TTP_{MYC/HIS} and PARP12_{HIS} were detected by using anti-MYC antibodies, and PARP12_{HIS} were purified using IMAC on HEK293T cells transfected with the corresponding expression vectors and incubated with 0.5 µg purified 20S proteasome for 45 min. (G) 20S proteasome for the indicated times. Labeled proteins were revealed by SDS-PAGE followed by sodium salicylate gel staining and fluorography.

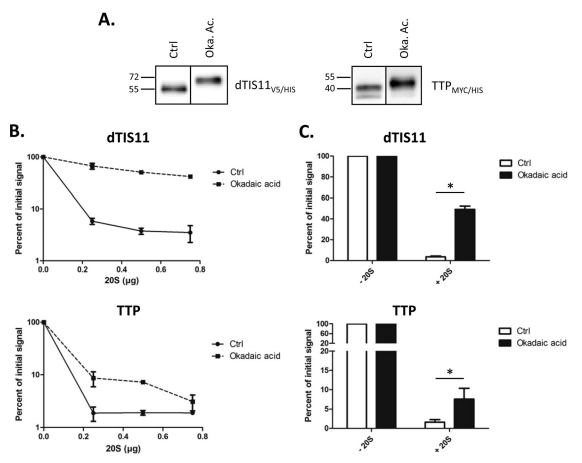


FIG 7 Phosphorylation regulates degradation of dTIS11 and TTP by the 20S proteasome *in vitro*. (A) Okadaic acid treatment induced phosphorylation of dTIS11 and TTP. S2 cells or HEK293T cells transfected with expression vectors for dTIS11_{V5/HIS} or TTP_{MYC/HIS}, respectively, were treated or not with okadaic acid (Oka. Ac.). His-tagged proteins were purified by IMAC and detected by Western blotting using anti-V5 or anti-MYC antibodies. Vertical bars indicate cuts between separated parts of the same blot. (B) Phosphorylation-regulated degradation of dTIS11 and TTP. Purified dTIS11_{V5/HIS} or TTP_{MYC/HIS} obtained as described for panel A were incubated for 1 h with the indicated amounts (B) or 0.5 μ g (C) of 20S proteasome and then detected by Western blotting. Two (B) and four (C) independent experiments were quantified using the ImageJ software; results are reported relative to the corresponding value in the absence of 20S proteasome and plotted on the graph. Error bars represent SD for each experimental condition. The Mann-Whitney U test was performed for statistical analysis. *, *P* < 0.05.

(Fig. 6D). To test whether ubiquitination is required for TTP proteasomal degradation, we compared the stability of wild-type TTP_{MYC/HIS} with its lysine-less mutant, TTP^{KR}_{MYC/HIS}. Of note, both proteins were doubly tagged with a Myc epitope for detection and 6×His repetition for purification. HEK293T cells were transfected with expression vectors for TTP_{MYC/HIS}, TTP^{KR}_{MYC/HIS}, or another zinc finger (ZC3H1) RNA-binding protein, $PARP12_{HIS}$, as a negative control. Translation inhibition experiments allow for the visualization of the degradation patterns. As observed for Drosophila dTIS11, conservative mutation of all 5 lysines into equally positively charged arginines did not prevent TTP efficient degradation, while the PARP12_{HIS} control remained stable (Fig. 6E). We next analyzed the ability of the 20S proteasome to directly recognize and degrade TTP_{MYC/HIS} purified from HEK293T extracts. Figure 6F shows efficient degradation of TTP_{MYC/HIS} when incubated with purified 20S proteasome, compared to the PARP12_{HIS} control. Interestingly, the disorder profile for PARP12 showed a much lower probability for destructuration, again correlating with the protein's insensitivity to 20S degradation (Fig. 6B and F). Finally, in vitro-translated TTP was degraded by purified 20S proteasome, indicating that TTP proteasomal degradation does not require ATP or ubiquitin (Fig. 6G).

Together, these results establish that as with *Drosophila* dTIS11, mammalian proteins of theTIS11/TTP family are targeted to the "degradation by default" pathway by ubiquitin-independent proteasome-dependent degradation.

Phosphorylation-dependent regulation of TTP and dTIS11 degradation by the 20S proteasome. In mammals, TIS11/TTP are substrates for phosphorylation events that result in stabilization of these factors (21). We tested whether phosphorylation affected TTP and dTIS11 degradation by the 20S proteasome. For this, S2 cells or HEK293T cells expressing dTIS11_{V5/HIS} and TTP_{MYC/HIS}, respectively, were treated or not with okadaic acid, a phosphatase inhibitor able to induce phosphorylation of TIS11/TTP proteins in mammalian cells (21, 59). Protein extracts obtained from these cells were then submitted to immobilized-metal affinity chromatography (IMAC) for HIS tag-based protein purification. Figure 7A confirms that okadaic acid treatment induced a shift toward higher apparent molecular weight bands, corresponding to phosphorylated forms for $dTIS11_{V5/2}$ HIS and TTP_{MYC/HIS}. Subsequent incubation with increasing amounts of 20S proteasome led to a significant decrease in the degradation efficiency for proteins obtained from okadaic acid-treated cells (Fig. 7B and C). These results suggest that phosphorylation prevents, at least in part, the degradation of both dTIS11 and TTP by the proteasome.

Notably, the stabilizing effect observed for TTP is much smaller than that seen for dTIS11. This could reveal intrinsic differences within regulatory aspects for the two proteins, or it could be due to the experimental setting used. Indeed, okadaic acid treatment may not affect dTIS11 and TTP phosphorylation to the same extent. Moreover, the associated hyperphosphorylation state may never even be achieved under normal conditions. Additionally, the proteasome concentration and the molecular environment in this assay do not reflect the physiological conditions of the cell, which could stress the differences observed between the two proteins. Nevertheless, our results clearly establish a phosphorylation-dependent effect on TIS11/TTP degradability by the 20S proteasome *in vitro*.

DISCUSSION

In order to maintain proper regulation of genetic expression, cells need to control the availability of main regulators, such as ARE-BPs. This can be achieved by adjusting the balance between protein production and degradation. In this study, we showed that, similar to other ARE-BPs, TIS11/TTP proteins possess high turnover rates, which ensure the setting of rapid transitions in protein levels. However, while ARE-BPs such as AUF1 or HuR are targeted to degradation by ubiquitin-dependent mechanisms (16, 17), we showed that TIS11/TTP proteins enter a ubiquitin-independent degradation-by-default pathway. The Drosophila dTIS11 protein and the mammalian TTP protein are both targeted for degradation by the same mechanism. Therefore, our results uncover a new evolutionarily conserved regulation mechanism for TIS11/TTP proteins. Drosophila dTIS11, as the sole member of the family in this organism, might be essential for regulation of a potentially broad spectrum of mRNAs, rather establishing dTIS11 as a general buffering protein that limits the expression of its targets. Evolutionary complexification by gene duplication may have then required fine-tuned and target-specific controls of the TIS11/TTP members by adding layers of regulation on each family member, such as implementing transcriptional and signaling-dependent regulation. Nevertheless, our results indicate that the mechanism controlling TIS11/TTP protein stability has been conserved across evolution from invertebrates to mammals.

Previous studies pointed to the proteasome as the main proteolytic complex responsible for TIS11/TTP proteins. However, this conclusion was most often drawn from pharmacological studies using drugs such as MG132 or lactacystin, which also possess inhibitory effects on lysosome activity (reviewed in reference 49). Thus, the potential implication of autophagy-driven lysosomal degradation, another pathway of cytosolic protein degradation (60), in TIS11/TTP turnover has not been clearly assessed. Here, we showed that in contrast to proteasomal inhibition, disturbance of lysosomal degradation by drugs targeting different steps of this process did not affect dTIS11 and TTP levels. In addition, a lysosome-deficient rabbit reticulocyte lysate programmed with either *dtis11* mRNA or *TTP* mRNA produced proteasome-sensitive dTIS11 and TTP, thus establishing these proteins as bona fide substrates of the proteasome.

Although proteasome dependent, we showed that TIS11/TTP degradation does not require ubiquitination. Rather, our results suggest that the destructuration states of these proteins are essential for proteasomal recognition. Intrinsically disordered regions

can be found as localized stretches in the primary sequence, linkers separating two folded domains, or even as extended parts of IDPs. Some of these regions have also been shown to serve as recognition domains for the 20S proteasome, interacting with the gate of the 20S and clearing the entrance to the degradation chamber (56). In addition, IDPs theoretically do not require active unfolding processes, as they do not present a stable 3-dimensional structure, thus allowing their entry in the degradation-by-default process.

While lacking tertiary structure under physiological conditions, many IDPs undergo induced folding upon binding to a partner. This process is thought to allow increased binding kinetics and structural plasticity (reviewed in reference 61). Furthermore, protein stabilization following binding/folding events has been described for several proteasome substrates (62-64). The overall 3-dimensional structure of TIS11/TTP proteins has not been elucidated, with the exception of two conserved regions, both predicted as folded in our study. First, the high-resolution crystal structure of the TTP C-terminal short stretch was crucial in understanding its function for the recruitment of the CCR4-NOT deadenylation complex (65). Second, nuclear magnetic resonance analysis of TTP and BRF1 tandem zinc finger domains showed that while folding of these regions depends on the availability of bivalent cations, their overall structuration is modified in the presence of target RNA sequences (66). In addition, complete structuration of the TTP second zinc finger requires proper binding to a target ARE (67), thus raising the possibility that target RNA availability controls the stability of TIS11/TTP proteins, as this mechanism has been described for the yeast RNA-BP YB-1 (63). Alternatively, other binding partners could prevent TIS11/TTP degradation by precluding the access of the proteasome to their disordered regions.

In mammals, the TIS11/TPP proteins are subject to extensive phosphorylation (21, 31, 68). A dephosphorylation assay revealed that the Drosophila dTIS11 is also heavily phosphorylated, which we then confirmed by mass spectrometry analysis (data not shown). Previous studies have shown that the degradation of TIS11/TTP factors is controlled by their phosphorylation status. In fact, several authors have proposed a model in which binding of phosphorylated forms of TIS11/TTP to the 14-3-3 adaptor protein would prevent their degradation (20, 21, 59). This model is mainly based on the fact that phosphorylation induces both TIS11/TTP stabilization and binding to 14-3-3. However, the causality link between the two events is not clear. In fact, addition of recombinant 14-3-3 in our in vitro degradation assay did not stabilize dTIS11 or TTP (data not shown). Our observations, though limited to our in vitro analysis, thus raise an alternate hypothesis in which phosphorylation would directly affect TIS11/TTP recognition by the proteasome, as such a process has been observed for at least one other protein (69). 14-3-3 adaptors are able to prevent TIS11/TTP dephosphorylation (70). Their stabilizing role could thus be strictly indirect with regard to proteasomal digestion.

The disordered state here associated with a degradation process is also known to provide additional properties to IDPs. Intrinsic disorder allows for an increased interaction surface and better exposure of interaction motifs, facilitating associations with several partners. This could explain the rather high number of interacting partners formally identified for TTP, including 14-3-3 proteins, the kinases p38, MK2, and MEKK4, and also members of the deadenylation machinery (reference 8 and references therein). Two-hybrid analysis also identified 31 potential interactions partners for TTP, 4 of which were later confirmed by coimmunoprecipitation experiments (71). Flexibility and adaptability are other crucial features of disordered regions. These would allow RNA-BPs to bind similar yet different RNA molecules as well as increase their "capture radius." In addition, the induced folding upon binding, common in IDPs, provides low-affinity/high-specificity binding (72), which is ideal for RNA-BP to target RNA recognition. These considerations may explain the observed enrichment in IDRs for RNA-BPs (73, 74). In accordance with this hypothesis, the disordered linker present in the tandem zinc finger domains of the BRF1 protein was shown to specifically contact AU-rich elements of mRNAs (66). This binding is controlled by an induced folding mechanism in which the tandem zinc finger becomes more rigid and stable upon mRNA contact (75). This mode of RNA recognition is correlated with a large diversity of mRNA targets identified for BRF1 (76).

RNA-BPs represent one of the largest protein families in eukaryotes (77). Although the average stability of RNA-BPs is higher than that of non-RNA-BPs (78), it remains to be determined whether the IDRs found to be enriched in RNA-BPs could serve as functional signals for proteasomal degradation for specific subsets of RNA-BPs.

ACKNOWLEDGMENTS

We thank Vincent Raussens for helpful discussions and William Rigby for providing CARP-3 anti-TTP antibody.

Work in our laboratory is supported by the Fund for Medical Scientific Research (Belgium; grant 2.4556.08), the Fonds Brachet, the Fonds Van Buuren, the Actions de Recherches Concertées (grants AV.06/11-345 and AV.12/17), and the FEDER Hainaut Biomed program. L. V. Ngoc is supported by a doctoral grant of the Fonds pour la Recherche en Industrie et Agriculture (Belgium).

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