DIFFERENTIAL EFFECTS OF SUMOYLATION ON TRANSCRIPTION AND ALTERNATIVE SPlicing BY TRANSCRIPTION ELONGATION REGULATOR 1 (TCERG1)

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Running Head: Sumoylation regulates TCERG1 activity

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Modification of proteins by small ubiquitin-like modifier (SUMO) is emerging as an important control of transcription and RNA processing. The human factor TCERG1 (also known as CA150) participates in transcriptional elongation and alternative splicing of pre-mRNAs. Here we report that SUMO family proteins modify TCERG1. Furthermore, TCERG1 binds to the E2 SUMO-conjugating enzyme Ubc9. Two lysines (K503 and K608) of TCERG1 are the major sumoylation sites. Sumoylation does not affect localization of TCERG1 to the splicing factor-rich nuclear speckles or the alternative splicing function of TCERG1. However, mutation of the SUMO acceptor lysine residues enhanced TCERG1 transcriptional activity, indicating that SUMO modification negatively regulates TCERG1 transcriptional activity. These results reveal a regulatory role for sumoylation in controlling the activity of a transcription factor that modulates RNAPII elongation and mRNA alternative processing, which are differently discriminated by this post-translational modification.

Splicing and transcriptional elongation are physically and functionally interconnected processes (1,2). While both processes can occur autonomously, their coupling and coordination may be important for regulation of gene expression. Coupling of these two processes may influence splicing and alternative splicing regulation. Indeed, promoter composition, transcriptional elongation efficiency, chromatin environment and recruitment of specific coregulators to the transcriptional complex have been shown to affect alternative splicing decisions in a number of experimental systems (3-6). Yet the molecular mechanisms at work are not understood. The unique C-terminal domain (CTD) of the large subunit of RNAPII seems to play a central role in the coupling of splicing, as well as other RNA processing functions, to transcription (7,8). To provide a framework, we consider two models: the recruiting and kinetic models, which are not mutually exclusive. In the “recruiting model” the CTD functions as a “landing pad” for specific subsets of RNA processing factors in a manner dependent of its phosphorylation pattern, and therefore on the functional state of the transcription complex (9). A number of independent research lines have suggested that RNA splicing factors can interact with RNAPII molecules that are hyperphosphorylated on their CTD (10-14), although these associations might be highly dynamic and transient in vivo. The integrity of the RNAPII CTD has also been shown to influence the recruitment of splicing factors to active transcription sites in the nucleus (15). The “kinetic model” (1) proposes that the rate of elongation of the nascent transcript affects specific alternative splicing decisions by modulating the probability of simultaneous presentation of competing splicing sites. Thus, modulation of transcription elongation efficiency at specific, alternatively spliced regions of genes, might constitute a mechanism to regulate splicing decisions (5). Several nuclear factors have been shown to play dual roles both in transcription and splicing regulation, and/or to exhibit physical interaction with components of both machineries, thus revealing themselves as potential “coupling mediators” or “crosstalk” factors; these include among others PSF, p54nrb/NONO, TAT-SF1, Prp40, FBP11, and TCERG1. PSF and p54nrb/NONO influence both transcription and splicing, and are components of high molecular weight complexes that include processive RNAPII as well as a broad subset of splicing factors (16,17). Tat-SF1 is an elongation regulator factor that interacts with UsnRNP complexes (18). Yeast Prp40 and
mammal FBP11 are related proteins that contain tandem repeats of WW and FF domains, a feature that might define a subset of transcription and splicing-related factors; consistent with this hypothesis, they both interact with the CTD of RNAPII and U1 snRNP (13,19-21).

TCERG1 (previously designated CA150) was first described as a transcriptional elongation regulator found in HIV-1 Tat-responsive HeLa nuclear extract fractions (22). Transient overexpression of TCERG1 reduces expression from the HIV-1 and α-4 integrin promoters by inhibiting elongation efficiency. This repression is promoter specific and it is also dependent on a specific functional TATA-box element (23). The primary sequence of TCERG1 contains three WW domains in the amino-terminal half and six FF repeat motifs in the carboxyl-terminal half. TCERG1 interacts directly with the phosphorylated CTD of RNAPII via its FF repeats (24) and binds to several elongation-related factors in HeLa nuclear extracts (25). Based on those data, TCERG1 appears to function in the elongation stage of transcription. However, accumulating evidence indicates a potential role of TCERG1 in splicing, and hence, in the coupling between transcription and splicing. TCERG1 copurifies with in vitro assembled spliceosomes, and was identified in spliceosomal subcomplexes (26-28). In addition, we and others have found multiple interactions with components of the splicing machinery (25,29-31). The subnuclear distribution of TCERG1 resembles that of an RNA-metabolism related factor with enrichment in the peripheral regions of the splicing factor-rich nuclear speckles (25). Importantly, TCERG1 can affect alternative pre-mRNA splicing of β-globin, β-tropomyosin, and CD44 splicing reporters (30,32,33) and in putative cellular targets identified upon TCERG1 knockdown by microarray analysis (33).

TCERG1 may be regulated at multiple levels. TCERG1 forms multiple protein complexes, subpopulations of which may differ in their functional properties and biochemical associations. Compartmentalization in the nuclear subdomains may control TCERG1 function. Post-translational modifications may also influence TCERG1 function: for instance, a recent report demonstrated that TCERG1 interacts with the spinal muscular atrophy protein SMN when methylated by CARM1. This modification modulated the functional interaction of TCERG1 and CARM1 to affect alternative splicing of a CD44 exon 4 (32). Phosphorylation of specific motifs on TCERG1 sequence has also been reported, although its functional significance remains unknown (34).

A number of Small Ubiquitin-like MOdifier paralogs have been described in higher eukaryotes: SUMO-1, SUMO-2, and SUMO-3 (SUMO-2 and SUMO-3 are approximately 96% identical and referred to us as SUMO2/3). They seem to modify different, partially overlapping subsets of cellular factors (35) and potentially could supply the absence of each other in vivo (36). In a cyclic process highly related to ubiquitination, SUMO modifiers are activated by an E1 activity (Uba/AOS heterocomplex), directed to the target substrate by an E2 activity (Ubc9), and covalently attached to the lysine residue that is usually embedded in a minimal motif ΨKxD/E in the substrate primary sequence, where Ψ stands for a large hydrophobic amino acid and x stands for any residue. This last step often requires an E3 group of substrate-specific ligases, at least in vivo (37). Unlike monoubiquitination, SUMO modification does not normally target proteins for proteolytic degradation. Instead, sumoylation modulates a wide range of properties of the protein substrates, including subnuclear localization, protein stability and functional interactions. Sumoylation has been shown to constitute a pivotal mechanism of transcriptional regulation, and its transcriptome-wide effect has been proposed to be transcription inhibition (38). The molecular basis of SUMO-driven transcriptional modulation is not well understood. Sumoylation of components of the transcription machinery promotes recruitment of chromatin remodelling complexes, such as histone deacetylases (HDACs) (39). Sumoylation can also modify the affinity of factors for target DNA sequences (40). RNA processing factors are also SUMO-targets, but the consequences have not been elucidated. For example, SUMO is implicated in the regulation of assembly of the 3’end processing machinery (41), thus revealing the potential importance of these modifiers as general mRNA metabolism regulators.

In this study, we investigated the role of sumoylation in modulating the function of TCERG1 in transcription and mRNA processing. We identified TCERG1 as a target for sumoylation in vivo. Sumoylation occurs on at least two sites in the amino terminus of TCERG1. Mutation of those lysines abolishes sumoylation both in vivo and in vitro. Arginine
substitution of those lysines did not interfere with TCERG1 localization to the splicing-factor rich nuclear speckles, nor was the activity of TCERG1 in mRNA processing affected. Instead, mutation of the sumoylation sites increased the transcriptional activity of TCERG1 in cell reporter assays. Thus, our data strongly support a role for SUMO modification in regulating specific aspects of TCERG1 function.

**Experimental Procedures**

**Plasmids.** The eukaryotic expression plasmids pEFBOST7-TCERG1 (1-1098) and (1-662), and bacterial expression vectors pGEX2TK-TCERG1 (234-662) and (631-1098) have been previously described (23,24). The plasmid pEFBOST7-TCERG1 (616-1098) was created by cloning a PCR product with BglII ends into the BamHI site of the parental vector pEFBOST7 (42). The pEFBOST7-TCERG1 full length single K503R, K608R and double K503,608R mutants were obtained by using the QuikChange XL II directed mutagenesis kit (Stratagene), and mutations were verified by sequencing. The enhanced cyan fluorescent protein (ECFP) fragment was obtained by PCR from pECFP-C1 plasmid (Invitrogen) and inserted in an EcoRI-digested pEFBOST7 fragment (4677 bp) yielding an intermediate vector pEF-ECFP. pEFBOS/ECFP/T7-TCERG1, pEFBOS/ECFP/T7-TCERG1/K503R, pEFBOS/ECFP/T7-TCERG1/K608R and pEFBOS/ECFP/T7-TCERG1/K503,608R were obtained using EcoRI fragments obtained from the corresponding pEFBOST7 parental vectors by standard cloning procedures. pGAL4-TCERG1 wild-type and mutants (K503R, K608R, and K503,608R) were constructed by inserting appropriate sequences into SacI/EcoRI-digested pSG424 vector (43). For expression of recombinant proteins, pGEX2TK-TCERG1 (234-662) and pGEX2TK-TCERG1 (631-1098) have been described previously (24). The TCERG1 deletion expression vectors pGEX2TK-TCERG1 (134-641), wild-type and SUMO mutants (K503R, K608R, and K503,608R), were cloned by inserting TCERG1 PCR fragments into the BamHI/EcoRI sites of pGEX2TK vector (Amersham). Expression vectors pcDNA3-HA-SUMO1 and SUMO2/3 were kindly provided by Dr. R.T. Hay (University of Dundee). For HA-Ubc9 and SUMO2 recombinant expression and native purification, BamHI/EcoRI fragments were obtained by PCR or enzymatic digestion from pSUbc9V5 (R.T. Hay) or pcDNA3-HA-SUMO2/3 expression vectors, respectively, and inserted in a BamHI/EcoRI digested pGEX2TK vector. EDI splicing reporter minigenes pSVEDAmFN (fibronectin promoter) and pSVEDATot (α-globin promoter) were a kind gift from A. R. Kornbluh (Universidad de Buenos Aires, Argentina) and and F. E. Baralle (ICGEB, Trieste), respectively, and have been described previously (44,45). The reporter construct pSVEDA-TK contains the herpes simplex virus (HSV) thymidine kinase (TK) promoter and was constructed by inserting a HindIII/BglII-digested pRL-TK (Promega) fragment into a Scal/BssHII-digested and blunted pSVEDATot plasmid. pSL3b-based Tau exon 10 splicing reporter has been previously described (46,47) and was kindly provided by Tom Misteli (NIH, Bethesda). p3X-κB-L and pRL-TK plasmids were kindly provided by M.A. Garcia-Blanco (Duke University, Durham).

**Antibodies.** Antibodies against TCERG1 have been described previously (25). Immunoglobulins (Igs) were purified from antiserum on protein A-sepharose columns (Pharmacia) by following standard procedures and used at dilutions of 1:10,000. To affinity purify the antibodies, antiserum was incubated with nitrocellulose containing immobilized TCERG1 protein as described by Lin and coworkers (30). Affinity-purified anti-GST antibodies were obtained by standard affinity chromatography through glutathione-sepharose columns covalently attached to purified GST (48). The anti-SUMO1 monoclonal antibody (Santa Cruz Biotechnology) was provided by M. Lafarga (Universidad de Cantabria), and used at dilutions of 1:100 in western blot analysis. The anti-T7 (Bethyl) or anti-HA 12CA5 (Roche) antibodies were used at dilutions of 1:30,000 and 1:4,000 to detect T7 or HA epitope-tagged proteins, respectively. Antibody against cyclin T1 (Santa Cruz Biotechnology) was used at a dilution of 1:500. For immunofluorescence studies, we used anti-T7 and anti-SC35 (Sigma) antibodies at dilutions of 1:1,000 and 1:2,000, respectively.

**Cell culture, transfection, and reporter gene assays.** HEK293T cells were grown in Dulbecco’s modified Eagle medium (DMEM) (Sigma) supplemented with 10% fetal bovine serum (FBS) (Gibco), L-glutamine at 4mM (Invitrogen), and penicillin-streptomycin to 100U and 100μg per ml, respectively.
For luciferase assays, transfections were performed in 35-mm-diameter plates (Nunc). Each plate was seeded with approximately $1 \times 10^5$ cells 20 h prior transfection. Cells were grown to approximately 60-70% confluence and transfected with the appropriate amounts of the indicated constructs by using calcium phosphate and/or Polyfect reagent (Qiagen Inc.) according to the manufacturer’s protocols. Transfections were carried out with at least two different preparations of each plasmid DNA purified using kits from Qiagen Inc. Approximately 48 h after transfection, cells were harvested and lysed in 0.2 ml of T7 buffer (20 mM HEPES, pH 7.9, 150 mM NaCl, 5 mM EDTA, 1% Nonidet P-40, 1 mM dithiothreitol [DTT], 1 mM phenylmethylsulphonyl fluoride [PMSF]) for 30 min at 4ºC. The luciferase activities (firefly luciferase for the reporter and renilla luciferase for the internal control) were measured by using the Dual-Luciferase Assay System (Promega) and a standard manual luminometer (Berthold). All experiments were performed in triplicate and at least 3 independent experiments were considered to calculate mean values and standard deviations. For RT-PCR analysis, total RNA was isolated from HEK293T cells by using the TRIzol reagent (Invitrogen) and then reverse transcribed by using M-MLV reverse transcriptase (Invitrogen). The resulting RT reaction product was analyzed for EDI and tau splicing products by radioactive PCR as previously described (44,49).

Quantification of EDI transcripts (Figure 4I) was carried out by real-time RT-PCR using the MasterMix SYBRgreen reagent (Bio-Rad), the iCycler thermal cycler station (Bio-Rad), and previously described protocols (50,51).

Immunofluorescence. Immunofluorescence studies were performed as described previously (25) using a spectral laser confocal microscope Leica SP5. Images were analyzed and digitally processed for presentation using LAS AF software.

In vivo sumoylation assays. Transfected cells were harvested in PBS and lysed in 0.2 ml of ice-cold RIPA-375 buffer (50mM TrisHCl, pH8, 375 mM NaCl, 1% Triton X-100, 0.5% sodium-deoxycholate, 0.1% sodium dodecyl sulphate [SDS], 5mM EDTA, 1mM DTT, 1mM PMSF, and protease inhibitor mixture [Complete, Roche Molecular Biochem.]) for 20 minutes. Chromatin was sheared by repeated passage of the extracts through 23G needles, and debris was cleared by centrifugation at 14,000 rpm for 5 minutes at 4ºC. Ten per cent of the lysate was saved as input samples (WCE) and the remaining extract was diluted to a 1-ml final volume with ice-cold RIPA-375. Thirty microliters of anti-T7 tag monoclonal antibody that was covalently coupled to cross-linked agarose beads (Novagen) was added to the diluted extract and then incubated with end-over-end rotation for 6 h at 4ºC. After five washes with 1.5 ml of RIPA-375 and once with 1.5ml of T7 buffer, proteins bound to the antibody resin were boiled in 30 µl of 2x SDS-PAGE sample buffer. Fifteen microliters of each input and pellet sample were subjected to SDS-PAGE and transferred to a polyvinylidene difluoride (Bio-Rad) or nitrocellulose (Amersham) membrane, and then incubated with the specific antibody. After being washed, the membrane was incubated with a peroxidase-conjugated secondary antibody, and bound antibodies were detected by enhanced chemiluminiscence (Perkin-Elmer Life Sciences).

To assess steady-state sumoylation of endogenous TCERG1, approximately $3 \times 10^5$ HEK293T transfected cells were lysed in buffer T7 with or without 25 mM N-ethylmaleimide (NEM) and 1% SDS. Lysates were subjected to Western blotting analysis as described above. Immunoprecipitation of endogenous TCERG1 was conducted as follows: approximately $5 \times 10^6$ HEK293T cells were lysed in SDS-HS buffer (1%SDS, 0.5% Triton-X100, 50mM TrisHCl pH 8, 5mM EDTA, 1mM DTT, 500mM NaCl, 0.5mM DTT, 25mM NEM, 1mM PMSF, and protease inhibitor mixture [Complete, Roche Molecular Biochem.]) and chromatin was sheared by repeated passage of the extracts through a 23G needle. The cell extracts were

Immunofluorescence studies were performed as described previously (25) using a spectral laser confocal microscope Leica SP5. Images were analyzed and digitally processed for presentation using LAS AF software.
diluted 10 times in non SDS-HS buffer (0.5% Triton-X100, 50mM TrisHCl pH 8, 5mM EDTA, 500mM NaCl, 0.5mM DTT, 25mM NEM, 1mM PMSF, and protease inhibitor mixture [Complete, Roche Molecular Biochem.]), and split in two equal volumes. Fifty microliters of anti-TCERG1 IgG antibodies (100 µg) that was covalently coupled (25) to cross-linked protein A sepharose beads (4FastFlow, Amersham) was added to the diluted samples and incubated with end-over-end rotation for 4 hours at 4°C. Purified non-specific rabbit IgGs were used as a control. After extensive washing with RIPA-375 buffer, immunoprecipitates were boiled in 50µl of 2x SDS-PAGE sample buffer and analyzed by Western blotting.

**Protein purification.** Proteins were expressed in BL21 (D3) cells and purified on glutathione-agarose beads (Sigma) following standard procedures. The release of the fusion protein and the cleavage to yield free proteins were carried out as described by Smith and Johnson (52). hUbc9 protein was dialyzed against PBS containing glycerol at 20%; SUMO-2 was dialyzed against IVS buffer (20 mM HEPES pH 7.5, 5mM Mg, 100 mM NaCl, 0.5 mM DTT, 0.5 mM PMSF); GST-TCERG1 proteins were dialyzed against a buffer described in Lin et al. (30).

**In vitro interaction and sumoylation assays.** In-solution interaction assay was performed essentially as previously described (53,54) with minor modifications. Briefly, 5 µg of HA-hUbc9 were incubated with 20 µg of the appropriate GST fusion protein pre-bound to 20 µl of glutathione agarose, in a 500-µl final volume of binding buffer (40 mM HEPES pH 7.9, 150 mM KCl, 5 mM MgCl2, 0.5 mM EDTA, 10% glycerol, 0.1%NP-40, 1 mM DTT, and 1 mM PMSF) with end-over-end rotation for 4 h at 4°C. Pellets were washed extensively with binding buffer, boiled in 2x SDS-PAGE buffer, and then separated on SDS-PAGE gels and analyzed by Coomasie staining and Western blotting.

**In vitro-SUMO modification of TCERG1.** was performed using a commercial sumoylation kit (LAE Biotech.). The reaction mixtures were performed in 20 µl of the suggested buffer (20 mM HEPES pH 7.5, 5mM Mg, 100 mM NaCl +/- ATP) with 500 ng of the GST-TCERG1 recombinant proteins, incubated at 37°C for 90 min, and terminated by addition of SDS-PAGE sample buffer. The samples were then subjected to SDS-PAGE and Western blotting with specific antibodies.

**RESULTS**

TCERG1 is modified by SUMO-1 and SUMO2/3 in vivo. TCERG1 potentially couples transcription and mRNA processing; however, the molecular mechanisms by which TCERG1 modulates transcriptional elongation and splicing remain to be elucidated. We postulated that the activities of TCERG1 may be regulated by posttranslational modification. We began our analysis by examining whether TCERG1 can be modified by sumoylation in vivo. The steady state level of most sumoylated proteins is very low, presumably caused by strictly regulated modification and/or rapid cycles of modification and de-modification. This often causes a detection problem of sumoylation in vivo. To overcome this obstacle, we have used protocols that involve enrichment of sumoylated proteins under strong denaturing conditions. HeLa whole cell lysates were obtained in the absence or presence of the cysteine protease inhibitor NEM (N-ethylmaleimide) or 1% SDS buffer, which are usually used to preserve the sumoylation of cellular proteins, and then subjected to immunoblotting with anti-TCERG1 specific antibodies. Along with the 150-kDa TCERG1, the antibodies detected a predominant slow migrating band with apparent mass of 200 kDa. This band is dependent on extracts prepared with NEM and 1% SDS buffer (Fig. 1A, lanes 2 and 4). Upon treatment with 1% SDS, a second retarded form of TCERG1, migrating at approximately 180 kDa, was detected. This 180 kDa form was unresponsive to NEM (Fig. 1A, lanes 3 and 4). Both the mobility and dependence of these modified TCERG1 species were consistent with potential sumoylation.

We also performed immunoprecipitation experiments under stringent conditions to preserve SUMO conjugates using affinity-purified anti-TCERG1 antibodies. Anti-SUMO-1 monoclonal antibody detected two specific bands in the anti-TCERG1 immunoprecipitates with apparent molecular weights similar to those of the previously identified putative SUMO-TCERG1 species (Supplemental Figure 1A). Those results further support that TCERG1 is a bona-fide target for in vivo SUMO modification.

As mentioned above, the level of sumoylation is very low in most of the target proteins (usually below 5%). To estimate the proportion of modified TCERG1, we carried out semiquantitative experiments in the linear range...
of the standard curve. We found that TCERG1 sumoylation accounts for a small percentage of total protein, approximately <5%, which is in agreement with most published data (Supplemental Figure 1B and data not shown).

Next, we asked if a T7-tagged human TCERG1 cDNA could also be modified and whether SUMO coexpression elicited TCERG1 sumoylation. First, we transfected HEK293T cells with a plasmid encoding T7-tagged TCERG1 and prepared whole cell extracts. Upon western blotting with anti-T7 monoclonal antibody, we observed the expected 150 kDa species of TCERG1. We also observed at least two slower migrating forms with apparent molecular mass of approximately 180 and 200 kDa, consistent with the preceding observations with anti-TCERG1 antibody (Fig. 1B, lane 1, WCE panel; grey and black arrows, respectively). To determine if the modified forms of TCERG1 could be enriched by sumoylation, we independently cotransfected each of the three SUMO isoforms with the T7-tagged TCERG1 into HEK293T cells and probed extracts from these cells (Fig. 1B). Coexpression of SUMO-1 with T7-tagged TCERG1 resulted in an increase of the 180 and 200 kDa species along with a third slower migrating form of apparent molecular size of 300 kDa (Fig. 1B, WCE panel; lane 2, grey and black arrows, and bracket, respectively). Coexpression of SUMO-2 or SUMO-3 with TCERG1 resulted in the appearance of a single slow migrating band of approximately 180 kDa immunoreactive to anti-T7 antibody (Fig. 1B, WCE panel; lanes 3 and 4, grey arrow). Higher molecular weight species, which likely represent polySUMO2/3 chains, were also detectable in most of the experiments (Fig. 1B, bracket).

To confirm that these higher molecular weight species were indeed SUMO-modified TCERG1 proteins, T7-tagged TCERG1 and HA-tagged SUMO-1 and SUMO-2/3 were again coexpressed. TCERG1 was immunoprecipitated from cells with anti-T7 antibody under highly stringent conditions, and then blotted with anti-T7 or anti-HA antibody. Similar higher molecular weight species were detected by those antibodies. Coexpression of SUMO-1 with T7-tagged TCERG1 resulted in an increase of the 180 and 200 kDa species (Fig. 1B, α-T7 IP and α-HA-tag panels; lane 2, grey and black arrows, respectively). Coexpression of SUMO-2 or SUMO-3 with TCERG1 resulted in the appearance of the single slow migrating band of approximately 180 kDa (Fig. 1B, α-T7 IP and α-HA-tag panels; lanes 3 and 4, grey arrow). Higher molecular weight species were also detectable in most of those experiments (Fig. 1B, α-T7 IP and α-HA-tag panels; lanes 2, 3 and 4, bracket). Finally, we analyzed the TCERG1 immunoprecipitates with specific SUMO-1 and SUMO-2 antibodies and found immunoreactivity that was overlapping with the signal obtained with anti-TCERG1 and anti-HA antibodies (Fig. 1B, α-SUMO-1 and α-SUMO-2 panels). Those data further support the results of the previous experiment. Therefore, both endogenous TCERG1 and transiently expressed TCERG1 can be covalently modified by SUMO paralogs in cells.

To map the sumoylation sites within TCERG1, two T7 epitope-tagged fragments encompassing the amino- and carboxyl-terminal halves of human TCERG1 (containing amino acid residues 1-662 and 616-1098, respectively) were assayed in a similar manner as described above. Both fragments retain the putative nuclear localization signal found in the middle of the protein and localize throughout the nucleus. Like wild-type, the TCERG1 carboxyl fragment localizes in the splicing factor-rich nuclear speckles ((25) and data not shown). HEK293T cells were transfected with either the amino- and carboxyl-TCERG1 fragments. SUMO-1, SUMO2, or SUMO-3 were coexpressed with TCERG1 fragments. Only the amino-terminal half of TCERG1 rendered a pattern of bands consistent with that of the full-length protein, either in whole cell extracts or in pellets of immunoprecipitations with anti-T7 antibody (Fig. 1C), whereas no higher molecular weight species were observed with the carboxyl fragment of TCERG1 (Fig. 1D). Thus, the major sumoylation sites reside within the first 662 amino acids of TCERG1.

Because most SUMO substrates interact with the E2 enzyme Ubc9 and this interaction is considered a strong indicator of sumoylation (54), we next determined whether TCERG1 interacts with Ubc9 using the GST pull-down assay. We first purified HA-tagged human Ubc9 recombinant protein from bacteria using native conditions (Figure 1E). The amino and carboxyl fragments of TCERG1 were purified from bacteria as GST fusions by binding to glutathione-agarose beads (Fig 1F, top panel). The bead-bound TCERG1 fragments were then used as baits in pull-down assays with the purified HA-hUbc9. As a negative control, we
used GST and beads alone. After extensive washing, the bead-bound proteins were eluted by boiling the samples with SDS-PAGE loading buffer and analyzed by SDS-PAGE and western blotting to detect HA-tagged hUbc9. The results of the in vitro binding assay revealed a direct interaction between hUbC9 and the amino-terminal 662 amino acids of TCERG1 (Figure 1F, bottom panel), thus supporting the conclusion that this region of TCERG1 contains required sequences for proper interaction with the sumoylation machinery and conjugation with SUMO.

K503 and K608 are the major acceptor sites of SUMO modification in TCERG1. Sumoylation usually takes place on a lysine embedded in the core consensus motif \( \Psi ^x Kx \) E/D, where \( \Psi \) represents a hydrophobic residue and \( x \) stands for any amino acid residue (37). Recent studies have defined an extended consensus motif \( \Psi ^x Kx/\Psi ^x E/D \) \( \Psi \) embedded in the core consensus motif \( \Psi ^x Kx \) E/D (54). This extended motif can be used to correctly predict targets for sumoylation. Two lysines within the amino region of TCERG1, K503 and K608, match this sumoylation motif (Fig. 2A). These lysines and surrounding sequences are highly conserved among TCERG1 homologs in metazoa, as shown in the amino acid alignment in Figure 2A.

To determine if these lysines were modified by sumoylation, we generated T7-tagged TCERG1 constructs containing lysine to arginine mutations at either position K503, K608, or both. We then carried out in vivo sumoylation assays by coexpressing the wild-type and mutated TCERG1 proteins with or without SUMO-1, SUMO-2 or SUMO-3 in transient transfection experiments. Both unmodified and sumoylated TCERG1 were detected by immunoblotting of these cell extracts with T7-specific antibodies (Fig. 2B-D, \( \alpha \)-T7-Tag panels). Coexpression of SUMO-1 with T7-tagged TCERG1 resulted in an increase of the 180 and 200 kDa species (Fig. 2B, lane 2, arrows). Mutation of lysine 503 abrogated the 200 kDa band associated with SUMO-1 conjugation (Fig. 2B, lane 3). Mutation of lysine 608 abrogated the formation of a shifted band corresponding to the faster migrating specie (180 kDa) modified specifically by SUMO-1 (Fig. 2B, lane 4). The K503R and K608R mutants abrogated the formation of the third shifted band of apparent molecular size of 300 kDa corresponding to the minor SUMO-1 modified form (Fig. 2B, lanes 3 and 4, bracket). The 300 kDa form may represent a double SUMO-1 modification of TCERG1 at different lysine residues. Mutations at lysines 503 and 608 caused changes in the sumoylation pattern, and no species associated with SUMO-1 conjugation were observed with the double mutant (Fig. 2B, lane 5), suggesting that both residues are the major sumoylation acceptor sites for SUMO-1 in vivo. The data also show that the carboxyl-terminal region of TCERG1 is poorly, if at all, modified by SUMO in vivo.

Coexpression of SUMO-2 or SUMO-3 with TCERG1 resulted in the appearance of the single slow migrating band of approximately 180 kDa immunoreactive to anti-T7 antibody (Fig. 2C and 2D, lane 2, grey arrow). The polySUMO2/3 chains were also detectable (Fig. 2D, bracket and data not shown). Mutation at lysine 503 and the double mutant abrogated sumoylation by SUMO2 and SUMO-3 (Fig. 2C and D, lanes 3 and 5). The sumoylated forms of the proteins produced by SUMO2/3 in transfected cells were unchanged upon mutating lysine 608 (Figure 2C-D, lane 4), supporting that modification by SUMO-2/3 in vivo occurs preferentially at position 503. Lysines residues can also be sites for ubiquitination, however, our transfection data indicate that the identified sumoylation sites do not affect protein stability and therefore unlikely to be critical for ubiquitination.

To further corroborate those results and to determine which slow mobility species bear SUMO-1 or SUMO-2/3, the HA-immunoreactivity of TCERG1 immunoprecipitates was also analyzed. The same specific bands described above were also detected in TCERG1 immunoprecipitates by using an anti-HA monoclonal antibody (Fig. 2E-G), which further suggest that the lower band of approximately 180 kDa likely corresponds to sumoylation at position 608 and the upper band of approximately 200 kDa to sumoylation at position 503.

Our findings indicate that K503 and K608 are the major SUMO acceptor sites within TCERG1 in vivo. To further confirm that K503 and K608 are SUMO acceptor sites, we carried out in vitro sumoylation assays. As a substrate, we used a GST-tagged TCERG1 fragment (134-662) that contained the major sumoylation sites. Wild-type TCERG1 or K503R, K608R and K503,608R mutant proteins were affinity
purified from bacteria and incubated in the presence of recombinant SUMO-1, and E1, and E2 (Ubc9) enzymes. Sumoylation is ATP dependent and therefore we performed the assay in the absence and presence of ATP. Higher molecular weight species of the amino-terminal domain of TCERG1 containing residues 134-662 were observed only when all components were added in the reaction in the presence of ATP (Fig. 2H, lanes 1 and 2), providing further evidence that the amino-terminal region of TCERG1 is a substrate for sumoylation by SUMO-1. Furthermore, the K503R and K608R mutants abrogated the formation of shifted bands corresponding to a single SUMO-1 molecule conjugated to a specific lysine residue (Fig. 2H, lanes 4 and 6), resembling closely the in vivo pattern shown for SUMO-1. The double mutant could not be sumoylated in vitro (Fig. 2H, lane 8), indicating that those lysines are also preferentially targeted for sumoylation in vitro.

We also carried out in vitro sumoylation experiments with SUMO-2 purified protein. Higher molecular weight species were again observed only when all components were added in the reaction in the presence of ATP (Fig. 2I, lanes 1 and 2). Mutation at position 503 abrogated the formation of the major sumoylated band (Fig. 2I, lanes 3 and 4) while mutation at position 608 eliminated a minor slow migrating band (Fig. 2I, lanes 5 and 6), thus confirming that those lysines are preferential SUMO acceptor sites. Simultaneous mutation of both lysines to arginine again abolished sumoylation (Fig. 2I, lanes 7 and 8). Here, we were able to detect modification of lysine 608 residue by SUMO-2, but still significant differences in the relative conjugation efficiency for each site between SUMO-1 and SUMO-2 were observed, supporting that the modification by SUMO-2 occurs preferentially at K503 (Figure 2I, bar graph).

The nuclear localization of TCERG1 is independent from modification at K503 and K608 by sumoylation. Because sumoylation is known to influence the cellular localization of target proteins, we wished to investigate whether loss of sumoylation at K503 or/and K608 modulates TCERG1 distribution. TCERG1 is distributed in dot-like structures throughout the nucleoplasm and colocalizes with the splicing-factor rich nuclear speckles (25). We expressed wild-type full-length TCERG1 and the lysine-to-arginine mutants tagged with the enhanced cyan fluorescence protein (ECFP) at the amino-terminus and examined their nuclear localization in HEK 293T cells using immunofluorescence microscopy. As shown in Figure 3, all the TCERG1 constructs tested exhibited a similar nucleoplasm distribution with an increased signal in organized granule-like sites and outside the highly compacted chromatin territories detected with the fluorescent dye TOPRO-3 (Fig. 3A). This pattern is similar to that of endogenous TCERG1 (25). TCERG1 also is enriched in the speckles (25); therefore, we investigated the effect of SUMO conjugation at K503 and K608 on the TCERG1 localization to nuclear speckles. Abrogation of SUMO conjugation at those lysines had no effect on the distribution of TCERG1 to nuclear speckles, which were visualized by co-staining with antibody against SC35, a splicing factor that is commonly used to define nuclear speckles (Fig. 3B). We analyzed the spatial relationship between wild-type and mutant TCERG1 variants relative to SC-35 by quantitatively scanning specific nuclear regions containing speckles (Fig. 3B, line scans: ECFP-TCERG1 and SC-35 in blue and red, respectively). We conclude that SUMO modification at amino acid residues K503 and K608 is dispensable for the natural nuclear distribution of TCERG1.

Sumoylation decreases TCERG1 transcriptional activity but does not affect its alternative splicing function. Many proteins involved at different steps of regulation of gene expression are modified by sumoylation. TCERG1 modulates the activity of RNAPII complexes, and appears to play a role in both transcription elongation and pre-mRNA splicing. We sought to determine the functional significance of TCERG1 sumoylation in the context of those processes.

Recent data have shown that TCERG1 modulates fibronectin extra domain I (EDI) exon inclusion (33). The fibronectin EDI minigene displays a weak 3’ splice site followed by a stronger splice site in the downstream intron. As a result, the EDI exon can be alternatively spliced in a regulated fashion; controlled, at least in part, by RNAPII elongation (4,45,50). We wished to confirm those results and use this assay to determine whether SUMO modification regulates the effect of TCERG1 on the alternative processing of the fibronectin EDI exon. Given that changes in promoter structure strongly affect EDI splice site selection (44), we performed the experiments using minigenes carrying different promoters (Fig. 4A).
HEK293T cells were transiently transfected with the minigenes containing the \(\alpha\)-globin, fibronectin, or HSV-thymidine kinase promoter (HSV-TK), and alternative splicing of EDI was assessed by RT-PCR. Inclusion of EDI exon was lower when transcription was driven by the \(\alpha\)-globin promoter compared with the fibronectin promoter (compare Fig. 4B and 4C), as previously demonstrated (44), thus confirming that the regulation of this system is linked to transcriptional control. A similar result was obtained when transcription was driven by the HSV-TK promoter (Fig. 4D). When TCERG1 cDNA was overproduced by cotransfection, EDI exon inclusion was significantly diminished from all three reporter minigenes (Fig. 4B-D), even in the case of a promoter previously reported to be resistant to TCERG1 influence such as HSV-TK (23). In our experiments, TCERG1 did not affect all three reporter minigenes (Fig. 4B-D), even in the case of a promoter previously reported to be resistant to TCERG1 influence such as HSV-TK (23). In our experiments, TCERG1 did not affect the EDI band (Fig. 4B-D). These results suggest a direct effect of TCERG1 on splicing decisions in a manner independent of transcription.

In those experiments, it is clear that for all three promoters, the decrease in EDI transcript is not fully compensated by an increase in the EDI transcript. To determine whether TCERG1 inhibits the overall transcription, we have used a previously described approach that quantitatively analyzes the amount of nascent transcript and total mRNA species derived from the EDI minigene by real-time PCR (50,51). No significant alteration in the level of nascent transcript or total mRNA was observed (Figure 4E). The results further support that TCERG1 influences alternative splicing and does not globally inhibit mRNA synthesis.

The effect of TCERG1 on the alternative splicing of the well-characterized tau minigene system was also determined. Human tau exon 10 is regulated by a stem-loop structure that is formed in the pre-mRNA at the downstream 5' splice site. Normally, exon 10 is included or excluded from the mRNA with roughly equal probability during pre-mRNA splicing. Mutations clustered around the 5' splice site of exon 10 result in predominant inclusion of exon 10, resulting in a severe neurological disorder (46,55). In contrast to the EDI situation, no specific change in the alternative splicing of exon 10 was observed upon TCERG1 overexpression (Fig. 4F-G). The very minor reduction of Tau 10 observed at 4 \(\mu\)g of TCERG1 (Fig. 4G) is not statistically significant. Those results are in agreement with previous observations that link TCERG1 to recognition of specific 3' splice sites of pre-mRNAs (30,56).

Having established that TCERG1 alters the alternative splicing of EDI exon, we investigated the potential functional consequences of the conjugation of SUMO to TCERG1. Cells were cotransfected with the \(\alpha\)-globin promoter construct and plasmids expressing the wild-type TCERG1 and the K503R, K608R, and K503,608R mutant versions. The splicing pattern elicited by the EDI minigene was similarly affected by the expression of the mutant variants that fail to be modified by SUMO (Figure 4H and I) even when the levels of TCERG1 used are moderate (Figure 4I). To exclude a contribution of the endogenous pool of sumoylated TCERG1, which might mask differences between overexpressed wild-type and mutant TCERG1, we examined the effects of overexpressing wild-type and sumoylation deficient TCERG1 in cells that have been previously depleted of endogenous TCERG1 by RNAi. Both wild-type and mutant TCERG1 decreased EDI inclusion the same magnitude. Therefore, SUMO modification does not modulate the ability of TCERG1 to regulate alternative splicing of EDI exon, which rules out a putative effect of endogenous TCERG1 sumoylation. (Supplementary Figure 2).

To investigate whether SUMO modification affects the transcriptional activity of TCERG1, we determined the effects of the single and double mutants (K503R, K608R, and K503,608R) compared with the wild-type protein. For this purpose, we employed a one-hybrid assay to measure TCERG1 transcriptional activity. This experiment uses a reporter construct that contains five copies of the GAL4 consensus DNA-binding site and a minimal adenovirus E1b promoter that drives the expression of luciferase (GAL4E1B-LUC in Fig. 5A). HEK293T cells were cotransfected with the reporter plasmid and the wild-type amino-terminal domain of TCERG1 (134-662) or the mutant variants fused to a putative effect of endogenous pool of sumoylated TCERG1, which might mask differences between overexpressed wild-type and mutant TCERG1, we examined the effects of overexpressing wild-type and sumoylation deficient TCERG1 in cells that have been previously depleted of endogenous TCERG1 by RNAi. Both wild-type and mutant TCERG1 decreased EDI inclusion the same magnitude. Therefore, SUMO modification does not modulate the ability of TCERG1 to regulate alternative splicing of EDI exon, which rules out a putative effect of endogenous TCERG1 sumoylation. (Supplementary Figure 2).

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single mutant (Fig. 5A). Thus, the lysine residue 503 that serves as SUMO-1 and SUMO-2/3 acceptor site appears to negatively regulate TCERG1 transcriptional activity. Expression of the various TCERG1 proteins was nearly identical, suggesting that the enhancement of TCERG1-K503R transactivation capacity was not due to different protein expression level (Fig. 5A). Furthermore, the activities exhibited by the different constructs seem to rely on a specific mechanism since the observed modulation of reporter gene activity was dependent upon GAL4 DNA-binding domain in the effector constructs and the presence of a functional TATA-box sequence in the reporter construct (Fig. 5B and 5C). The observed effects are consistent with the fact that effects of SUMO on transcriptional regulation are generally much more evident at promoters bearing multiple binding sites (57).

To lend additional support to the observation that SUMO modification represses transcriptional activation by TCERG1, we examined the transcriptional activity of the full-length wild-type TCERG1 and the sumoylation-impaired mutants in an independent system that does not imply recruitment through GAL4-DBD. Toward this end, we took advantage of a system to study TCERG1 activated NF-kB-dependent transcription originally developed by the Garcia-Blanco’s laboratory (Duke University) that uses a luciferase-reporter construct consisting of a minimal murine c-fos promoter and three upstream copies of the MHC class I NF-kB binding site termed p3X-kb-L (Fig. 5D), previously described by Mitchell and Sugden (58). Transient overexpression of wild-type TCERG1 in HEK293T cells stimulated the expression from p3X-kb-L reporter plasmid in a dose-dependent manner (Fig. 5D). We next measured the transcriptional response to the mutant versions of TCERG1. Both the K503R and K608R mutants exhibited greater transactivation activity than wild-type TCERG1 (Fig. 5E). This enhancement was not the result of differential TCERG1 expression, as confirmed by western blotting (Fig. 5E). The induction level of p3X-kb-L reporter plasmid by the double mutant (K503,K608R) was similar to that of the single mutants (Fig. 5E). Together, these findings support our previous data and suggest that the SUMO modification sites negatively modulate TCERG1 transcriptional activity.

DISCUSSION

In this study, we report sumoylation as a novel regulator mechanism of TCERG1 function. We demonstrate that TCERG1 is a substrate for SUMO-1 and SUMO2/3 conjugation. We observed that sumoylation occurs within the amino-terminal region of TCERG1 (Fig. 1). Furthermore, the amino region interacts directly with the SUMO E2-conjugating enzyme Ubc9 (Fig. 1F). No additional active sumoylation sites were detected within the carboxyl moiety of TCERG1 in vivo, however we cannot rule out other sites. In fact, several strong putative SUMO acceptor sites are predicted in the carboxyl region of the protein (data not shown). These might reflect other SUMO conjugation pathways regulated in a cell- or environment-specific manner. Consistent with our results, TCERG1 has two motifs that conform to the extended consensus sequence recognized by Ubc9 and are located at the amino region. Here, we provide in vivo and in vitro evidence that human TCERG1 is modified by SUMO-1 and SUMO-2/3 on lysine residues 503 and 608 (Fig. 2) following specific patterns and displaying distinct paralog specificities, which may reflect an additional functional specialization of each acceptor site. In relation to this, we have detected a shift from SUMO-1 to SUMO2/3 conjugation upon heat shock treatment (data not shown). Interestingly, a very recent report describing the heat shock-induced global changes in the sumoylation state of the proteome identified TCERG1 as a target for SUMO-2 modification (59). To note, proteins involved in mRNA transcription and RNA-binding proteins were significantly overrepresented in the SUMO-2 modified proteome (59). In this context, it will be of interest to investigate TCERG1 effects on splicing upon cellular stress and whether it depends on TCERG1 sumoylation.

The algorithms that we used to locate putative SUMO acceptor residues predicted other potential sumoylation sites in the amino region of TCERG1. Among those, a lysine at position 478 significantly scores with the sumoylation consensus sequence. We carried out in vivo and in vitro sumoylation experiments with TCERG1 constructs containing lysine to arginine mutation at position 478 and we found no evidence of SUMO modification on this particular amino acid residue (data not shown).

Having identified sumoylation sites in TCERG1, we investigated its effect on the
protein. One documented function of sumoylation is to control targeting of proteins to different cell compartments. We described previously that TCERG1 accumulates in the splicing factor-rich nuclear speckles and that the FF-domains, which are located at the carboxyl region of the protein, are required for this localization (25). We investigated whether sumoylation of TCERG1 affects its spatial distribution in the nuclei. Our results show that mutant forms of TCERG1 that are not sumoylated localize to the nucleus similarly to the wild-type protein (Fig. 3). Thus, it is unlikely that sumoylation of TCERG1 is required for the spatial distribution of this protein in the cell.

TCERG1 modulates alternative pre-mRNA splicing ((30,33) and Fig. 4) and transcription ((23) and Fig. 5), and a role for this protein in coordinating transcription elongation and pre-mRNA processing has been proposed (25,56). We show here that TCERG1 specifically influences fibronectin EDI splicing independently of the promoter used and without altering the overall levels of full-length transcripts (Fig. 4), which might argue against a coupling mechanism to alter splicing decisions at the EDI exon. Very recently, studies have also reported observations that demonstrate that these two functions can occur independently of each other for regulators that function as transcriptional coactivators that can also regulate alternative splicing (60,61). SUMO addition could influence these processes. Recently, several hnRNPs were found in proteomic studies to be modified by SUMO (62) suggesting a role of sumoylation in regulating mRNA metabolism and perhaps splicing regulation. In support of this hypothesis, the list of spliceosome components that are substrates for SUMO conjugation has been recently extended (35). Moreover, sumoylation alters ADAR1 editing activity and the assembly and activity of the pre-mRNA 3’ processing complex (41,63) emphasizing the importance of this post-translational modification in RNA-processing events. However, no differences were found between wild-type TCERG1 and the mutant forms that are not modified by SUMO in the alternative processing of the fibronectin EDI exon (Fig. 4). Thus, we conclude that post-translational modification of TCERG1 at lysines 503 and 608 by SUMO may not be essential for TCERG1 in mediating these specific alternative splicing decisions, although we cannot rule out potential effects on other mRNAs. In our experiments, TCERG1 did not affect the EDI band (Figure 4B-E). Although this finding remains to be fully investigated, this observation might point towards a role of TCERG1 in the modulation of the spliceosome assembly after exon definition, a poorly understood mechanism for alternative splicing regulation that has been observed recently (64) and that could explain a previously reported intron-retention event with one of the endogenous targets of TCERG1 (33).

We have also taken several approaches to define the consequences of TCERG1 sumoylation for transcription. TCERG1 can activate transcription in cell assays by a mechanism not fully understood but that may include elements that regulate transcriptional control such as P-TEFb (43,65,66). In fact, TCERG1 interacts with protein complexes that are essential for establishing a processive transcription (22,25). Mutating the SUMO acceptor sites enhanced the transactivation capacity of TCERG1. In other examples, sumoylation negatively impacts transcription factor activity (67,68) by multiple mechanisms (38). At least two repressive mechanisms are consistent with low steady-levels of transcription-factor sumoylation (69). First, sumoylation of the transcription factor can lead to the recruitment of repressive factors with chromatin-remodelling activity. If this occurs, a repressive chromatin state will remain even after sumoylation of the transcription factor is lost. Alternatively, sumoylation of a transcription factor can be necessary to initiate the recruitment of inhibitory factors to the promoter. After a stable repressor complex has formed, this complex will remain stable even after SUMO is removed from the initial transcription factor. In both cases, the transcription factor only needs to be sumoylated for a short period of time, during which gene silencing is initiated. Although the precise consequence of sumoylation on TCERG1 function remains to be determined, we speculate that this modification might alter or disrupt protein interactions. Sumoylation may alter TCERG1 association with transcriptional complexes, including RNA polymerase and elongation factors. In fact, structural elements adjacent to the SUMO acceptor motifs participate in interactions with elongation factors and play a role in the transcriptional activity of TCERG1 (25,29). It is tempting to speculate that SUMO modification might serve as a molecular switch to rearrange transcription processing.
complexes. Future work should document that SUMO modification affects the transcript level of an endogenous gene whose expression is directly affected by TCERG1 and clarify the precise role of TCERG1 sumoylation in the coupling between transcription and alternative splicing, the regulation of those processes, and the molecular mechanisms that are involved.

There are indications that TCERG1 is involved in the pathogenesis of Huntington’s disease (70, 71). Given the recent findings implicating sumoylation as an important regulator of aggregation or toxicity of polyglutamine expanded proteins (72, 73), it is tempting to speculate that TCERG1 sumoylation may influence its role in this disease.

Other proteins related to TCERG1 by virtue of their domain architecture (specifically the WW- and FF-domain-containing proteins FBP11, HYPC and FBP21) also possess predicted sumoylation target sites. These putative sumoylation sites match the expanded consensus motif for sumoylation NDSM and are structurally located in a similar position as K503 and K608 in TCERG1 (data not shown). Whether those proteins, which like TCERG1 are involved in transcription and mRNA processing, are regulated by sumoylation remains to be determined.

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The abbreviations used are: TCERG1, transcription elongation regulator 1; RNAPII, RNA polymerase II; CTD, carboxyl-terminal domain; SUMO, small ubiquitin-like modifier; EDI, fibronectin extra domain I.

FIGURE LEGENDS

Fig. 1. TCERG1 is a sumoylation substrate in vivo and interacts directly with human Ubc9. (A) Western blotting analyses of endogenous TCERG1 from HeLa cells with or without 25 nM NEM or/and 1% SDS. Specific antibodies against TCERG1 and cyclin T1 (as a loading control) were used to localize the proteins. The bracket indicates the positions of the two predominant slow migrating bands in the samples treated with 25 nM NEM and 1% SDS buffer. Molecular masses in kDa are indicated to the left.

(B) HeLa cells were cotransfected with a plasmid encoding T7-tagged TCERG1 along with empty vector (lane 1) or vectors encoding HA-SUMO-1 (lane 2), HA-SUMO-2 (lane 3), or HA-SUMO-3 (lane 4). Whole-cell lysates (WCE) were directly analyzed by western blotting or subjected to immunoprecipitation with T7-specific antibodies followed by SDS-PAGE and western
blotting with anti-T7, anti-HA, anti-SUMO-1, anti-SUMO-2, or anti-cyclin T1 antibodies. Specific higher-molecular-weight forms of TCERG1 are visible above the position of the 175 kDa marker, and are indicated by arrows and bracket (see description in text). The unmodified form of TCERG1 is indicated by a close arrowhead. (C and D) The same experiment described for panel B was repeated but using plasmids encoding T7-tagged amino (C) and carboxyl (D) regions of TCERG1. *, nonspecific band present in WCE and reactive to anti-T7 antibodies. **, immunoglobulin fraction eluted from the matrix. (E) Purification of HA-tagged Ubc9. Lanes 1 and 2: crude E. coli lysates with (+) or without (-) IPTG; lane 3: elution fraction after thrombin cleavage (HA-hUbc9). (F) The amino region of TCERG1 interacts directly with human Ubc9. HA-tagged Ubc9 was incubated with equal amounts of GST or GST-TCERG1 (234-662) and GST-TCERG1 (631-1098) bound to glutathione Sepharose beads. Bound proteins (lanes 2-5) were eluted with SDS-PAGE loading buffer and resolved in a 15% SDS-PAGE gel along a sample of the input (lane 1). The upper part of the gel was stained with Coomassie Blue R250 to visualize the eluted GST-fusion proteins and the lower part was transferred to a membrane and incubated with an anti-HA-specific antibody to detect ubc9. Molecular masses (M) in kDa are indicated to the left.

Fig. 2. K503 and K608 are the major sites of sumoylation of TCERG1 and are conserved among species. (A) Panel shows a CLUSTALW amino acid alignment of TCERG1 homologs that include the putative SUMO acceptor motifs. Conserved acceptor lysines within the core consensus motif (in yellow) and NDSM consensus motif (in blue) features are highlighted. Hs, Homo sapiens; Mm, Mus musculus; Gg, Gallus gallus; Xl, Xenopus laevis; Dr, Danio rerio; Sm, Schistosoma mansoni; Ae, Aedes aegypti. (B-D) HEK293T cells were cotransfected with expression constructs of HA-SUMO-1 (A), HA-SUMO-2 (B) or HA-SUMO-3 (D) and T7-tagged wild-type (WT) TCERG1 or the indicated mutants, and lysates subjected to immunoblotting with anti-T7 and anti-Cyclin T1 antibodies. The sumoylated forms of TCERG1 are indicated by arrows (see description in text). The unmodified form of TCERG1 is indicated by a close arrowhead. (E-G) HEK293T cells were cotransfected with expression constructs of HA-SUMO-1 (E), HA-SUMO-2 (F) or HA-SUMO-3 (G) and T7-tagged WT TCERG1 or the indicated mutants, and lysates subjected to immunoprecipitation with anti-T7 antibodies and western blotting analysis with anti-HA antibodies. The sumoylated forms of TCERG1 are indicated by arrows. (H) In vitro-expressed GST-TCERG1 (134-662) fusion protein, either WT or the indicated mutants, were incubated with SUMO-1 reaction components and the reaction products were analyzed by SDS-PAGE and immunoblotting with specific antibodies against GST. The bands corresponding to the SUMO-conjugated forms of TCERG1 (134-662) is indicated by arrows. The unmodified form of recombinant TCERG1 is indicated by a close arrowhead. (I) The same experiment described in panel H was carried out with SUMO-2. Quantification of the experimental data obtained with the double mutant variant from two independent experiments were quantified and are shown in graphic form at the right of the panel.

Fig. 3. The nuclear localization of TCERG1 is independent from modification at K503 and K608 by sumoylation. (A) Immunofluorescence analysis of HEK293T cells transfected with ECFP-TCERG1-WT or the indicated mutant constructs (ECFP, blue). TOPRO-3 labelling was used to stain chromatin (red). Individual and merge images are shown. (B) Colocalization of the indicated TCERG1 constructs with the essential splicing factor SC35 that commonly serves to define nuclear speckles. Dual labeling of cells with either ECFP-TCERG-WT or SUMO mutants (ECFP, blue) and with the anti-SC35 antibody (SC-35, red) was performed. Line scans showing local intensity distributions of TCERG1 and of SC35 are shown to the right of the panels. Bars indicate the position of the line scans.

Fig. 4. Effect of wild-type TCERG1 and sumoylation mutants on EDI alternative splicing. (A) Schemes of the minigenes carrying the different promoters. Exons are represented by boxes and introns by lines. Alternative EDI isoforms generated by inclusion (EDI+, 500 bp) or skipping (EDI-, 230 bp) of EDI exon are indicated. Also indicated are the position of the primers used to amplify the mRNA splicing variants. α-globin, human alpha-globin; mFN, human fibronectin; HSV-TK, herpes simplex virus-thymidin kinase. (B-D) Effect of TCERG1 overexpression on alternative splicing elicited by minigenes carrying the α-globin (A), fibronectin (B), and HSV-TK (C) promoters. RNA splicing variants were detected by radioactive RT-PCR and the products of amplification were
separated by polyacrylamide gels. β-actin was amplified as a control. A sample of cell lysate was immunoblotted with anti-TCERG1 antibody to demonstrate the expression levels. Ratios between radioactivity in EDI+ bands and EDI- bands from two independent experiments are shown below the panels. (E) Quantitative analysis of the amount of nascent transcript and total mRNA species derived from the α-globin EDI minigene upon cotransfecting with wild-type TCERG1 by real-time PCR. Quantification of the experimental data from four independent experiments is shown in graphic form. (F) Schematic representation of the tau exon 10 minigene system. Alternative spliced mRNAs containing tau exon 10 (Tau 10+, 245 bp) and mRNAs with tau exon 10 skipped (Tau 10-, 153 bp) are shown in the figure. (G) Effect of TCERG1 overexpression in the regulation of tau exon 10. RT-PCR analysis of the cotransfections of tau 10 minigene with TCERG1. β-actin was amplified as a control. A sample of cell lysate was immunoblotted with anti-TCERG1 antibody to demonstrate the expression levels. Ratios between radioactivity in tau 10+ bands and tau- bands from two independent experiments are shown below the panel. (H) Cotransfection assays were carried out with wild type and sumoylation mutants of TCERG1 and the α-globin EDI minigene. Patterns of EDI minigene splicing were analyzed as in previous panels. (I) Curve dose-response for wild-type (WT) TCERG1 and double mutant variant (K503,608R) on EDI alternative splicing. The α-globin EDI minigene and the indicated TCERG1 constructs were cotransfected at 100 (lanes 2 and 4) and 500 (lanes 3 and 5) ng. RNA splicing variants were detected by radioactive RT-PCR and the products of amplification were separated by polyacrylamide gels (top panel). Experimental data from two independent experiments were quantified and are shown in graphic form (middle panel). A sample of cell lysates were immunoblotted with anti-TCERG1 antibody to demonstrate the expression levels (bottom panel).
Figure 3.
Figure 4.
Figure 5.
