Inhibition of G₁ to S Phase Progression by a Novel Zinc Finger Protein P58\(^{\text{TFL}}\) at P-bodies

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Abstract

We recently reported the translocation of the immunoglobulin (lg) light chain \(\kappa\) locus gene with a possible tumor suppressor gene, \(\text{TFL}\), in transformed follicular lymphoma. However, the functional significance in cell transformation remains to be elucidated. Here, we first identified two gene products, \(\text{P58}^{\text{TFL}}\) and \(\text{P36}^{\text{TFL}}\), derived by alternative splicing. The expression was prominent in normal human lymphocytes but defective in some leukemia/lymphoma cell lines. Overexpression of either protein in a mouse pro-B cell line, Ba/F3, and a human leukemia cell line, Jurkat, inhibited \(\text{G}_1\) to \(\text{S}\) phase progression through suppression of retinoblastoma protein (Rb) phosphorylation. The dominant gene product, \(\text{P58}^{\text{TFL}}\), colocalized with mRNA-processing body markers, eukaryotic translation initiation factor 2C and DCP1 decapping-enzyme homolog A, but not with a stress granule marker, T-cell intracellular antigen 1, in the cytoplasm. Taken together with the unique CCCH-type zinc finger motif, the present study suggests that \(\text{P58}^{\text{TFL}}\) could play an important role in the regulation of cell growth through posttranscriptional modification of cell cycle regulators, at least partially, upstream of Rb. (Mol Cancer Res 2009;7(6):880–9)

Introduction

Cytogenetic abnormalities contribute not only to cancer initiation but also to tumor progression for many types of malignancies derived from hematopoietic, mesenchymal, and epithelial tissues (1-5). Neither the \(\text{BCL}-2\) nor \(\text{BCL}-2\)-positive follicular lymphoma in aggressive lymphoma, many cytogenetic abnormalities have been reported (10). Among them, loss of regions in the long arm of chromosome 6 (6q), which is frequently observed not only in hematologic but also in epithelial malignancies (11, 12), has been reported to be associated with poor prognosis (13, 14). These findings suggest the existence of an as yet uncharacterized tumor suppressor gene on 6q21-25.

To pave the way for identification of tumor suppressor gene candidates, microarrays such as array based-comparative genomic hybridization, which can visualize gene amplification and/or other defects in cancer, have been used by several groups around the world. Indeed, loss of 6q21-q27 was reported to be frequent in B-cell lymphoma (15, 16).

Alternatively, a rare cytogenetic translocation sometimes reveals a novel oncogene or cryptic mutation important for tumorigenesis or leukemogenesis. For example, the tumor suppressor gene \(\text{HACE1}\) was first identified from a t(6;15) translocation in a Wilms’ tumor sample. Thereafter, \(\text{HACE1}\) was shown to be involved in multiple cancers (17, 18). The universal cryptic defect of the \(\alpha\)-type platelet-derived growth factor receptor gene (\(\alpha\text{PDGF}\)) in chronic eosinophilic leukemia was also first identified from a rare case with \(t(1;4)\) (q44;ql2) (ref. 19). Thereafter, the genetic change established a new clinical entity defined in the updated WHO classification (20).

Recently, we identified a possible tumor suppressor gene at the break points of the t(2;6)(p12;q25) translocation in a transformed follicular lymphoma, designated \(\text{TFL}\) (21, 22). The break point was located at the putative second intron of \(\text{ZC3H12D}\), an orphan gene previously defined by the human genome project (GenBank accession no. NM 207360). The gene presumably encodes a single “Cys-x8-Cys-x5-Cys-x3-His” (CCCH)-type zinc finger protein, and was recently reported to be a novel tumor suppressor gene candidate. Its loss of heterozygosity was found in sporadic lung cancer, and a single-nucleotide polymorphism at codon 106 was shown to affect tumor growth (23).

However, the \(\text{TFL}\) gene structure, its gene products, or their physiologic function has not yet been determined. Reports of the exon organization of \(\text{ZC3H12D}\) transcripts have been frequently rewritten on the web site of National Center for Biotechnology Information (NCBI). To clarify the functional significance of this tumor suppressor gene candidate in cell transformation, we investigated the \(\text{TFL}\) gene products and their expression profiles and biological function.
Results

Identification of Two Alternatively Spliced Transcripts

At the beginning of our cDNA cloning, two independent genes, ZC3H12D and “an orphan gene encoding FLJ00361 protein (accession #AK090441),” were listed at the NCBI database. RNA blot analysis using a DNA fragment of the TFL gene cloned from a genomic library constructed from a patient’s lymph node with t(2;6)(p12;q25) (ref. 22) and a 5.5-kb cDNA clone, FLJ00361, provided by KAZUSA DNA Research Institute (Tsukuba, Japan), supported the possibility that TFL1 and TFL2 encode 58TFL and P36TFL, respectively. P58TFL and P36TFL had a common amino-terminal sequence (white boxes) including a CCCH-type zinc finger domain (dotted box) and specific carboxyl-terminal sequences (white box with vertical or horizontal lines) and a Pro-rich domain (checked box).

Expression Profiles in Normal Tissues and Leukemic Cell Lines

We first examined mRNA expression in human blood to gain insight about the functional significance of TFL in malignant lymphoma (Fig. 2A). The 5-kb TFL transcript was abundantly expressed in human lymph nodes as well as in normal peripheral T and B lymphocytes. In contrast, human bone marrow expressed less of this transcript. Human B-cell (Daudi and Nalm6) and some myeloid leukemia/lymphoma cell lines (THP-1) expressed TFL as well as did lymph nodes. However, it was less abundant in another myeloid leukemic cell line, HL60. In contrast, mRNA expression in T-cell leukemia/lymphoma cell lines (Jurkat, Molt14, and CCRF/CEM) was undetectable. A smaller transcript (~3.2 kb) was also seen in THP-1 and Daudi cells, which was compatible in size with our cDNA clone TFL1.

We next examined mRNA expression in other normal tissues. The 5-kb transcript was prominently expressed in mouse lymphoid organs, spleen, and thymus (Fig. 2B, left). The mRNA was also detectable in lung, stomach, and kidney, whereas it was undetectable in brain, heart, liver, and muscle. Mouse TFL was also expressed in both splenic T and B lymphocytes (Fig. 2B, right). Although the transcript was not detectable in mouse bone marrow, a faint expression was seen not only in the mouse pro-B-cell line Ba/F3 but also in the myeloid progenitor cell line 32D.

These results indicate that TFL is predominantly expressed in normal lymphoid tissues and that some lymphoid malignancies lack expression.

Identification of the Major Protein Expressed in Human Blood

We next confirmed the expression of TFL-encoded protein by developing polyclonal antisera against recombinant protein derived from a TFL2 cDNA expression vector. As shown in Fig. 1, the first 298 amino acids encoded by TFL1 are identical to those of TFL2. Thus, the antisera could recognize a 58-kDa protein, P58TFL, in Ba/F3 cells transfected with a TFL1 expression vector (Fig. 3A). P58TFL was preferentially expressed in human peripheral mononuclear cells (MNC) but not in polymorphonuclear cells (PMN). The expression pattern in human
hematopoietic cell lines was completely consistent with the results of RNA blot analyses (Figs. 2A and 3A).

To further confirm the specificity of the antisera, we performed immunoblot analysis using immunoprecipitates. The P58TFL was specifically immunoprecipitated, but a nonspecific signal (indicated by a star just above P58TFL) that was detected in cell lysates was absent (Fig. 3). With regard to protein encoded by TFL2, the antisera could recognize recombinant P36TFL expressed in a mammalian cell line (Fig. 3B). Both molecular weights (MW) observed experimentally by SDS-PAGE analysis were compatible with those predicted from their primary structures. P36TFL seems to be similar to a provisionally named p34, although there is a four-amino-acid difference from codon 298 to codon 301 (P34, Pro-Val-Leu-Pro; P36TFL, Leu-Gly-Val-Arg, respectively) by possible alternative splicing of exons 6a and 6b (23). However, the endogenous P36TFL was...
not evident by immunoblot using whole-cell lysates. Some diffuse bands that comigrated with the recombinant P36<sub>TFL</sub> in MNC, HL60, and THP-1 immunoprecipitates are not likely to represent endogenous P36<sub>TFL</sub> because the recombinant P36<sub>TFL</sub> was tagged with hemagglutinin (HA). Thus, the major protein expressed in normal human lymphocytes was P58<sub>TFL</sub>. Moreover, aberrant expression of P58<sub>TFL</sub> was found in some human leukemic cell lines.

**TFL Inhibits DNA Synthesis and Progression to S Phase**

To clarify the biological function of TFL, a green fluorescent protein (GFP)-TFL fusion protein was expressed in an interleukin 3 (IL-3)--dependent mouse normal pro-B-cell line, Ba/F3, where the TFL mRNA expression level was less than that of mature B lymphocytes (Fig. 2B). Interestingly, although the numbers of cells expressing GFP-TFL1 or GFP-TFL2 were equivalent to the number expressing GFP alone at 4 hours after electroporation, the number of cells expressing either TFL1 or TFL2 was drastically decreased by 48 hours (Fig. 4A). We speculated that this might be due to TFL-induced growth arrest and/or cell death because the population of nontransfected Ba/F3 cells soon overwhelmed that of TFL transfectants.

To examine whether TFL could suppress DNA synthesis, we performed bromodeoxyuridine (BrdUrd) assays under G0-G1 synchronization conditions as described in Materials and Methods. As expected, the number of S-phase cells incorporating BrdUrd was decreased by introducing TFL1 or TFL2 cDNA (Fig. 4B). The number of G0-G1-phase cells among transfectants was increased proportionately. These results indicate that the G1 to S phase progression was inhibited by TFL. Inhibition of DNA synthesis by TFL was also confirmed by tritiated thymidine uptake (data not shown). The observation that growth inhibition was observed after introducing either TFL1 or TFL2 cDNAs suggests that a common protein domain between P58<sub>TFL</sub> and P36<sub>TFL</sub> might play a crucial role in cell growth control.

**TFL Suppresses Phosphorylation of Retinoblastoma Protein**

Cell cycle progression is regulated by many signaling molecules. Phosphorylation of retinoblastoma protein (Rb) plays a key role in the G1 to S phase transition (24). To determine whether TFL could inhibit Rb phosphorylation or not, we first assessed Rb phosphorylation status in Ba/F3 transfectants by intracellular flow cytometry assay (Fig. 5A). As expected, introduction of either TFL1 or TFL2 cDNA significantly restrained Rb phosphorylation compared with a control vector alone (mock). These results indicated that TFL inhibited cell cycle progression into S phase through the suppression of Rb phosphorylation. To confirm this finding in another cell line, NIH3T3 cells were transfected with the GFP-tagged TFL vectors. Rb phosphorylation was then assessed by immunofluorescence staining using a specific antibody for phospho-Rb. Cells expressing GFP-TFL1 fusion protein completely lacked Rb phosphorylation.

![Figure 4](image-url)

**Figure 4.** TFL interfered with progression to S phase. **A.** Ba/F3 cells were nucleofected with vectors containing GFP-tagged mock or TFL1 or TFL2 cDNA. GFP-positive cells were analyzed by flow cytometry at 4, 24, and 48 h after electroporation. Left, representative cytograms at 4 and 48 h. The bar graph shows time-dependent decreases of GFP-TFL--positive cells. Columns, mean surviving cells (% of control) compared with mock transfectants from three experiments; bars, SD. **B.** Representatives of three independent BrdUrd assays using mock, TFL1, and TFL2 cDNA transfectants (left). After IL-3 starvation for 16 h, Ba/F3 transfectants were cultured in the presence of IL-3 for an additional 16 h. Then, cells were assessed for BrdUrd incorporations. The bar graph shows percentages of gated cells in G0-G1, S, or G2-M phase. Columns, mean of triplicate samples; bars, SD.
TFL Promotes Caspase-3-Dependent Apoptosis

Apoptotic pathways are also important for the development of cancers. To examine the effect of TFL on cell apoptosis, Ba/F3 cells carrying TFL vectors were stained with Annexin V. As shown in Fig. 6A, both types of TFL increased the number of Annexin V–positive cells by 24 hours after electroporation. To confirm whether cell death was caspase-3 dependent or not, we also studied the activation of caspase-3 by flow cytometry. Ba/F3 cells carrying TFL1 as well as TFL2 vectors had significantly increased cleaved caspase-3 compared with mock transfectants (Fig. 6B). These results suggest that both P58TFL and P36TFL induced caspase-3–dependent apoptosis.

Overexpression of TFL Alters the Carcinogenic Properties of Human Leukemia Cells

It is interesting that the predominant lymphocytic splice form, TFL1, was not expressed in certain human leukemia cell lines as described above (Fig. 3A). To examine whether the deficiency is only correlative or causative for leukemogenesis, we tried some functional assays by introducing the TFL cDNA expression vectors into the T-cell leukemia cell line Jurkat.

Overexpression of TFL1 as well as TFL2 inhibited the growth of Jurkat cells (Fig. 7A). Moreover, TFL vectors suppressed BrdUrd incorporation and Rb phosphorylation, and it increased cleaved caspase-3–positive apoptotic cells (Fig. 7B–D). These results indicate that TFLs serve as a tumor suppressor in certain human leukemia cells.

Differential Subcellular Localization of P58TFL and P36TFL

We also examined the subcellular localization of P58TFL and P36TFL (Fig. 8A). P58TFL localized as discrete granules in the cytoplasm, but not at all in the nuclei, of HeLa cells. On the other hand, P36TFL localized to both cytoplasm and nuclei (but not nucleoli); nuclei were stained more intensely than the cytoplasm. The distinctive localization of P58TFL and P36TFL was confirmed in NIH3T3 and Ba/F3 transfectants (data not shown). There are some differences in the cytoplasmic terminal sequences between P58TFL and P36TFL (Fig. 1). These differences may contribute to specific subcellular localization through as yet unidentified sequence motifs present within this region.

TFL has a single CCCH-type zinc finger motif that binds nucleic acids including RNA. mRNA processing bodies (P-bodies), where mRNA degradation or stabilization occurs, are localized to the cytoplasm as discrete granules. We predicted that P58TFL granules might colocalize with P-bodies. Thus, we cotransfected GFP-tagged P58TFL with HA-tagged P-body markers, eukaryotic translation initiation factor 2C (EIF2C2; also known as AGO2; ref. 25) or DCP1 decapping enzyme homolog A (DCP1A; ref. 26). Almost complete colocalization of P58TFL and EIF2C2 was confirmed in NIH3T3 and Ba/F3 transfectants (data not shown). There are some differences in the carboxyterminal sequences between P58TFL and P36TFL (Fig. 1). These differences may contribute to specific subcellular localization through as yet unidentified sequence motifs present within this region.

Both P-bodies and stress granules are distinct cytoplasmic aggregates involved in mRNA storage and/or degradation, but they are spatially, compositionally, and functionally linked (27, 28). It is possible that P58TFL might be also accumulating in foci near P-bodies much like stress granules. Therefore, we further examined the subcellular localization by cotransfection with a HA-tagged stress granule marker, T-cell intracellular antigen-1 (TIA-1; Fig. 8B, top). Most P58TFL and TIA-1 granules are distinctively localized in the cytoplasm. A few P58TFL granules were juxtaposed with TIA-1 granules, but the two granules never overlapped. Interestingly, overexpression of mouse TFL in HeLa cells promoted the fusion of endogenous TIA-1–positive stress granules with TFL granules. Some stress granules were incompletely engulfed by TFL (Supplementary Fig. S1). In contrast, other granules such as endosomes or lysosomes were not colocalized with P58TFL granules, as assessed with anti-EA1 antibody and LysoTracker, respectively (refs. 29, 30; data not shown).

FIGURE 5. TFL suppressed Rb phosphorylation. A, Phosphorylation levels of Rb in GFP-positive Ba/F3 cells were analyzed by intracellular flow cytometry using Cy5-conjugated phospho-Rb–specific antibody. Dashed lines, isotype control. * P < 0.01, phospho-Rb–positive cells in TFL1 and TFL2 transfectants compared with mock transfectants. Results are expressed as mean ± SD of triplicate samples. B, Immunofluorescence staining of phospho-Rb for NIH3T3 cells transfected with designated GFP fusion vectors. Almost all Rb proteins of control transfectants (Mock GFP–positive cells indicated by arrows) were phosphorylated as shown by yellow nuclei in a merged panel, whereas TFL1 and TFL2 transfectants indicated by arrows showed less Rb phosphorylation as shown by faint red nuclei in Rb-Cy5 staining in both transfectants. There are no yellow signals in the merged image of TFL2.
P58TFL Regulates Cell Growth at P-body

FIGURE 6. Apoptosis induced by TFL. A. Ba/F3 transfectants of mock, GFP-TFL1, and GFP-TFL2 cDNA vectors were stained with Annexin V-Cy5 at 24 h after transfection. *, P < 0.01. Annexin V-positive cells were significantly increased in GFP-positive TFL1 and TFL2 transfectants versus mock. Results are expressed as mean ± SD of triplicate samples. B. Representatives cleaved caspase-3-positive cell analyses. *, P < 0.01, percentages of active caspase-3-positive cells in GFP-positive TFL1 or TFL2 transfectants compared with that in mock transfectants (GFP alone). Results are expressed as mean ± SD of triplicate samples.

These results suggest that P58TFL may be a component of the P-body, which spatially interacts with stress granules and might suppress G1 to S phase progression through posttranscriptional modification of an indispensable cell cycle regulator.

Discussion

ZC3H12D belongs to a gene family including three members, ZC3H12A, ZC3H12B, and ZC3H12C. Among them, monocyte chemoattractant protein-1-induced (MCPIP) was first identified as a product of ZC3H12A in monocyte chemoattractant protein-1-activated monocytes (31). ZC3H12B and ZC3H12C mRNAs were expressed in normal tissues with unique expression profiles, but their expression at the protein level has not yet been confirmed. Here, we first identified two ZC3H12D-derived splicing variants encoding possible tumor suppressors, P58TFL and P36TFL. In contrast to the dominant expression of ZC3H12A in macrophages, ZC3H12D was expressed preferentially on normal lymphoid tissues. In particular, it was very strongly expressed on mature T and B lymphocytes rather than immature thymocytes. Whereas TFL was undetectable in some immature hematopoietic cell lines such as Jurkat, MOLT-14, and CCRF/CEM, the mature adult T-cell leukemia cell line HUT102 expressed it intensively (Fig. 2A). MCPIP is inducible in macrophages by inflammatory stimuli, such as lipopolysaccharide, and regulates proinflammatory activation of macrophages by a negative feedback loop (32). A prototype CCCH-type zinc finger protein, tristetraprolin, encoded by the Zfp36 gene, also regulates tumor necrosis factor α in macrophages through a negative feedback loop (33). Therefore, TFL proteins might play some feedback roles in lymphocyte activation as well as maturation. It is of interest how P58TFL and P36TFL are regulated under T-cell receptor or Ig stimulus.

We originally identified the TFL gene ZC3H12D by cloning the breakpoint of a t(2;6)(p12;q25) translocation in a BCL-2–positive transformed follicular lymphoma (22). The 5′ region of ZC3H12D including exon 1 and exon 2 was replaced by the Vκ and Cκ regions of the Igκ gene. Originally, another 6q allele had been lost in the follicular lymphoma before the transformation. However, it remains to be elucidated whether deregulation of the gene or deletion is relevant to transformation. The growth inhibitory activities of P58TFL and P36TFL on a human leukemia cell line shown here support the hypothesis that loss of the biological function by translocation is implicated in transformation as a tumor suppressor. Deregulated antiapoptotic protein BCL-2, preventing cellular apoptosis in follicular lymphoma, causes massive lymphoid hyperplasia, a premalignant state (34, 35). A secondary hit at ZC3H12D in lymphocytes might result in complete loss of their normal function, facilitating a release from the growth control into tumor progression. Moreover, loss of heterozygosity of ZC3H12D was found in sporadic lung cancer, whereas normal lung expressed the mRNA. By using a human lung carcinoma cell line, Wang et al. (23) showed that a single amino acid change due to single-nucleotide polymorphism contributes to tumor progression. However, there was no significant difference in the growth inhibition between mouse pro-B-cell lines overexpressing TFL with an exchanged codon at 106, from lysine to arginine, according to the single-nucleotide polymorphism (data not shown). Whether this discrepancy is due to the cell lines used or reflects different tissue specificity is not yet clear. In addition, the four-amino-acid difference between p34 and P36TFL might affect biological activity.

Suppression of Rb phosphorylation is thought to precede G0-G1 arrest in P58TFL and P36TFL transfectants. Activated cyclin-dependent kinase phosphorylates multiple sites in Rb, and phosphorylated Rb up-regulates cell growth–related proteins (36). Microarray screening of the expression profiles of cell cycle–related genes in the transfectants revealed the down-regulation of PCNA, E2F3, and E2F4, as expected. Just as MCPIP could contribute to the suppression of tissue inflammation, TFL also might prevent from excessive lymphocyte proliferation following tissue damage, an origin of cancer development.

MCPIP is reported to induce apoptosis in monocytes as well as in cardiac myocytes (37). In addition to G0-G1 arrest, caspase-3–dependent apoptosis was confirmed in Ba/F3 cells overexpressing P58TFL or P36TFL. There are two possibilities about how apoptosis was induced in the transfectants. First, TFL may directly promote apoptosis through a different signaling pathway from that inducing G0-G1 arrest. Alternatively, it may be possible that cell cycle arrest caused apoptosis in the...
transfectants. It is meaningful to determine whether or not the apoptosis is induced in a p53-dependent manner because p53 is involved in cell cycle arrest as well as apoptosis. Apoptosis is characterized by chromatin condensation, internucleosomal degradation of the DNA, cell shrinkage, and disassembly into membrane-enclosed vesicles as a consequence of caspase-3 activation (38). Because apoptosis is generally accompanied by a reduction in cell volume, this parameter was used to discriminate between morphologically apoptotic and viable cell populations (39). Thus, we examined cell cycles as well as Rb phosphorylation by discerning such shrinkage, so-called morphologically apoptotic cells, with flow cytometry. Indeed, cleaved caspase-3-positive cells were not increased in morphologically viable cell populations at all (data not shown). In addition, the number of apoptotic cells accumulated in a time-dependent manner. These results suggest that G0-G1 arrest preceded apoptosis in Ba/F3 transfectants.

Thus far, no differences in the biological functions of P58TFL and P36TFL have been found. However, their subcellular distributions were distinctive. P58TFL was shown to be colocalized with P-body–related proteins, EIF2C2 and DCP1A. Posttranscriptional mechanisms are gradually becoming better defined and are gaining attention (40). For example, EIF2C2 is a major RNA-induced silencing complex component where microRNA as well as short interference RNA is processed. RNA-induced silencing complex components play an important role for posttranscriptional pathways (25, 41). In addition to mRNA processing, sites for RNA editing are very unique. When cellular stress triggers mRNA processing, mRNA and RNA binding proteins are gathered in cytoplasmic granules called P-bodies and stress granules (42). The tandem CCCH-type zinc finger domain originally identified in tristetraprolin binds to AU-rich elements, regulating mRNA stability (33). A similar zinc finger domain–containing protein, Roquin, was also shown to be an RNA-binding protein localized in P-bodies (43, 44). Recent surveys indicate that posttranscriptional pathways and mRNA processing pathways are indispensable to the regulation of protein expression as well as its deregulation in tumorigenesis. With this in mind, P58TFL might bind to certain RNA AU-rich elements and be involved in posttranscriptional modification. More comprehensive assays will be required to identify target molecules of P58TFL and define its role as an RNA binding protein.

In contrast, MCPIP is thought to work as a transcription factor (45). Similarly, P36TFL contains the same zinc finger domain and might be a transcriptional factor because of its unique localization in the nucleus but not nucleoli. The identification of molecular targets of P58TFL and P36TFL would help address the question of how these proteins control cell cycle and apoptosis.

The data presented here show that P58TFL and P36TFL regulate cell growth likely by suppressing Rb phosphorylation. Although future studies are needed to ascertain whether the deregulation P58TFL and/or P36TFL is actually involved...
in tumor progression in vivo, it is of interest to study their potential utility as diagnostic as well as potential therapeutic targets in various malignancies.

Materials and Methods

**Human cDNA Cloning**

mRNAs were extracted from human lymph nodes with Trizol (Invitrogen). After DNase I digestion, cDNA was prepared using PrimeScript RTase (Takara, Japan). PCR was done with high-fidelity PrimeSTAR HS DNA Polymerase (Takara) using several primer sets designed using the NCBI database. Sequencing of PCR products was done using an ABI PRISM 310 Genetic Analyzer (Applied Biosystems). PCR using just one primer set showed several bands, and sequence analyses showed at least two patterns of splice variants. To clone the TFL1 and TFL2 human cDNAs into expression vectors, the following primer sets were used: a common forward primer, 5'-ATGGAGCACCCCAGCAAGAT-3', and reverse primers for TFL1, 5'-TTAGGGCTTGCCCAGGGGCGCCC-3', and TFL2, 5'-CTAGGGCGGTGTTCGCCCCG-3'. To confirm the 3'UTR, we also screened a cDNA library from human peripheral blood (Clontech).

**FIGURE 8.** Distinct subcellular localization of P58TFL and P36TFL. A. Each expression vector of GFP alone or GFP-fused P58TFL or P36TFL was transfected into HeLa cells. Cell nuclei are stained with 4',6-diamidino-2-phenylindole (DAPI). B. HA-tagged EIF2C2, DCP1A, or TIA-1 vector was cotransfected with a GFP-fused P58TFL vector into HeLa cells. HA-tagged P-body and stress granule markers were visualized with Cy5-conjugated antibody.
PCR using total RNA of splenocytes and a forward primer, mouse cDNA was ligated into the reverse primer, 5′cDNAs including complete ORF were ligated into the TIA-1 HA-tagged vector described above using the 5′ vectors were produced by linker ligation of oligonucleotides, 600-bp mouse cDNA probe, which includes the homologous refluoride. For immunoprecipitation, total cell lysates were treated with 1 mmol/L phenylmethylsulfonyl fluoride. For immunoprecipitation, total cell lysates were washed thrice with lysis buffer and thrice with PBS.

Human and mouse T and B lymphocytes (MNC) were isolated by Dynabeads using specific monoclonal antibodies (Dynal Biochem). Mononuclear and polymorphonuclear cells were isolated by Lympholyte-H (Cedarlane) and Histopaque-1119 (Sigma).

RNA Blot Analyses
RNA blots were hybridized with 32P-labeled human and mouse TFL CDNA probes. A 963-bp human TFL probe, which includes the complete ORF of TFL2, was amplified using a forward primer, 5′-ATGGAGCACCACCCAGCAAGATGGAATTTC-3′, and a reverse primer, 5′-CTAACCACATTAGGCAATGCTGC-3′. A 600-bp mouse cDNA probe, which includes the homologous region of human TFL2 cDNA ORF, was amplified from mouse spleen lymphocyte RNA using a forward primer, 5′-ATGAGCCATGGAATAAAAGACCT-3′, and a reverse primer, 5′-ACCAGCCAGGCCGCTTTGCA-3′.

Expression Vectors and Transfection
Mammalian cell expression vectors containing human TFL1 and TFL2 cDNAs were constructed by using a pQBI25 GFP vector (Takara, Japan). Each cloned cDNA containing its complete ORF was introduced using the XbaI sites of the vector. HA-tagged vectors were produced by linker ligation of oligonucleotides, 5′-CTAGATACCTTTATGTGTTTCTGACTAGCTGCG-3′ and 5′-CTAGCGCCATAGTCAGAACATCTAAGGGTGAT-3′, using the Xhol site of pQBI25 vector, E1F2C2, DCP14, and TIA-1 cDNAs including complete ORF were ligated into the HA-tagged vector described above using the Xhol site.

A mouse TFL cDNA was cloned by reverse transcription-PCR using total RNA of splenocytes and a forward primer, 5′-GCTCGAGATGAGCCCATGAAATAAGA-3′, and a reverse primer, 5′-CTAAGGATCCCCACGGAGGAG-3′. The mouse cDNA was ligated into the Xhol and BamHI sites of a DsRed monomer vector (Clontech).

Ba/F3 cells were electroporated with the Nucleofector system (Amazka). Adherent cell lines were transfected with Lipofectamine 2000 (Invitrogen) following the manufacturer’s protocols.

Immunoblot Analysis
Cells were lysed in lysis buffer containing 50 mmol/L Tris-HCl (pH 7.5), 150 mmol/L NaCl, 0.5% NP40, 1 mmol/L sodium vanadate, 50 mmol/L sodium fluoride, protease inhibitor cocktail (Sigma), and 1 mmol/L phenylmethylsulfonyl fluoride. For immunoprecipitation, total cell lysates were incubated with antihuman TFL antisera, followed by the addition of protein G-Sepharose (Amersham Biosciences), on a rotating shaker for 4 h at 4°C. The immunoprecipitates were washed thrice with lysis buffer and thrice with PBS.

Total cell lysates as well as immunoprecipitates were separated on 10% SDS-polyacrylamide gels and transferred onto polyvinylidene difluoride membranes (Millipore). The membranes were first incubated with TBST [20 mmol/L Tris-HCl (pH 7.5), 150 mmol/L NaCl, 0.1% Tween 20] containing 5% nonfat dried milk and probed with antihuman TFL or rabbit anti–β-actin (Sigma), followed by incubation with secondary horseradish peroxidase. Blots were visualized by enhanced chemiluminescence (Millipore).

Brdu, Phospho-Rb, and Apoptosis Assays
For BrdUrd assays, Ba/F3 cells were synchronized in G0-G1 phase by IL-3 deprivation for 16 h (48). After TFL expression vectors were introduced, the cells were incubated in complete medium supplemented with IL-3 for an additional 16 h, and the BrdUrd incorporation was assayed using the APC BrdU Flow Kit (Becton Dickinson).

For flow cytometric analyses of phospho-Rb, cells were fixed with 4% formaldehyde and permeabilized with 90% methanol, and then incubated with rabbit anti-mouse serine 807/811 phosphorylated Rb (Cell Signaling Technology) followed by incubation with antirabbit secondary antibody conjugated with Cy5 (Jackson ImmunoReserch). Annexin V-Cy5 (Becton Dickinson), biotinylated anti–active caspase-3 (Becton Dickinson), and streptavidin-allophycocyanin (e-Biosciences) were used to evaluate cell apoptosis.

Immunofluorescence Staining
Cells were grown on eight-chamber slides (Nunc). After transfection with TFL vectors, cells were fixed with 4% formaldehyde and permeabilized with 0.3% Triton X-100. After blocking procedures, cells were incubated with mouse anti- HA antibody (Covance) overnight at 4°C followed by incubation with antimouse secondary antibody conjugated with Cy5 (Jackson ImmunoReserch) before mounting. For detection of endogenous human TIA-1, we used anti–TIA-1 antibody (Santa Cruz) and secondary antibody conjugated with Alexa488 (Molecular Probes). Fluorescence was analyzed by confocal microscopy (510META, Carl Zeiss).

Statistical Analysis
Student’s t test was used for comparisons of numerical values in cell growth and apoptosis assays.

Disclosure of Potential Conflicts of Interest
No potential conflicts of interest were disclosed.

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