

# Long non-coding RNA *gadd7* interacts with TDP-43 and regulates *Cdk6* mRNA decay

Xuefeng Liu<sup>1,2</sup>, Dan Li<sup>1</sup>, Weimin Zhang<sup>1</sup>,  
Mingzhou Guo<sup>2</sup> and Qimin Zhan<sup>1,\*</sup>

<sup>1</sup>State Key Laboratory of Molecular Oncology, Cancer Institute and Hospital, Chinese Academy of Medical Sciences and Peking Union Medical College, Beijing, China and <sup>2</sup>Department of Gastroenterology and Hepatology, Chinese PLA General Hospital, Beijing, China

Long non-coding RNAs (lncRNAs) transcribed extensively from the genome have been proposed to be key regulators of diverse biological processes. However, little is known about the role of lncRNAs in regulation of the cell-cycle G1/S checkpoint following DNA damage, a key step in the maintenance of genomic fidelity. Here we show that growth-arrested DNA damage-inducible gene 7 (*gadd7*), a DNA damage-inducible lncRNA, regulates the G1/S checkpoint in response to UV irradiation. Interestingly, UV-induced *gadd7* directly binds to TAR DNA-binding protein (TDP-43) and interferes with the interaction between TDP-43 and cyclin-dependent kinase 6 (*Cdk6*) mRNA, resulting in *Cdk6* mRNA degradation. These findings demonstrate a role for *gadd7* in controlling cell-cycle progression and define a novel mechanism by which lncRNAs modulate mRNA expression at the post-transcriptional level by altering mRNA stability.

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## Introduction

Mammalian genomes transcribe a larger number of non-coding RNAs than previously expected (Mattick, 2009). In addition to small regulatory RNAs (e.g., microRNAs, small interfering RNAs and others), surprisingly numerous long non-coding RNAs (lncRNAs) have been identified (Mattick and Makunin, 2006; Guttman *et al.*, 2009). lncRNAs are generally considered as non-protein-coding transcripts varying in length from several hundred bases to tens of kb. Similar to protein-coding genes, lncRNAs have been implicated in diverse biological processes, including invasiveness and metastasis (Gupta *et al.*, 2010), apoptosis (Huarte *et al.*, 2010), dosage compensation (Deng and Meller, 2006) and embryogenesis (Pauli *et al.*, 2011). Furthermore, lncRNAs have been reported to regulate gene expression

\*Corresponding author. State Key Laboratory of Molecular Oncology, Cancer Institute and Hospital, Chinese Academy of Medical Sciences and Peking Union Medical College, Beijing 100021, PR China.  
Tel.: +86 10 67762694; Fax: +86 10 67715058;  
E-mail: zhanqimin@pumc.edu.cn

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through a wide range of mechanisms (Mercer *et al.*, 2009). For example, lncRNAs can serve as a repressor or an activator to regulate the process of transcription. *LincRNA-p21*, a lncRNA downstream of p53, acts as a repressor in p53-dependent transcriptional responses by physically associating with heterogeneous nuclear ribonucleoprotein (hnRNP) K and modulating its localization (Huarte *et al.*, 2010). In addition, the ability of lncRNAs to recognize complementary sequences also allows lncRNAs to regulate various steps in the post-transcriptional processing of mRNAs, as in the case of the *Zeb2* antisense RNA, which complements an intron in the 5'-untranslated region (5'UTR) of *Zeb2* and inhibits its splicing (Beltran *et al.*, 2008). Recently, a handful of studies have implicated lncRNAs in a variety of disease states, especially in neurodegenerative diseases and cancers (Huarte and Rinn, 2010; Qureshi *et al.*, 2010; Wapinski and Chang, 2011). Although lncRNAs have been recognized as a critical functional expression in complex organisms, there are still huge gaps in our understanding of lncRNAs.

*gadd7* (growth-arrested DNA damage-inducible gene 7) is a 754-nt polyadenylated lncRNA that was isolated from Chinese hamster ovary (CHO) cells on the basis of increased expression following UV irradiation (Fornace *et al.*, 1988; Hollander *et al.*, 1996). Induction of *gadd7* is also observed after several other types of DNA damage and growth arrest signals (Fornace *et al.*, 1988; Hollander *et al.*, 1996). In addition, overexpression of *gadd7* leads to suppression of cell growth (Hollander *et al.*, 1996), suggesting that *gadd7* may be involved in cell-cycle control. More recently, it has been reported that *gadd7* is also a regulator of lipid-induced oxidative and endoplasmic reticulum stress (Brookheart *et al.*, 2009).

Here we further demonstrate that *gadd7* is induced by both growth arrest and DNA damage signals, and plays a key role in regulation of the G1/S checkpoint after DNA damage. Moreover, we find that *gadd7* regulates *Cdk6* mRNA decay by dissociating TAR DNA-binding protein (TDP-43) from cyclin-dependent kinase 6 (*Cdk6*) mRNA upon UV irradiation. Collectively, this study describes *gadd7* as a cell-cycle regulator, and explores the underlying mechanism by which *gadd7* regulates the cell-cycle checkpoint in response to DNA damage, which provides novel insights into the understanding of the biological functions of lncRNAs.

## Results

### *gadd7* is induced in response to genotoxic and non-genotoxic stresses

It has been reported that *gadd7* is a DNA damage-inducible transcript (Fornace *et al.*, 1988; Hollander *et al.*, 1996). We here further investigated the induction of *gadd7* by DNA damage and growth arrest signals. The results demonstrated that expression of *gadd7* RNA was substantially induced in a dose-dependent manner following CHO-K1 cells' exposure to UV irradiation (Figure 1A; Supplementary Figure S1A).

Furthermore, cisplatin, a chemotherapeutic drug that causes DNA damage, also induced *gadd7* expression in a dose-dependent manner (Figure 1B; Supplementary Figure S1B). Similar to the induction by DNA damage, increased *gadd7* expression was seen following treatment of CHO-K1 cells with growth arrest signals such as media depletion (Figure 1C; Supplementary Figure S1C) and serum reduction (Figure 1D; Supplementary Figure S1D). In these assays, *Gadd45a*, a well-known DNA damage-inducible transcript (Rosemary Siafakas and Richardson, 2009), was included as positive control. These results further confirm that *gadd7* is induced upon various types of genotoxic and non-genotoxic stresses that elicit growth arrest.

### ***gadd7* plays a role in regulation of the G1/S checkpoint following DNA damage**

Growth arrest is a well-known and critical cellular event after DNA damage. We aimed to determine the physiological role of *gadd7* in this process. First, the approach of RNA interference (RNAi) was employed to knock down endogenous *gadd7* in CHO-K1 cells, and significant decrease of *gadd7* expression was achieved (Figure 2A). We found that depletion of *gadd7* resulted in increased cell proliferation compared with cells treated with non-specific siRNA control as measured by growth curves and MTT assay (Figure 2B and C), suggesting that *gadd7* plays a negative role in the control of cell growth.

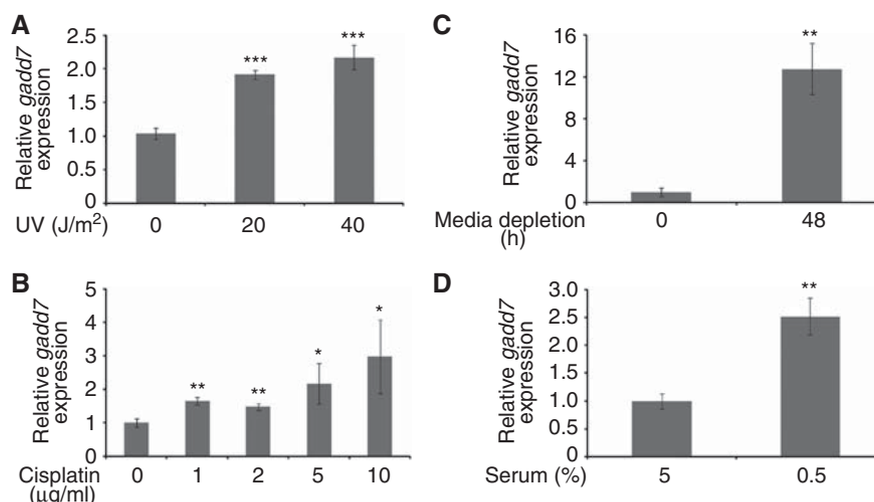
Next we examined whether *gadd7* regulates cell-cycle progression. We observed that knockdown of *gadd7* led to the changes in cell-cycle distribution. The cell population in G1 phase was reduced, but that in S phase was increased after depletion of *gadd7* (Figure 2D, top; Supplementary Figure S2A), suggesting that *gadd7* may affect G1/S transition. To better understand the *gadd7*-regulatory function in G1/S transition, cell-cycle distribution analyses were further conducted in the presence of nocodazole, which blocks cells in mitosis (Zieve *et al*, 1980). After treatment with nocodazole

for 6 h, a remarkable reduction in the G1 population and an accumulation of G2/M phase were observed in *gadd7*-depleted cells compared with siRNA control (Figure 2D, middle; Supplementary Figure S2A). At 12 h, nearly all *gadd7*-knockdown cells progressed through G1 phase and entered S and G2/M phase, while many of the cells treated with control siRNA were still in G1 phase (Figure 2D, bottom; Supplementary Figure S2A). Thus, *gadd7* might be essential for the control of an orderly G1/S transition.

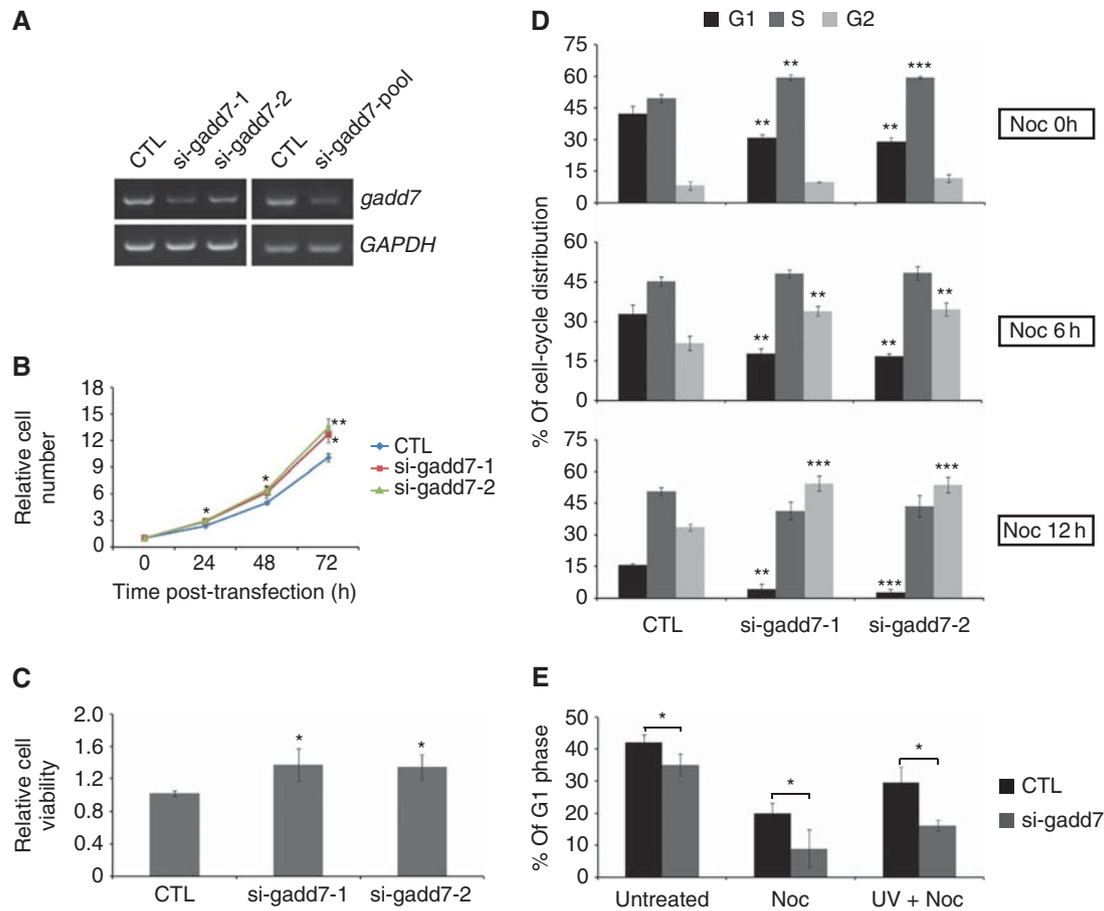
Since *gadd7* is induced by various types of DNA damage signals, we examined the effect of *gadd7* on the G1/S checkpoint in response to DNA damage. In the control cells, UV irradiation resulted in a large accumulation of the G1 population in the presence of nocodazole treatment, indicating that there is an intact G1/S checkpoint after DNA damage. In contrast, the accumulation of G1 phase was substantially reduced in cells silenced for *gadd7* after UV irradiation (Figure 2E; Supplementary Figure S2B). These observations demonstrate that *gadd7* is a key regulator of the G1/S checkpoint after UV irradiation. In addition, we also investigated the effect of *gadd7* overexpression on G1 arrest in the absence of UV treatment. However, overexpression of *gadd7* alone appeared not to generate a G1 arrest (Supplementary Figure S3A and B), suggesting that other factors induced by UV irradiation may act in concert with *gadd7* to regulate the G1/S checkpoint.

### ***gadd7* specifically binds to TDP-43**

Recent studies have shown that lncRNAs perform their functions through interaction with proteins (Rinn *et al*, 2007; Huarte *et al*, 2010). Therefore, we assumed that the role of *gadd7* in cell-cycle control might be possibly mediated by such a mechanism. To identify the proteins that are associated with *gadd7*, biotin RNA pull-down assay was performed. The biotinylated *gadd7* and the two negative control transcripts, biotinylated antisense-*gadd7* and biotinylated *GAPDH*, were transcribed *in vitro* (Supplementary Figure S4A) and incubated



**Figure 1** Induction of *gadd7* following DNA damage and growth arrest signals in CHO-K1 cells. (A, B) Induction of *gadd7* expression upon DNA damage. Exponentially growing CHO-K1 cells were treated with different doses of UV (A) or cisplatin (B). The expression of *gadd7* was detected by qRT-PCR and normalized to glyceraldehyde-3-phosphate dehydrogenase (*GAPDH*). Data are from one of three independent experiments and are represented as mean  $\pm$  s.d. with  $n = 3$ . \* $P < 0.05$ , \*\* $P < 0.01$  or \*\*\* $P < 0.001$  compared with the untreated groups. (C, D) Induction of *gadd7* expression upon growth arrest signals. CHO-K1 cells were treated with the two growth arrest signals, media depletion (C) and serum reduction (D). The expression of *gadd7* was detected and analysed as in (A, B). \*\* $P < 0.01$  compared with the untreated groups.



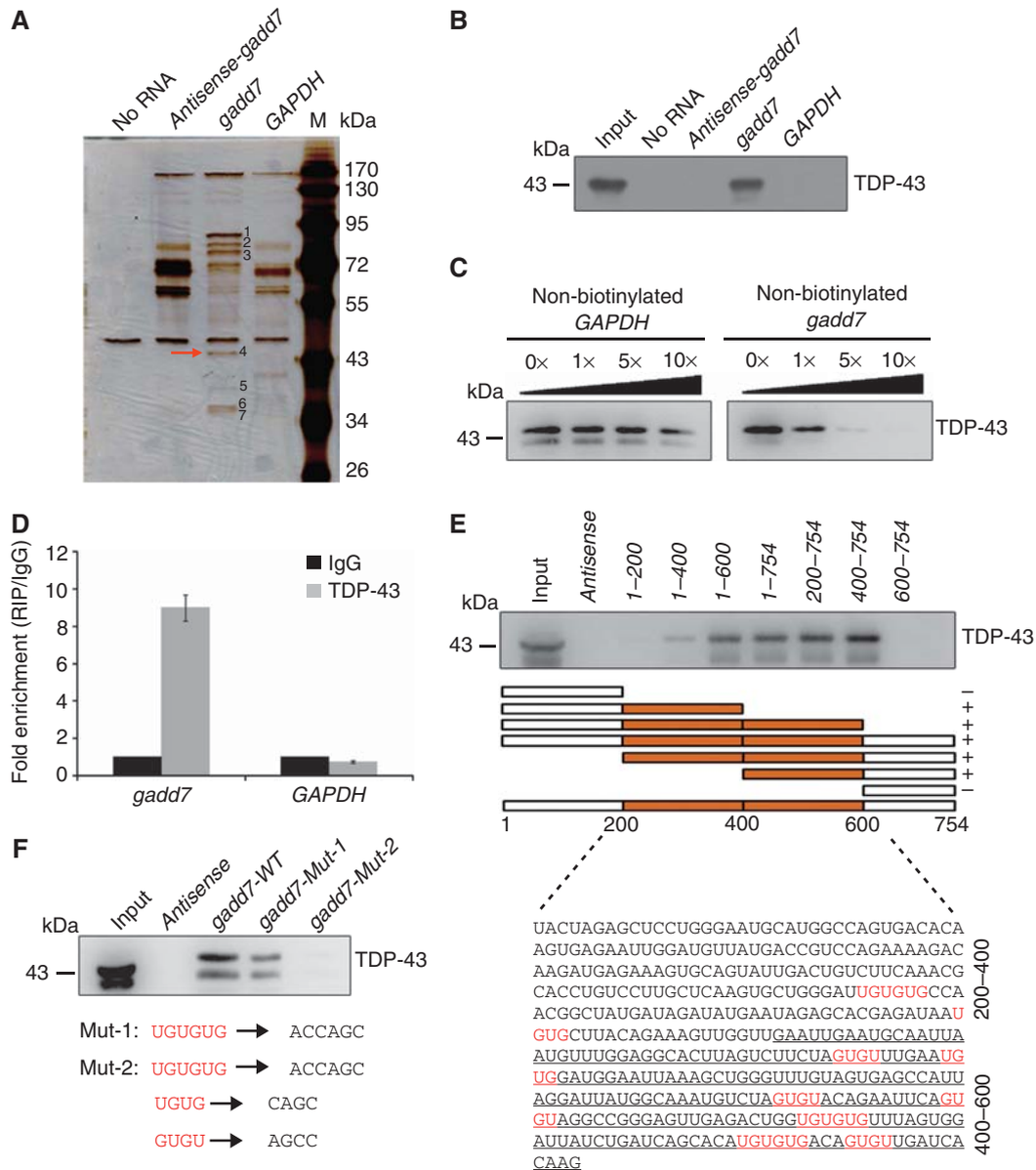
**Figure 2** Defect in the G1/S checkpoint in *gadd7*-depleted cells. (A) RNAi-mediated knockdown of *gadd7* in CHO-K1 cells. CHO-K1 cells were transfected with control siRNA (CTL) or two individual siRNAs targeting *gadd7* separately or combined. The expression of *gadd7* was detected by semi-quantitative RT-PCR after transfection for 48 h. *GAPDH* was used as internal reference. (B, C) Knockdown of *gadd7* increases cell growth. The relative numbers of CHO-K1 cells transfected with control siRNA or two individual siRNAs against *gadd7* were counted daily for 3 days (B), and the relative cell viability was determined by MTT assay after transfection for 72 h (C). Data are represented as mean  $\pm$  s.d. from three independent experiments. \* $P < 0.05$  or \*\* $P < 0.01$  compared with control siRNA. (D) Depletion of *gadd7* causes a rapid G1/S transition. CHO-K1 cells transfected with the indicated siRNAs for 48 h were treated with nocodazole (Noc) for 0 h (top panel), 6 h (middle panel) or 12 h (bottom panel). Cell-cycle distribution was measured by PI staining followed by flow cytometry. The percentage of cells in G1, S or G2 phase transfected with control siRNA is defined as control. Data are represented as mean  $\pm$  s.d. from three independent experiments. \*\* $P < 0.01$ , \*\*\* $P < 0.001$ . (E) *gadd7* knockdown results in G1/S checkpoint defect following UV irradiation. CHO-K1 cells transfected with control siRNA or *gadd7* siRNA pool for 48 h were treated with nocodazole alone or in combination with UV irradiation just prior to the addition of nocodazole. After 12 h, the distribution of cell cycle was analysed as in (D). Data are represented as mean  $\pm$  s.d. from three independent experiments. \* $P < 0.05$ .

with whole CHO-K1 cell lysates. Biotin-labelled transcripts and their associated cellular proteins were captured by streptavidin beads and then subjected to sodium dodecyl sulphate-polyacrylamide gel electrophoresis (SDS-PAGE) analysis. The results of biotin RNA pull-down assay are presented in Figure 3A. Compared with antisense-*gadd7* (line 2) and *GAPDH* (line 4) RNAs, there were seven bands that were specifically associated with *gadd7* (line 3). The protein bands specific to *gadd7* were then extracted, digested with trypsin and subjected to mass spectrometry analysis. Interestingly, this analysis showed that the majority of these proteins binding to the *gadd7* transcript were hnRNPs (Supplementary Table S1). To focus on the role of *gadd7* in the control of cell cycle, TAR DNA-binding protein (TDP-43) (Figure 3A, red arrow) was chosen for further studies given its function in regulating cell-cycle distribution (Ayala *et al*, 2008). First, the association of *gadd7* with TDP-43 was validated by immunoblotting assay and the results showed that TDP-43 was clearly detected in *gadd7*-pull-down protein

complexes, but not in the complexes associated with antisense-*gadd7* or *GAPDH* (Figure 3B). To further confirm the specificity of this interaction, competition assay was performed by adding different amounts of non-biotinylated *gadd7* or *GAPDH*. The interaction between *gadd7* and TDP-43 was competed by non-biotinylated *gadd7* in a dose-dependent manner, but not by *GAPDH* (Figure 3C).

To confirm the interaction between *gadd7* and TDP-43 *in vivo*, RNA immunoprecipitation (RIP) was performed. An antibody against TDP-43 was incubated with CHO-K1 cell lysates, and co-precipitated RNAs were then analysed by RT-polymerase chain reaction (RT-PCR) or qRT-PCR using primers for *gadd7* or *GAPDH* (negative control). As expected, we observed an enrichment of *gadd7*, but not *GAPDH*, with TDP-43 IP when compared with non-specific IgG IP (Figure 3D; Supplementary Figure S4B).

We then aimed to map the regions of *gadd7* that are required for the interaction. For this purpose, a series of *gadd7* deletion mutants, as displayed in Figure 3E, were



**Figure 3** *gadd7* directly interacts with TDP-43. (A) Identification of cellular proteins associated with *gadd7* *in vitro*. Proteins from CHO-K1 cell extracts were pulled down with the biotin-RNAs, subjected to SDS-PAGE and visualized by silver staining. The bands specific to *gadd7*, as pointed out by the numbers (bands 1–7), were subjected to mass spectrometry. The TDP-43 (band 4) is indicated by the red arrow. (B) Immunoblotting analysis of the specific interaction of *gadd7* with TDP-43. Proteins pulled down with the biotin-RNAs as in (A) were analysed by immunoblotting with TDP-43 antibody. (C) Specific association of *gadd7* with TDP-43 is confirmed by competition assay. Various amounts of unlabelled *gadd7* were added to compete with biotin-labelled *gadd7* in interacting with TDP-43 (right panel). Unlabelled *GAPDH* RNA was used as negative control (left panel). TDP-43 associated with the biotin-*gadd7* was analysed by immunoblotting. (D) *gadd7* interacts with TDP-43 *in vivo*. TDP-43 was immunoprecipitated from CHO-K1 cells and co-precipitated RNAs were detected by qRT-PCR using primers for *gadd7* and *GAPDH* (as negative control). IP enrichment is determined as the amount of RNA associated to TDP-43 IP relative to IgG control. Data are from one of three independent experiments and are represented as mean  $\pm$  s.d. with  $n = 3$ . (E) Two regions in the middle of *gadd7* are necessary to associate with TDP-43. A series of *gadd7* deletion mutants and *gadd7* antisense were used to perform RNA pull-down assay as in (A), and followed by immunoblotting with TDP-43 antibody. The regions bound by TDP-43 are indicated using the boxes shown in orange. UG/GU repeats in *gadd7* transcript are shown in red. The ranges of *gadd7* sequence are indicated on the right with 400–600 nt underlined. (+) binding; (–) unbinding. (F) The interaction of *gadd7* with TDP-43 is GU/UG-repeat dependent. Two point mutants of *gadd7* were treated as in (A) and associated TDP-43 was detected by immunoblotting. Mut-1 denotes three UGUGUG elements mutated to ACCAGC. Mut-2 denotes UGUG and GUGU repeats mutated to CAGC and AGCC, respectively, in addition to UGUGUG elements mutation. WT, wild type. Figure source data can be found with the Supplementary data.

synthesized by *in vitro* transcription and labelled with biotin (Supplementary Figure S4C). RNA pull-down assay was carried out and followed by immunoblotting assay to measure the interactions. We found that deletion mutants that contain the regions from nucleotides 200 to 400 or 400 to 600 were able to bind to TDP-43 (Figure 3E). Interestingly,

UG/GU (UG or GU) repeats, the canonical TDP-43 target sequences (Polymenidou *et al*, 2011; Tollervey *et al*, 2011), are present in the regions from nucleotides 200 to 400 and especially in 400–600, indicating that these two regions with UG/GU repeats might be necessary for *gadd7*'s association with TDP-43.

To confirm the requirement of these UG/GU repeats for binding to TDP-43, two point-mutant versions, namely *gadd7-Mut-1* and *gadd7-Mut-2*, in which three UGUGUG elements or all UG/GU repeats were mutated, respectively (Figure 3F), were transcribed *in vitro* for RNA pull-down experiments (Supplementary Figure S4D). The results showed that mutation of three UGUGUG elements in *gadd7* RNA substantially reduced its interaction with TDP-43, and additional mutation of UGUG and GUGU sites almost completely abolished such interaction (Figure 3F), indicating that UG/GU repeats are responsible for the interaction between *gadd7* and TDP-43. Collectively, all these observations indicate that *gadd7* directly interacts with TDP-43 through UG/GU repeats.

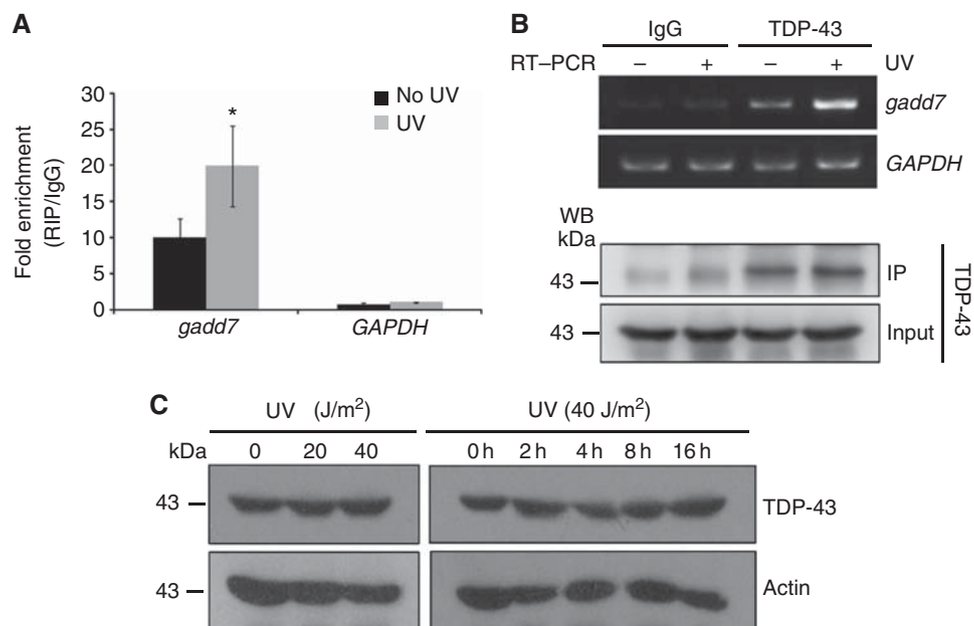
### The interaction between *gadd7* and TDP-43 is increased following UV irradiation

Based on the induction of *gadd7* following UV irradiation, we examined the interaction between *gadd7* and TDP-43 in response to UV irradiation. To this end, RIP with an antibody against TDP-43 was performed in CHO-K1 cells subjected to UV irradiation, followed by qRT-PCR or RT-PCR analysis. As expected, UV irradiation led to increased interaction between *gadd7* and TDP-43, while the interaction between *GAPDH* and TDP-43 showed no significant change (Figure 4A and B). To exclude the possibility that increased interaction between *gadd7* and TDP-43 may be the result of changed levels of TDP-43 upon UV irradiation, the expression of TDP-43 was examined. We observed that the expression of TDP-43 was unchanged in response to UV irradiation either in a dose-dependent manner or in a time course-dependent manner

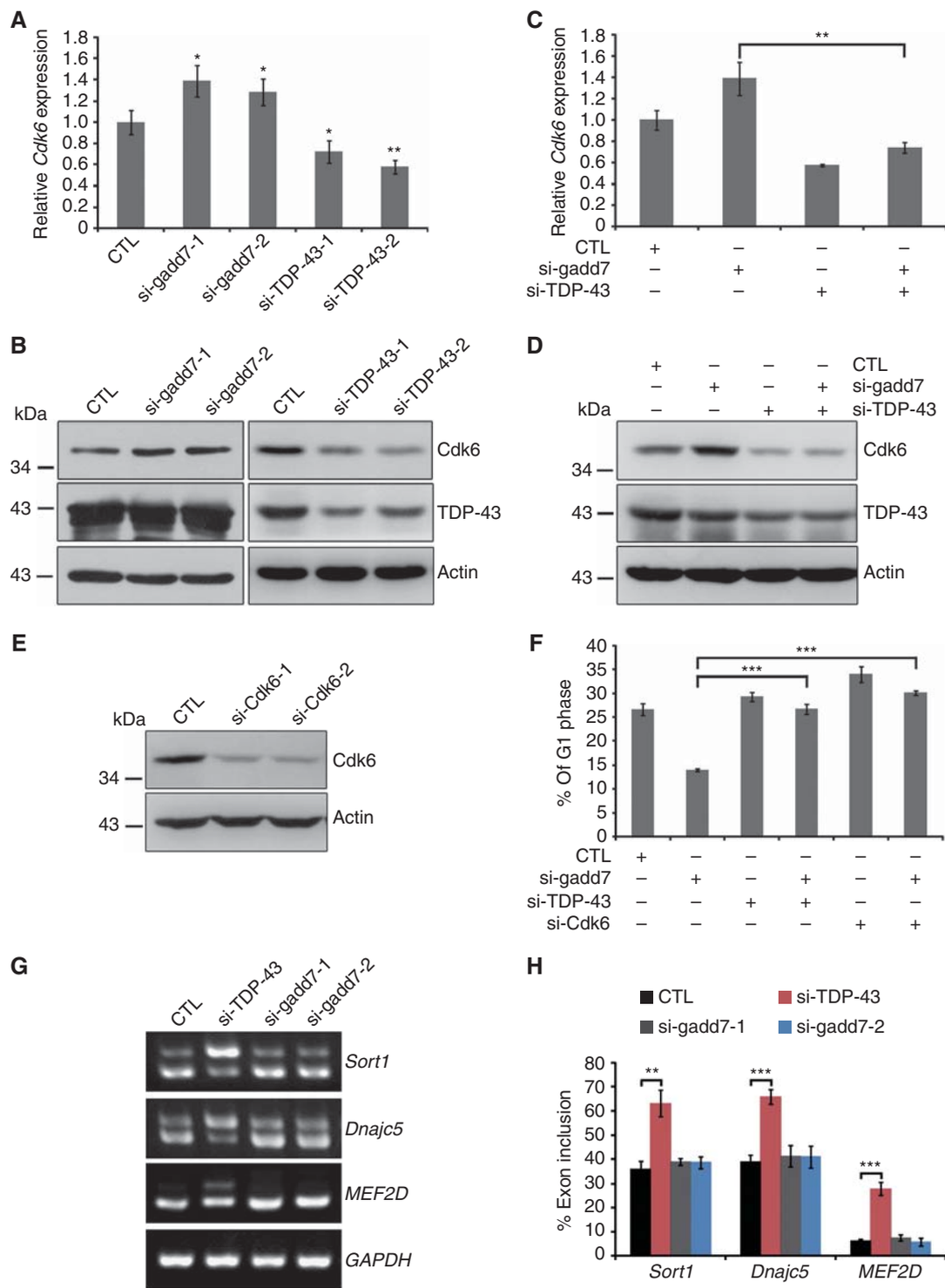
(Figure 4C). Together with the observation that *gadd7* expression is increased after UV irradiation, these findings suggest that upon UV irradiation *gadd7* is substantially induced and then binds to TDP-43.

### *gadd7* and TDP-43 co-regulate *Cdk6* mRNA expression

We next investigated the functional relevance of the interaction between *gadd7* and TDP-43. Recent studies by Ayala *et al* (2008) demonstrate that TDP-43 inhibits *Cdk6* expression through its recruitment to the UG-rich transcript of *Cdk6* in HeLa cells. *Cdk6* is known to associate with Cyclin D and regulate G1/S transition of the cell cycle (Lee and Yang, 2003). Thus, we speculated that *gadd7* might be involved in the regulation of *Cdk6* expression. Indeed, we observed upregulation of *Cdk6* transcript and protein levels upon knockdown of *gadd7* in CHO-K1 cells (Figure 5A and B). However, to our surprise, knockdown of TDP-43 in CHO-K1 cells significantly decreased the expression of *Cdk6* in both RNA and protein levels (Figure 5A and B), which was not consistent with the findings of Ayala *et al* (2008). To further confirm the effect of TDP-43 on *Cdk6* expression, we introduced the expression vector of human full-length TDP-43 into CHO-K1 cells and then examined the expression of *Cdk6*. Consistent with the observation that silencing of TDP-43 in CHO-K1 cells downregulates *Cdk6*, overexpression of TDP-43 resulted in upregulated expression of *Cdk6* (Supplementary Figure S5A and B). These findings indicate that TDP-43 might act as an activator rather than a repressor in the expression of *Cdk6* in CHO-K1 cells. Furthermore, simultaneous knockdown of *gadd7* and TDP-43 substantially attenuated the induction of *Cdk6* by *gadd7* depletion at both



**Figure 4** UV irradiation results in increased interaction between *gadd7* and TDP-43. **(A, B)** Exponentially growing CHO-K1 cells subjected to UV irradiation (40 J/m<sup>2</sup>) were cultured for additional 4 h and then harvested for RIP assay with TDP-43 antibody and IgG. RNAs present in the immunoprecipitates were detected by qRT-PCR **(A)** or semi-quantitative RT-PCR **(B, upper panel)** using primers for *gadd7* or *GAPDH* (as negative control). IP of TDP-43 was confirmed by immunoblotting using the cognate antibody **(B, lower panel)**. IP enrichment is determined as the amount of RNA associated to TDP-43 IP relative to the corresponding IgG control. Data are from one of two independent experiments and are represented as mean  $\pm$  s.d. with  $n = 3$ . \* $P < 0.05$  compared with the no UV-treated group. **(C)** The expression of TDP-43 is unchanged following UV irradiation. Exponentially growing CHO-K1 cells were treated with different doses of UV, and then harvested at the indicated times. The expression of TDP-43 and actin was detected by immunoblotting with the corresponding antibodies. Actin was used as loading control. Figure source data can be found with the Supplementary data.



**Figure 5** The expression of *Cdk6* is regulated by *gadd7* and TDP-43. (A, B) Effect of knockdown of *gadd7* or TDP-43 on *Cdk6* mRNA and protein levels. Control siRNA (CTL) or the indicated siRNAs were transfected into CHO-K1 cells. Forty-eight hours after transfection, *Cdk6* mRNA level (A) was detected by qRT-PCR and normalized to *GAPDH*. The protein level (B) was detected by immunoblotting and Actin was used as loading control. Data are from one of three independent experiments and are represented as mean  $\pm$  s.d. with  $n = 3$ . \* $P < 0.05$  or \*\* $P < 0.01$  compared with control siRNA. (C, D) Induction of *Cdk6* by *gadd7* depletion was attenuated when TDP-43 was simultaneously knocked down. CHO-K1 cells were transfected with control siRNA or an siRNA pool targeting *gadd7* or TDP-43 (either alone or in combination). After 48 h, *Cdk6* mRNA level (C) and protein level (D) were detected as in (A) and (B), respectively. Data are from one of three independent experiments and are represented as mean  $\pm$  s.d. with  $n = 3$ . \*\* $P < 0.01$ . (E) Knockdown of *Cdk6* in CHO-K1 by RNAi. *Cdk6* expression was examined by immunoblotting. (F) Co-transfection of *gadd7* siRNA with *Cdk6* or TDP-43 siRNA rescues the defective G1/S checkpoint. The G1/S checkpoint of CHO-K1 cells treated with siRNAs against *gadd7*, *Cdk6* or TDP-43, singly or together, was analysed as in Figure 2E. Data are represented as mean  $\pm$  s.d. from three independent experiments. \*\*\* $P < 0.001$ . (G) Effects of TDP-43 and *gadd7* on alternative splicing. The indicated siRNAs were co-transfected into CHO-K1 cells. Forty-eight hours after transfection, total RNA was extracted, and splicing patterns were examined by semi-quantitative RT-PCR. (H) Quantification of inclusion rates from three independent experiments by measuring the intensity of each band shown in (G). \*\* $P < 0.01$ , \*\*\* $P < 0.001$ . Figure source data can be found with the Supplementary data.

RNA and protein levels (Figure 5C and D). Given the direct interaction of *gadd7* with TDP-43 and loss of Cdk6 induction with simultaneous knockdown of *gadd7* and TDP-43, we conclude that *gadd7* downregulates the expression of Cdk6 mainly through its inhibition of TDP-43.

Since *gadd7* is involved in regulation of the G1/S checkpoint, we hypothesized that this effect might be mediated through its interaction with TDP-43 and inhibition of Cdk6. To this end, Cdk6 was knocked down by RNAi in CHO-K1 cells (Figure 5E), and the effect on the G1/S checkpoint of CHO-K1 cells upon co-transfection of *gadd7* siRNA with either TDP-43 or Cdk6 siRNA was analysed. Clearly, defect in the G1/S checkpoint in *gadd7*-depleted cells was rescued by TDP-43 or Cdk6 depletion (Figure 5F).

### ***gadd7* interferes with the interaction between TDP-43 and *Cdk6* mRNA**

TDP-43 plays an important role in RNA processing, especially in pre-mRNA splicing, as target sequences (UG-rich sequences) are preferentially localized in long intronic regions and near splice site acceptors (Polymenidou *et al*, 2011; Sephton *et al*, 2011; Tollervey *et al*, 2011). Hence, several selected targets of TDP-43, such as *MEF2D* (Tollervey *et al*, 2011), *Sort1* and *Dnajc5* (Polymenidou *et al*, 2011), were used to investigate whether *gadd7* is also involved in the splicing event. We found that knockdown of TDP-43 led to more inclusion of exons in *MEF2D*, *Sort1* and *Dnajc5* compared with control siRNA, while depletion of *gadd7* had no effect on alternative splicing of these genes (Figure 5G and H). Since the function of TDP-43 in CFTR exon 9 alternative splicing is well characterized (Buratti and Baralle, 2001; Buratti *et al*, 2001), we subsequently analysed the role of *gadd7* in this important event. CHO-K1 cells were co-transfected with TG13T5 minigene and siRNAs targeting TDP-43 or *gadd7*. Similar to what was observed in *MEF2D*, *Sort1* and *Dnajc5*, knockdown of TDP-43, but not *gadd7*, led to considerably increased inclusion of CFTR exon 9 (Supplementary Figure S6A–C). These data demonstrate that TDP-43 is indeed a powerful splicing modulator in CHO-K1 cells, but that *gadd7* has no role in regulating this process.

In addition to participation in pre-mRNA splicing, TDP-43 has also been implicated in mRNA decay because of its binding to target 3'UTRs (Colombrita *et al*, 2012). Several investigations have shown that TDP-43 stabilizes the human neurofilament (hNFL) mRNA by binding to UG motifs present on the stem loops of the 3'UTR (Strong *et al*, 2007; Volkening *et al*, 2009). UG-rich sequences are also present in the *Cdk6* 3'UTR and are bound by TDP-43, as determined by individual-nucleotide resolution CLIP (iCLIP)-seq (Tollervey *et al*, 2011). To test the hypothesis that TDP-43 modulates *Cdk6* expression in a similar manner in CHO-K1 cells, we first performed RIP assay to analyse the interaction between TDP-43 and *Cdk6* mRNA. TDP-43 IP specifically retrieved *Cdk6* mRNA but not *GAPDH* or other G1 phase-related genes, such as *Cdk4*, *Cdk2* and *Cyclin D1*, when compared with non-specific IgG IP (Figure 6A), indicating that TDP-43 binds to *Cdk6* mRNA in CHO-K1 cells. Furthermore, to validate whether the *Cdk6* 3'UTR is responsible for its binding to TDP-43, RNA pull-down assay was carried out with different regions of *Cdk6* mRNA: coding sequence (CDS), 3'UTR or full length (including both CDS and 3'UTR). Clearly, both 3'UTR and full length of *Cdk6* were able to bind to TDP-43 strongly,

but the CDS region failed to bind to TDP-43 (Figure 6B). The interaction specificity was examined by competition experiment. The binding of TDP-43 to *Cdk6* mRNA was efficiently competed by increasing amounts of non-biotinylated *Cdk6* mRNA (Figure 6C, second panel), but not *GAPDH* (Figure 6C, first panel). Taken together, these observations demonstrate that TDP-43 specifically associates with *Cdk6* 3'UTR in CHO-K1 cells.

We then sought to determine whether *gadd7* represses *Cdk6* expression by affecting the binding of TDP-43 to *Cdk6* mRNA. To this end, we first conducted RIP analysis in *gadd7*-depleted cells and observed that *gadd7* depletion led to increased interaction between TDP-43 and *Cdk6* mRNA (Figure 6D). The increased interaction with *Cdk6* mRNA was not the result of increased TDP-43 levels in *gadd7*-knockdown cells (Figure 5B), suggesting that *gadd7* antagonizes the ability of TDP-43 to bind to the *Cdk6* mRNA. The *gadd7* inhibitory effect is further supported by competition experiment (Figure 6C). Non-biotinylated *gadd7* RNA significantly reduced the association of TDP-43 with *Cdk6* mRNA in a dose-dependent manner, while *gadd7-Mut-2* RNA with mutated TDP-43 binding sites failed to interfere with the binding even at a high dose.

Moreover, since large sets of putative TDP-43 RNA targets, including 3'UTR targets, have been identified recently through RIP-seq or cross-linking and immunoprecipitation (CLIP)-seq, we also selected several 3'UTR targets, such as its own 3'UTR (Ayala *et al*, 2011; Polymenidou *et al*, 2011), *MEF2D* 3'UTR, *CSNK1A1* 3'UTR, *CSNK2A1* 3'UTR, *SLC1A2* 3'UTR (Tollervey *et al*, 2011) and *Gm* 3'UTR (Colombrita *et al*, 2012), to examine whether *gadd7* affects the binding of TDP-43 to other 3'UTRs. We observed that *gadd7* depletion led to an increase in the binding of TDP-43 to most of these targets (Figure 6D), coincident with the changed expression of these targets on *gadd7* depletion (Figure 6E), although the binding to its own 3'UTR or *SLC1A2* 3'UTR was not changed and the binding to *Gm* 3'UTR was reduced (Figure 6D). To avoid the interference from changed expression and re-association of mRNAs in native RIP, RIP was carried out in UV cross-linked CHO-K1 cells and normalized to transcript levels. Similar to what was observed in native RIP, depletion of *gadd7* was sufficient to increase the binding of TDP-43 to most of these 3'UTRs, including *Gm* 3'UTR, but not to its own 3'UTR or *SLC1A2* 3'UTR (Supplementary Figure S7). Since RIP with UV cross-linking could avoid the re-association of mRNAs, the binding of TDP-43 to *Gm* should have been increased *in vivo* when *gadd7* was knocked down.

Additionally, treatment with UV irradiation resulted in dismissal of TDP-43 from *Cdk6* mRNA, whereas this dismissal was significantly attenuated after depletion of *gadd7* (Figure 6F). These results correlate with the finding that the expression of Cdk6 was downregulated upon UV irradiation (Figure 6G; Supplementary Figure S8A and B), but depletion of *gadd7* attenuated the reduction of *Cdk6* by UV (Figure 6G). Taken together, these findings indicate that the interaction between TDP-43 and *Cdk6* mRNA is disrupted by *gadd7*, which at least partially contributes to downregulated expression of Cdk6 upon UV irradiation.

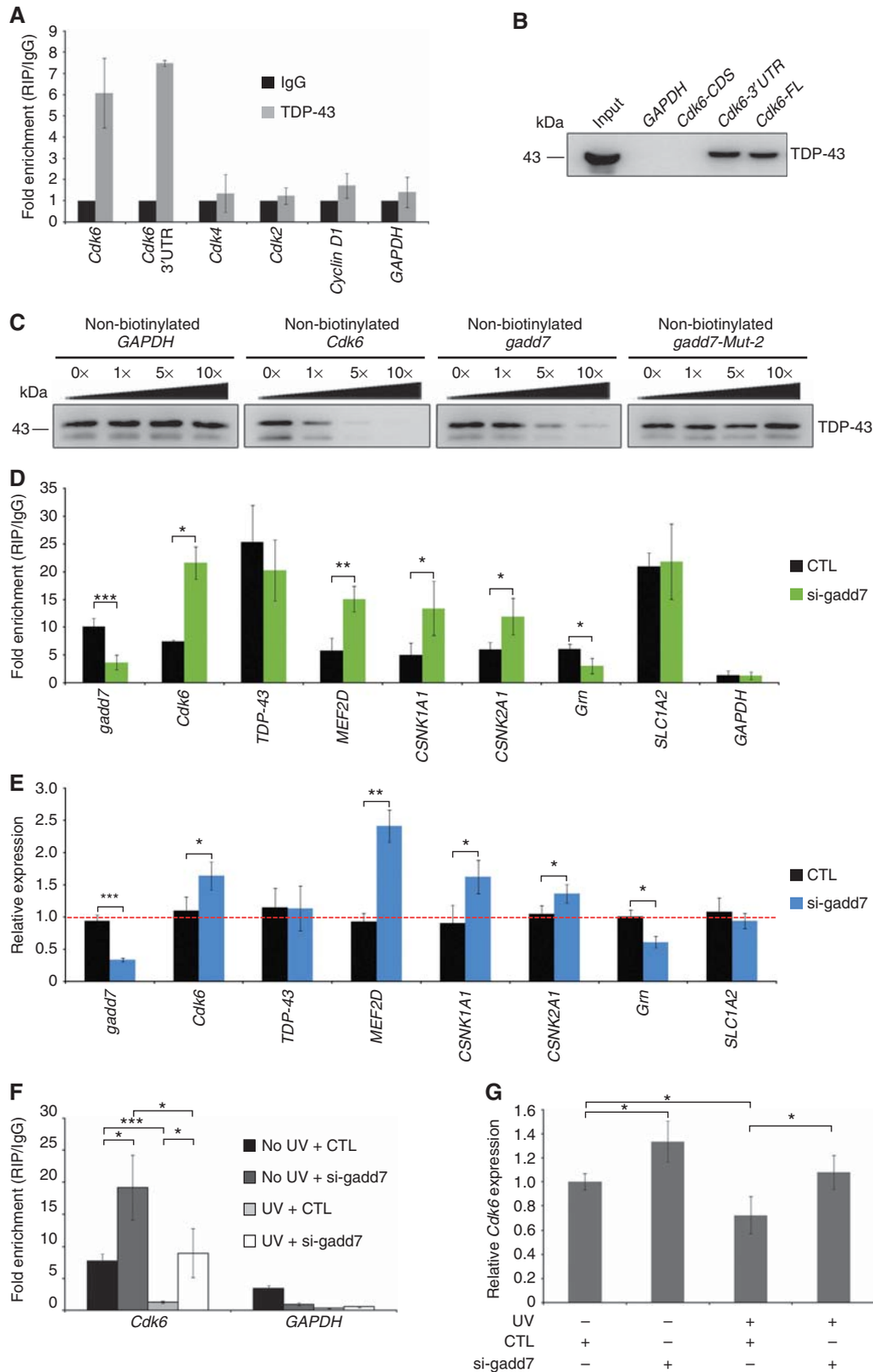
### ***gadd7* modulates *Cdk6* mRNA decay through TDP-43**

As the interaction between TDP-43 and *Cdk6* 3'UTR was disrupted by *gadd7*, we next aimed to determine whether

the stability of *Cdk6* mRNA is regulated by *gadd7*. We examined the decay of *Cdk6* mRNA following treatment of CHO-K1 cells with actinomycin D, a RNA Pol II inhibitor. Depletion of TDP-43 accelerated the degradation of *Cdk6* mRNA in the presence of actinomycin D relative to negative control. In contrast, depletion of *gadd7* showed a pattern of

much slower degradation of *Cdk6* mRNA upon actinomycin D treatment (Figure 7A), indicating that *gadd7* is involved in regulation of *Cdk6* mRNA decay.

We further conducted luciferase reporter assay to address the role of *gadd7* in the *Cdk6* mRNA stability. A 2026-bp fragment of *Cdk6* 3'UTR containing TG repeats was cloned



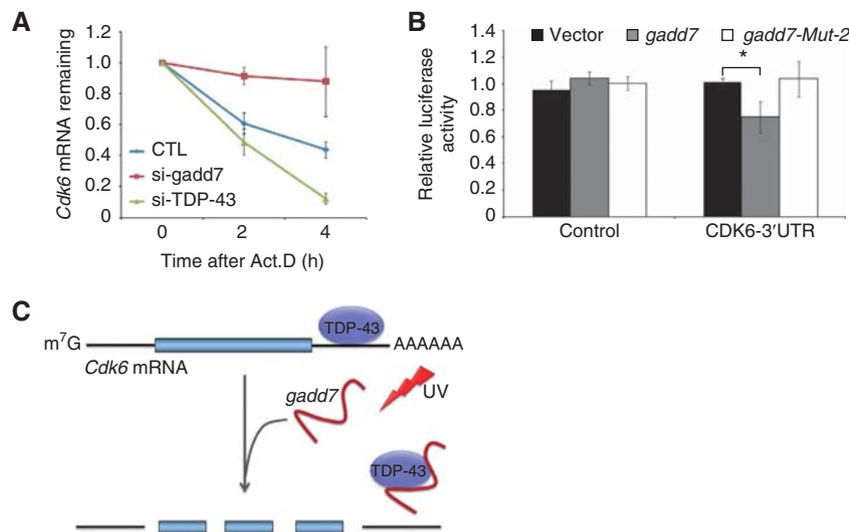
into the region downstream of the firefly luciferase gene of pGL3-Control vector. Both pGL3-Control vector and pGL3-*Cdk6*-3'UTR vector were transfected into CHO-K1 cells along with *gadd7* wild-type vector (pcDNA3.1-*gadd7*) or mutant vector (pcDNA3.1-*gadd7*-Mut-2). The results showed that overexpression of *gadd7* resulted in a reduction of luciferase activity, while the *gadd7* mutant did not have any influence on the activity relative to the control empty vector (Figure 7B). These findings demonstrate that the effect of *gadd7* on *Cdk6* mRNA stability is dependent on TDP-43.

## Discussion

Although lncRNAs are transcribed pervasively from mammalian genomes, the current understanding on the functions of lncRNAs is largely limited. Here we have identified a novel mechanism by which *gadd7*, a DNA damage-inducible

lncRNA, plays an important role in regulation of the cell-cycle G1/S checkpoint in response to DNA damage. We demonstrate that following UV irradiation *gadd7* is greatly induced and interacts with TDP-43, which in turn destabilizes *Cdk6* mRNA and promotes the degradation of *Cdk6* mRNA, thus leading to the inhibition of cell-cycle progression.

The cell-cycle G1/S checkpoint following DNA damage is an important event that maintains genomic integrity, and so disruption of the G1/S checkpoint may result in instability of the genome (Bartek and Lukas, 2001). To our knowledge, the G1/S checkpoint in the mammalian is governed by a series of protein factors, and there is little understanding on the functions of lncRNAs in the control of such an important event. In this study we find that as the lncRNA *gadd7* is implicated in regulation of the G1/S checkpoint. Following CHO-K1 cells' exposure to genotoxic stress such as UV irradiation or cisplatin treatment *gadd7* is rapidly induced,



**Figure 7** *gadd7* modulates *Cdk6* mRNA decay via TDP-43. (A) Knockdown of *gadd7* or TDP-43 affects *Cdk6* mRNA decay. CHO-K1 cells were transfected with siRNA control (CTL) or siRNA pool targeting *gadd7* or TDP-43. Forty-eight hours after transfection, cells were treated with actinomycin D (Act.D) for 0, 2 and 4 h. Total RNA was prepared and *Cdk6* mRNA expression was analysed by qRT-PCR and normalized to *GAPDH*. The transcript remaining is defined as relative expression at the indicated times compared with the expression level at 0 h. Data are from one of three independent experiments and are represented as mean  $\pm$  s.d. with  $n = 3$ . (B) Relative luciferase activity of the reporter containing the *Cdk6* 3'UTR is reduced in cells overexpressing *gadd7* wild type but not mutant compared with the control vector. Relative luciferase activity is the ratio of firefly luciferase to Renilla luciferase. Data are represented as mean  $\pm$  s.d. from three independent experiments. \* $P < 0.05$ . (C) Proposed model for the function of *gadd7* in regulating *Cdk6* mRNA stability. DNA damage-induced *gadd7* binds to TDP-43, and inhibits TDP-43 occupancy to *Cdk6* mRNA, thus leading to the instability of *Cdk6* mRNA.

**Figure 6** *gadd7* abrogates the interaction between TDP-43 and *Cdk6* mRNA. (A) TDP-43 binds to *Cdk6* mRNA *in vivo*. RIP with TDP-43 antibody was performed using CHO-K1 lysates and subsequently subjected to qRT-PCR analysis for the indicated RNAs. IP enrichment is determined as the amount of RNA associated to TDP-43 IP relative to IgG control. Data are from one of three independent experiments and are represented as mean  $\pm$  s.d. with  $n = 3$ . (B) *Cdk6* 3'UTR is responsible for its binding to TDP-43. RNA pull-down assay was performed with different regions of *Cdk6* mRNA. FL, full length. (C) Competition assay analysis of the binding of TDP-43 to *Cdk6* mRNA in the presence of various amounts of unlabelled *GAPDH* (first panel), *Cdk6* (second panel), *gadd7* (third panel) or *gadd7*-Mut-2 (fourth panel). (D) The binding of TDP-43 to its 3'UTR targets is affected by *gadd7* loss. CHO-K1 cells transfected with control siRNA (CTL) or siRNA pool targeting *gadd7* for 48 h were harvested for RIP assay with TDP-43 antibody and IgG. Binding of TDP-43 to the indicated RNA targets was determined by qRT-PCR with primers against their 3'UTRs. IP enrichment is determined as the amount of RNA associated to TDP-43 IP relative to the corresponding IgG control. Data are represented as mean  $\pm$  s.d. from three independent experiments. \* $P < 0.05$ , \*\* $P < 0.01$ , \*\*\* $P < 0.001$ . (E) The expression of indicated RNA targets as in (D) is changed upon *gadd7* depletion as analysed by qRT-PCR. Data are from one of three independent experiments and are represented as mean  $\pm$  s.d. with  $n = 3$ . \* $P < 0.05$ , \*\* $P < 0.01$ , \*\*\* $P < 0.001$ . (F) RIP analysis of the interaction between TDP-43 and *Cdk6* mRNA upon treatment with control siRNA or siRNA pool targeting *gadd7* in the presence or absence of UV irradiation (40 J/m<sup>2</sup>). Binding of TDP-43 to *Cdk6* mRNA was determined by qRT-PCR. IP enrichment is determined as the amount of RNA associated to TDP-43 IP relative to the corresponding IgG control. Data are represented as mean  $\pm$  s.d. from three independent experiments. \* $P < 0.05$ , \*\*\* $P < 0.001$ . (G) qRT-PCR quantification of *Cdk6* mRNA expression in CHO-K1 cells as treated in (F). Data are from one of three independent experiments and are represented as mean  $\pm$  s.d. with  $n = 3$ . \* $P < 0.05$ . Figure source data can be found with the Supplementary data.

while depletion of *gadd7* via the RNAi approach substantially abrogates cell-cycle G1 arrest after UV treatment, indicating that *gadd7* is required for regulation of the G1/S checkpoint after DNA damage. However, overexpression of *gadd7* alone in the absence of UV treatment appears not to induce a G1 arrest, suggesting that the well-established G1/S checkpoint may involve not only *gadd7*, but also other factors that are activated by DNA damage. This observation is consistent with the previous demonstration that *gadd7* overexpression alone does not increase cellular ROS levels or induce ER stress (Brookheart *et al*, 2009).

The mechanism by which *gadd7* regulates the G1/S checkpoint after DNA damage is investigated in this study. It has been shown that lncRNAs bind to and modulate the functions of proteins (Rinn *et al*, 2007; Huarte *et al*, 2010). Through RNA pull-down assay and mass spectrometry analysis, TDP-43, an hnRNP protein, was identified to associate with *gadd7*. This interaction is further supported by RIP, competition assay, deletion-mapping experiment and point-mutation experiment. We find that UG/GU repeats located at the two regions in the middle of *gadd7* are required for the interaction with TDP-43. We also investigated the interaction between *gadd7* and TDP-43 following UV treatment. As expected, in response to UV irradiation, the interaction of *gadd7* with TDP-43 was increased.

TDP-43 belongs to the family of hnRNPs, which is a multifunctional RNA/DNA-binding protein involved in transcription, splicing, mRNA transport and stability (Buratti and Baralle, 2008; Cohen *et al*, 2011). More importantly, recent studies have shown that dominant mutations in the TDP-43 cause a series of neurodegenerative diseases, including frontotemporal lobar degeneration (FTLD) and amyotrophic lateral sclerosis (ALS) (Kabashi *et al*, 2008; Sreedharan *et al*, 2008). Both the capacity of RNA binding to UG repeats and the interaction with other hnRNPs through the C-terminal region are critical for the function of TDP-43 (Buratti *et al*, 2005; D'Ambrogio *et al*, 2009; Kuo *et al*, 2009). Until now, many targets of TDP-43 have been identified, among which *Cdk6* is reported to be repressed by TDP-43 through recruitment to the UG-rich *Cdk6* transcript in HeLa cells (Ayala *et al*, 2008). *Cdk6* is a well-known member of the CDK family that is important for G1-phase progression (Lee and Yang, 2003). However, we here find that the expression of *Cdk6* is activated by TDP-43, which is not consistent with the previous report. The reason for this discrepancy may be that the mechanisms involved in control of TDP-43-mediated *Cdk6* expression are various in cell lines of different species and backgrounds. As is known, HeLa is a HPV-positive cancer cell line derived from human cervical cancer cells, but CHO-K1 is an immortalized cell line derived from CHO. Therefore, the mechanisms by which TDP-43 regulates *Cdk6* expression may be different in the two cell lines, thus leading to different outcomes. Indeed, as reported by Ayala *et al* (2008), the loss of TDP-43 has no effect on *Cdk6* protein levels in chicken cells (DF-1), since chicken *Cdk6* lacks TG repeats completely. Although all of human, rat, mouse and hamster *Cdk6* have the TG repeats, the distribution and amounts of TG repeats are quite different. Until now, to our knowledge, the regulatory effect of TDP-43 on *Cdk6* expression has not been well studied in other cell lines, although the binding of TDP-43 to *Cdk6* has been observed, as measured by RIP-seq or iCLIP-seq (Sephton *et al*, 2011;

Tollervey *et al*, 2011). The positive role of TDP-43 in *Cdk6* expression in CHO-K1 cells may be mediated by stabilizing its mRNA rather than inhibiting pre-mRNA splicing. Indeed, as proof of principle, TDP-43 binds to *Cdk6* 3'UTR, which leads to increase in its mRNA half-life, while *gadd7* decreases the *Cdk6* mRNA abundance by affecting the binding of TDP-43 to *Cdk6* mRNA. Of note, the *in vitro* binding affinity of TDP-43 to *Cdk6* is larger than to *gadd7* in competition experiments, although the *in vivo* binding of TDP-43 to *Cdk6* mRNA is weaker than to *gadd7* RNA in RIP assay. The reason for this discrepancy may be due to the differences between these two kinds of experiments, and also because RNA secondary structures probably mediate the association between TDP-43 and these targets. The secondary structures of biotin-labelled RNA transcripts produced *in vitro* may be different from those of *in vivo* transcripts. Furthermore, following UV irradiation, *gadd7* is substantially induced and more *gadd7* binds to TDP-43, thus leading to decreased interaction between TDP-43 and *Cdk6* mRNA and thus downregulation of *Cdk6* expression.

In addition to *Cdk6* mRNA, *gadd7* could also reduce the binding of TDP-43 to some other 3'UTR targets, such as *MEF2D*, *CSNK1A1*, *CSNK2A1* and *Grn*. However, *gadd7* did not affect the interaction between TDP-43 and another 3'UTR target, *SLC1A2*. Furthermore, *gadd7* is also not sufficient to affect the binding of TDP-43 to its own 3'UTR. TDP-43 has been reported to autoregulate its own protein level by directly binding and enhancing splicing of intron 7 in its own 3'UTR, thereby triggering degradation (Polymenidou *et al*, 2011; Avendano-Vazquez *et al*, 2012). Hence, splicing of intron 7 in its own 3'UTR is a critical event for TDP-43 autoregulation. *gadd7*, however, seems to have no effect on alternative splicing of TDP-43 targets as measured by splicing assay and CFTR reporter assay, which may be the reason for the unchanged binding to its own 3'UTR and expression level upon *gadd7* depletion. Collectively, we conclude that *gadd7* is able to affect the interactions between TDP-43 and some of its 3'UTR targets. One of the possible mechanisms by which *gadd7* affects the interaction between TDP-43 and its RNA targets is by directly binding to and sequestering TDP-43 from its 3'UTR targets. Furthermore, other indirect effects may also exist. TDP-43 has extensive interactions with RNA-binding proteins (Freibaum *et al*, 2010; Sephton *et al*, 2011). The changed expression of these RNA-binding proteins or interaction with TDP-43 upon *gadd7* depletion may also affect the binding of TDP-43 to its targets. Moreover, under stress, TDP-43 is localized to stress granules (SGs) (Colombrita *et al*, 2009; Freibaum *et al*, 2010; McDonald *et al*, 2011), which are dynamic triage centres for mRNA storage, decay or re-initiation during stress conditions (Anderson and Kedersha, 2008); thus, the localization of TDP-43 to SGs may likely contribute to the changed interaction with its RNA targets upon *gadd7* depletion.

Like other lncRNAs, such as *Xist* and *rncs-1*, *gadd7* is poorly conserved in sequence, since sequence searches have not revealed homologues in human or mouse species (Pang *et al*, 2006; Hellwig and Bass, 2008). In fact, the secondary structure of lncRNAs is more highly conserved than the nucleotide sequence (Washietl *et al*, 2005; Torarinsson *et al*, 2006). Thus, orthologues with similar secondary structure or functional elements to *gadd7* may exist in humans or mice and exert their function by

modulating TDP-43 function. Identification of *gadd7* orthologues in humans will be critical for exploring the pathogenesis of FTLD and ALS that is correlated with TDP-43.

In summary, we demonstrate that *gadd7*, as the first DNA damage-inducible lncRNA, is implicated in regulation of the G1/S checkpoint. Hereby, we propose a model in which UV-induced *gadd7* binds to TDP-43 and disrupts the interaction of TDP-43 with *Cdk6* mRNA, thus resulting in the instability of *cdk6* mRNA (Figure 7C). Our findings indicate that lncRNAs may serve as regulators by dissociating the protein–RNA interaction, thereby modulating the mRNA expression at the post-transcriptional level.

## Materials and methods

### Plasmids

The full-length *gadd7* sequence lacking a poly A tail (on the basis of the *gadd7* sequence reported as L40430 in GenBank) was amplified by PCR on the complementary DNA (cDNA) of CHO-K1 cells, and cloned into the *HindIII/XbaI* sites of the mammalian expression vector pcDNA3.1(+) (Invitrogen) to generate pcDNA3.1-*gadd7* vector. pcDNA3.1-antisense-*gadd7* was constructed by subcloning the antisense of *gadd7* into *XbaI/HindIII* sites of the pcDNA3.1(–) vector (Invitrogen). Two mutant versions of *gadd7* cDNA with *HindIII/XbaI* sites and three mutated TGTGTG elements or all mutated TG/GT repeats, respectively, were chemically synthesized by Invitrogen and inserted into pcDNA3.1(+) to create pcDNA3.1-*gadd7*-Mut-1 and pcDNA3.1-*gadd7*-Mut-2. A 2026-bp fragment of *Cdk6* 3'UTR containing TG repeats (on the basis of the *Cdk6* mRNA sequence reported as XM\_003496965.1 in GenBank) was PCR amplified from CHO-K1 cells' cDNA, and cloned into the *XbaI* site of pGL3-Control vector (Promega) to generate pGL3-*Cdk6*-3'UTR vector.

### Cell culture and treatment

CHO-K1 cells were purchased from the Cell Culture Center, Chinese Academy of Medical Sciences (Beijing, China). CHO-K1 cells were grown in Dulbecco's modified Eagle's medium/Ham's F-12 (DMEM/F12, 1:1) medium supplemented with 5% fetal bovine serum (Gibco) and 2 mM L-glutamine at 37°C with 5% CO<sub>2</sub>. For UV treatment, exponentially growing CHO-K1 cells were rinsed with phosphate-buffered saline (PBS) and irradiated with UV radiation at different doses. After cells' exposure to UV irradiation, fresh medium was added and the cells were cultured for 4 or 6 h till harvest. For cisplatin treatment, exponentially growing CHO-K1 cells were grown in medium containing cisplatin at different concentrations for 4 h till harvested. For media depletion, CHO-K1 cells were grown to confluence, then held without refeeding for an additional 48 h. For serum reduction, exponentially growing CHO-K1 cells were cultured with fresh DMEM/F12 media with 5% or 0.5% serum for 24 h. For actinomycin D (Serva) treatment, exponentially growing CHO-K1 cells were treated with 500 ng/ml actinomycin D for 0, 2 and 4 h. For nocodazole (Sigma) treatment, exponentially growing CHO-K1 cells were treated with 200 ng/ml nocodazole for 6 or 12 h.

### RNA interference

In all, 20–30% confluent CHO-K1 cells were transfected with 50 nM of siRNAs using Lipofectamine 2000 (Invitrogen) following the manufacturer's direction. All siRNAs were obtained from Invitrogen (25-mer duplex Stealth siRNAs for *gadd7* and TDP-43; 21-mer duplex siRNA for *Cdk6*). The target sequences are the following: siRNA-1, 5'-GAGCAGGAGAAUUGUCUACAGA-3' and siRNA-2, 5'-CCAGUGACACAAGUGAGAAUUGGAU-3' for *gadd7*; siRNA-1, 5'-UGGUUCAGGUCAAGAAAGAUUUUA-3' and siRNA-2, 5'-CAGCGUGCAUUAUCCAAUGCUGAA-3' for TDP-43; siRNA-1, 5'-CCUAAAGCCACAGAACAUUTT-3' and siRNA-2, 5'-GCAAGAGUGAUUGCAGCUUTT-3' for *Cdk6*.

### Viability and cell-cycle analysis

CHO-K1 cells grown in 24-well plates were transfected with 50 nM of control siRNA or siRNAs targeting *gadd7*. At 24, 48 and 72 h after

transfection, cell numbers were counted and plotted for the growth curves. MTT absorbance of CHO-K1 cells was detected by using a microplate reader at 570 nm 72 h post transfection. For cell-cycle analysis, 48 h after transfection, cells grown in six-well plates were harvested, washed with PBS and fixed with ice-cold 70% ethanol at –20 °C overnight. Samples were then washed with PBS and stained with propidium iodide (PI) (Sigma) containing RNase A (Sigma) for 30 min at 37°C. Cell-cycle distribution was measured by flow cytometry (Becton Dickinson).

### Luciferase reporter assay

CHO-K1 cells were transfected by Lipofectamine 2000 (Invitrogen) in 96-well plates with 100 ng pGL3-Control vector or pGL3-*Cdk6*-3'UTR vector, 2 ng pRL-CMV control vector and 100 ng pcDNA3.1-*gadd7* or pcDNA3.1-*gadd7*-Mut-2. Forty-eight hours after transfection, cells were harvested and analysed using a dual-luciferase assay kit (Promega). Firefly activity was normalized to Renilla luciferase activity.

### RNA isolation, real-time RT-PCR and semi-quantitative RT-PCR

Total RNA from CHO-K1 cells was extracted with TRIzol (Invitrogen). First-strand cDNA was synthesized by using the Superscript II-reverse transcriptase kit (Invitrogen) according to the manufacturer's instructions. Real-time PCR (qPCR) was conducted using SYBR Premix Ex Taq (Takara) on an ABI 7300 Real-Time PCR System (Applied Biosystems). Gene-specific qPCR primer pairs are provided in Supplementary Table S2. All samples were normalized to *GAPDH*.

Semi-quantitative RT-PCR was conducted using the Taq polymerase (Takara). The gene-specific PCR primer pairs used are as follows: *gadd7*, forward 5'-GGGAAGCTGAGGTTTTTC-3', reverse 5'-CACACCAGTCTCAACTCCC-3'; *GAPDH*, forward 5'-GCTGAGAA CGGAAGCTTGT-3', reverse 5'-GCCAGGGTGCTAAGCAGTT-3'; *Cdk6*, forward 5'-GCTGAGGCACCTGGAGACCT-3', reverse 5'-GTC CAGACCTCGGAGAAGCTG-3'. *GAPDH* was used as the internal control.

### Immunoblotting

Cell protein lysates were prepared using 1 × PBS supplemented with 1% Nonidet P-40, 1 × protease inhibitors cocktail (Roche) and 50 µg/ml phenylmethylsulphonyl fluoride. Proteins were resolved by 10% SDS-PAGE and transferred to polyvinylidene difluoride membrane. Membranes were incubated with the indicated primary antibodies and anti-mouse or anti-rabbit secondary antibodies conjugated to horseradish peroxidase (HRP). After addition of HRP substrate, the chemiluminescence signal was detected with X-ray film or Luminescent Image Analyzer LAS-4000 (Fujifilm). Antibodies used for immunoblotting are as follows: rabbit anti-TDP-43 (Proteintech Group, 10782-2-AP, 1:1000), mouse anti-*Cdk6* (Santa Cruz Biotechnologies, sc-56282, 1:300), rabbit anti-Cyclin D1 (Santa Cruz Biotechnologies, sc-753, 1:500) and mouse anti-Actin (Sigma-Aldrich, A1978).

### RNA pull-down assay

RNA pull-down assay was performed as described previously, with some modifications (Rinn *et al*, 2007). The plasmids, pcDNA3.1-*gadd7*/pcDNA3.1-*gadd7*-Mut-1/pcDNA3.1-*gadd7*-Mut-2/pcDNA3.1-antisense-*gadd7*, were linearized with the corresponding restriction enzymes used to clone the vector at the 3', in order to prepare the template DNAs for *in vitro* transcription of *gadd7* wild type, point mutants or antisense. Template DNAs for *in vitro* transcription of *GAPDH*, various deletion mutants of *gadd7* or different fragments of *Cdk6* mRNA were PCR amplified from CHO-K1 cells' cDNA using the primers containing T7 promoter sequence in the sense primer. The primers used are listed in Supplementary Table S3.

All biotin-labelled RNA transcripts were produced *in vitro* using the MEGAscript T7 Kit (Ambion) with biotin-16-UTP (Ambion), and purified by MEGAclear Kit (Ambion) according to the protocol provided by the manufacturer. Ten picomoles of biotinylated RNAs was heated to 56°C for 5 min and 37°C for 10 min, and then slowly cooled to 4°C. RNAs were mixed with 500 µg of pre-cleared CHO-K1 cell extracts in binding buffer (10 mM HEPES pH 7.0, 50 mM KCl, 10% glycerol, 1 mM EDTA, 1 mM DTT, 0.5% Triton X-100) supplemented with tRNA (0.1 µg/µl), heparin (0.5 µg/µl)

and RNasin (1 unit), and incubated at room temperature for 30 min. Fifty microlitres of washed streptavidin agarose beads (Invitrogen) was added to each binding reaction and further incubated at room temperature for 15 min. Beads were washed five times with the binding buffer, and boiled in 1 × loading buffer for 10 min. The retrieved proteins were subjected to SDS-PAGE, and further visualized by silver staining or immunoblotting assay. Protein bands were excised and identified by in-gel trypsin digestion and analysed by mass spectrometry.

#### Native RIP and UV cross-linking RIP

Native RIP experiment was carried out with the EZ-Magna RIP Kit (Millipore) according to the manufacturer's protocol using 5 µg of rabbit anti-TDP-43 antibody (Proteintech Group, 10782-2-AP) or rabbit IgG (Millipore). The co-precipitated RNAs were extracted with phenol:chloroform:isoamyl alcohol and detected by semi-quantitative RT-PCR or qRT-PCR. Proteins isolated from the beads were detected by immunoblotting. For UV cross-linking RIP, cells were irradiated with 100 mJ/cm<sup>2</sup> UV light before performing RIP assay as described previously (Anko *et al*, 2012).

#### Statistical analysis

Data are presented as mean ± s.d. Statistical analyses were carried out with Student's *t*-tests. Differences were considered statistically significant at *P* < 0.05.

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#### Supplementary data

Supplementary data are available at *The EMBO Journal* Online (<http://www.embojournal.org>).

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*Author contributions:* XL, MG and QZ designed the experiments. XL, DL and WZ performed the experiments and analysed the data. XL and QZ wrote the paper.

## Conflict of interest

The authors declare that they have no conflict of interest.

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