Alternative Expression of Platelet Glycoprotein Ibβ mRNA from an Adjacent 5’ Gene with an Imperfect Polyadenylation Signal Sequence

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Abstract

Glycoprotein (GP) Ib is a major component of the platelet membrane receptor for von Willebrand factor, designated the GP Ib-IX–V complex. GP Ib is composed of two subunits (GP Ibα and GP Ibβ) each synthesized from separate genes. The 206 amino acid precursor of GP Ibβ is synthesized from a 1.0-kb mRNA expressed by megakaryocytes and was originally characterized from cDNA clones of human erythroleukemic (HEL) cell mRNA, a cell line exhibiting megakaryocytic-like properties. The cell line CHRF-288–11 also exhibits megakaryocytic-like properties, but synthesizes two related GP Ibβ mRNA species of 3.5 and 1.0 kb. We performed cDNA cloning experiments to identify the origin of the 3.5-kb transcript and determine its relationship to the 1.0-kb GP Ibβ mRNA found in megakaryocytes, platelets, and HEL cells. Our cloning experiments demonstrate that the longer transcript results from a nonconsensus polyadenylation recognition sequence, 5’-AACAAT–3’, within a separate gene located upstream to the platelet GP Ibβ gene. In the absence of normal polyadenylation the more 5’ gene uses the polyadenylation site within its 3’ neighbor, the platelet GP Ibβ gene. This newly identified 5’ gene contains an open reading frame encoding 369 amino acids with a high degree of sequence similarity to an expanding family of GTP-binding proteins. (J. Clin. Invest. 1997. 99:520–525.) Key words: gene expression regulation • cloning, molecular • blood platelets • platelet membrane glycoproteins • cell division cycle proteins

Introduction

The GP Ib-IX–V complex is an essential platelet membrane receptor supporting platelet adhesion through its interaction with von Willebrand factor, an adhesive ligand of the blood and subendothelium (1, 2). The expression of the GP Ib-IX–V complex is dependent upon the coordinated assembly of at least three gene products, the α- and β-subunits of GP Ib, and GP IX (3). cDNA cloning experiments for these three subunits of the complex have utilized megakaryocytic-like cell lines and without exception the reported coding sequences have been representative of the mRNA species present in the megakaryocyte and platelet (4–6). However, the megakaryocytic cell line CHRF-288–11 (7) is an exception containing two related mRNA species from the locus encoding the β-subunit of GP Ib (GP Ibβ) (8). A short 1.0-kb transcript is consistent with the size of the megakaryocytic mRNA encoding platelet GP Ibβ, whereas a longer transcript approximating 3.5 kb appears to be present in a wide variety of cell types but noticeably absent in megakaryocytes and other megakaryocytic-like cell lines (8, 9). Kelly et al. reported the cloning of a cDNA hybridizing to the platelet GP Ibβ cDNA and suggested the 3.5-kb transcript expressed by endothelial cells possesses the potential to synthesize a novel 45-kD protein containing a carboxyl terminus of platelet GP Ibβ polypeptide sequence (8). However, a number of fundamental concerns were raised by their report. First, the cDNA sequence lacked a 3’ polyadenylated sequence and contained ~600 base pairs of sequence 3’ to the polyadenylation recognition signal utilized by megakaryocytes. It was unclear why endothelial cells would fail to recognize a consensus polyadenylation signal sequence unless the reported sequence was derived from a pre-mRNA species. A second concern was the presence of the single GP Ibβ intron sequence within the proposed open reading frame of the related endothelial cell mRNA. This situation suggested an intron of a gene expressed by megakaryocytes is a translated exon when expressed by endothelial cells, an unusual form of alternative processing.

We sought to investigate the origin of the longer transcript in CHRF-288–11 cells. Now we report a number of key differences from the sequence reported by Kelly et al. (8) and in the fundamental conclusions derived from their work. We show that a unique gene located 5’ to the platelet GP Ibβ gene utilizes two different polyadenylation sites. The more 5’ polyadenylation site results from an imperfect polyadenylation recognition sequence, whereas the more 3’ polyadenylation event results from a utilization of the consensus polyadenylation sequence within the platelet GP Ibβ gene. The use of the platelet gene’s polyadenylation sequence results in a 3.5-kb transcript containing an open reading frame encoding a polypeptide related to GTP-binding proteins and a downstream open reading frame encoding platelet GP Ibβ polypeptide. The potential to express platelet GP Ibβ by cells other than megakaryocytes and platelets is discussed along with the identity of a unique gene residing 5’ to the platelet GP Ibβ gene.

Methods

RNA isolation and northern transfer. CHRF-288–11 cells were obtained from Dr. Michael Lieberman (Children’s Hospital Medical Center, Cincinnati, OH) and were cultured as described in the original description of the cell line (7). Total RNA was isolated by ultracentrifugation through cesium chloride cushions using an established procedure (10). Poly (A)+ RNA was prepared by affinity chromatography using oligo-dT cellulose columns purchased from Life Technologies, Inc. (Bethesda, MD).
Gel electrophoresis of poly (A)⁺ RNA was performed through denaturing formaldehyde gels prepared as described by Lehrah et al. (11). Northern transfer and hybridization were performed using standard methodologies (10). The hybridization probe for the Northern analyses illustrated in Figs. 1 and 4 was a full-length platelet GP Ibβ cDNA fragment isolated from a human erythroleukemia cell cDNA library prepared by the authors. The coding sequence for platelet GP Ibβ within this cDNA fragment is identical to that described by Lopez et al. (5). After hybridization, the nitrocellulose filter was washed three times in 0.3 M sodium chloride/0.03 M sodium citrate/0.1% SDS (10 min, RT) and once in 0.15 M sodium chloride/0.015 M sodium citrate/0.1% SDS (30 min, 55°C). Less stringent washing conditions resulted in a relatively high background due to crosshybridization with 28S RNA. After washing the nitrocellulose was analyzed by autoradiography using Kodak X-OMAT film.

cDNA library construction. CHRF-288-11 poly (A)⁺ RNA was converted into cDNA and cloned into the bacteriophage AZAP/Ex-press available from Stratagene Cloning Systems (La Jolla, CA). First-strand cDNA was synthesized using Stratascript RNase H-deficient reverse transcriptase (Stratagene) primed by poly(dT). The unamplified cDNA library consisted of 1.5 × 10⁹ plaque-forming units. Random analysis of 20 independent clones revealed cDNA inserts in 18 clones ranging in size from 0.8-kb pairs to ~4 kb. 1 × 10⁶ plaque-forming units were expanded into a high-titer stock of the CHRF-288-11 cDNA library.

Two additional cDNA libraries were also used in this project. The first was a library prepared by the authors from poly(A)⁺ mRNA isolated from human endothelial cells. These cells were a gift from Dr. Eugene Levin (The Scripps Research Institute, La Jolla, CA) and were third-passage cells prepared from a donor umbilical vein. This library was also constructed in the AZAP/Express vector and contained more than 1 × 10⁶ recombinant bacteriophage. An additional fetal brain cDNA library was purchased from Stratagene.

Library screening and clone analysis. Screening of the library was performed using standard methodologies. A fragment of platelet GP Ibβ cDNA was radiolabeled using a random priming system available from Stratagene. Positive clones were plaque-purified via four rounds of screening before performing analyses on the cloned cDNA fragments. Each cDNA insert was rescued from its recombinant AZAP/Express vector via the in vivo excision protocol provided by Stratagene. DNA sequence analysis was performed on single-stranded DNA templates from the rescued filamentous phagemids using dideoxyribonucleotides and Sequenase (U.S. Biochemical, Cleveland, OH; 12). All DNA sequence reported in this manuscript was determined by analysis of both strands and from multiple runs of individual sequencing reactions. The majority of sequence was generated using sequence-specific primers synthesized after each individual DNA sequencing gel. In a few cases some confirming sequence data was obtained after subcloning individual restriction fragments in the filamentous phage, M13mp18 or M13mp19, and sequencing using standard dideoxy methodologies (12).

Computer analysis of the DNA sequence was performed using the PC-based software, Lasergene (DNASTAR Inc., Madison, WI) and sequence comparison programs available through the Internet from the National Center for Biotechnology Information (http://www.ncbi.nlm.nih.gov/).

Results

Origin of the 3.5-kb platelet GP Ibβ-related transcript expressed by CHRF-288–11 cells. CHRF-288–11 cells were established from a metastatic tumor in an infant with acute megakaryoblastic leukemia (7). These cells contain two mRNA species hybridizing with a full-length (1.0 kb) platelet GP Ibβ cDNA probe (Fig. 1). The smallest transcript is ~1.0 kb and most likely corresponds to the mature GP Ibβ mRNA transcript seen in other megakaryocytic cell lines, megakaryocytes, and platelets (5, 9). A larger, less abundant species is ~3.5 kb and is not an expressed transcript seen in megakaryocytic RNA (8). To investigate the origin of the 3.5-kb transcript we prepared a CHRF-288–11 cDNA library and screened the library with the full-length platelet GP Ibβ cDNA. Characterization of 17 hybridizing clones revealed that 14 contained cDNA fragments ranging from 0.9 to 1.0 kb corresponding to near full-length copies of the platelet GP Ibβ mRNA. Of the remaining three clones, two contained inserts ~1.9 kb and another was ~3.2 kb. The clone containing the longest insert, bacteriophage λC, was chosen for further DNA sequence analysis.

DNA sequence analysis of λC revealed a cDNA insert of 3,172 nucleotides (Fig. 2). The 3’ portion of the insert, nucleotides 1,866–3,172 corresponds to the GP Ibβ genomic gene sequence as reported by Yagi et al. (9), yet lacking the single GP Ibβ intron. Within this cDNA sequence is the major megakaryocytic transcription start site of the platelet GP Ibβ gene (nucleotide 2,218; 9), the complete coding sequence for the GP Ibβ precursor protein (nucleotides 2,245–2,862), the platelet GP Ibβ polyadenylation recognition sequence (nucleotides 3,153–3,158) and a polyadenylated tail (Fig. 2).

The more 5’ sequence within λC contain an open reading frame encoding a 385 residue polypeptide sequence and represented the only significant open reading frame within the more 5’ sequence. Thus, this fragment contained an unusual arrangement of cDNA sequence beginning with a unique 5’ sequence contiguous with the platelet GP Ibβ gene lacking its single intron. The possibility that the unique 5’ sequence might be a cloning artifact resulting from the analysis of a single clone was dismissed with the characterization of the two remaining clones purified during the initial screening of the same library, λ11C, and λ17C. Each of these clones contains varying lengths of the same cDNA sequence beginning at nucleotides 1,224 and 1,521 of λC, respectively. Clone λ17C contained an additional uncommon feature, the complete intron sequence of the platelet GP Ibβ gene, suggesting that in the context of the longer transcript, the single intron within the platelet GP Ibβ gene was not efficiently removed.

A gene residing 5’ to the platelet glycoprotein Ibβ gene utilizes two alternative polyadenylation recognition sequences. To obtain additional information on the newly identified 5’ sequence, the CHRF-288–11 cDNA library, a fetal brain cDNA library and a human endothelial cell cDNA library were each
The utilization of this imperfect polyadenylation site suggested a related mRNA transcript of ~2.3 kb would be present in the poly(A) mRNA from CHRF-288-11 cells. Indeed, Northern analysis identified a 2.3-kb transcript utilizing the 5' probe demonstrating that post-transcriptional modification of the primary transcript for the more 5' gene involves two different polyadenylation recognition sequences, its own imperfect site or the polyadenylation recognition site within its 3' neighbor, the platelet glycoprotein Ibβ gene (Figs. 3 and 4). Figure 3 conceptually illustrates the origin of these clones as they align with the derived sequence from individual cDNA clones.

Figure 2. Nucleotide sequence of the platelet GP Ibβ related-transcript of CHRF-288-11 cells. The complete DNA sequence of a bacteriophage λ clone designated, λ6C, is shown and contains two long open reading frames (ORFs). The first ORF, nucleotides 2–1,156, encodes 385 amino acids sharing homology with a family of GTP-binding proteins. The second ORF, nucleotides 2,245–2,862, is the complete ORF encoding the platelet GP Ibβ polypeptide. A polyadenylated tail resulting from the polyadenylation signal sequence in the downstream platelet GP Ibβ gene (underlined, nucleotides 3,153–3,158) represents the most 3' sequence. The transcription initiation site of the platelet GP Ibβ precursor transcript is highlighted by a double arrow (⇑) at nucleotide 2,218 (9). Thus, the longer transcript expressed by CHRF-288–11 cells contains a unique 5' sequence adjoining the platelet GP Ibβ gene. An imperfect polyadenylation site within the more 5' sequence is double-underlined (nucleotides 1,976–1,981). Accession number U59632 has been assigned by GenBank.

Several of the characterized clones contained a polyadenylated tail in which the last nucleotide preceding the polyadenylated tail corresponded to nucleotide 1,994 within the sequence presented in Fig. 2. Since the 3' boundary of these new clones lacked adenines, these clones were not the result of oligo d(T) mispriming in the first-strand reaction, and most likely represented cDNA products from an imperfect polyadenylation site. Since the 3' end of each clone (nucleotides 1,976–1,981) was identified by Northern analysis (Fig. 4), the location of each probe depicts its position relative to each transcript.

The utilization of the 1.0-kb platelet GP Ibβ transcript is expressed by megakaryocytes and megakaryocytic-like cells lines and is shown as it aligns within the longer transcript. Two radiolabeled probes representing 5' sequence (5' probe) and 3' sequence (3' probe) were used to confirm the presence of the three transcripts (A–C) by Northern analysis (Fig. 4). The location of each probe depicts its position relative to each transcript.

The isolation of multiple clones containing a polyadenylated tail suggested the longer transcript depicted in (a) is a result of the utilization of the consensus polyadenylation recognition sequence within the downstream platelet GP Ibβ gene. The transcript depicted in (B) is ~2.3 kb and was identified by Northern analysis (see Fig. 4) and by the characterization of cDNA clones (Results and see Fig. 5). The 1.0-kb platelet GP Ibβ transcript is expressed by megakaryocytes and megakaryocytic-like cells lines and is shown as it aligns within the longer transcript. Since the 3' boundary of these new clones lacked adenines, these clones were not the result of oligo d(T) mispriming in the first-strand reaction, and most likely represented cDNA products from an imperfect polyadenylation sequence (3′AACAAAT′) 13 nucleotides upstream from the 3' end of each clone (nucleotides 1,976–1,981 of Fig. 2). Figure 3 conceptually illustrates the origin of these clones as they align with the derived sequence from individual cDNA clones.

The utilization of this imperfect polyadenylation site suggested a related mRNA transcript of ~2.3 kb would be present in the poly(A)+ mRNA from CHRF-288-11 cells. Indeed, Northern analysis identified a 2.3-kb transcript utilizing the 5' probe demonstrating that post-transcriptional modification of the primary transcript for the more 5' gene involves two different polyadenylation recognition sequences, its own imperfect site or the polyadenylation recognition site within its 3' neighbor, the platelet glycoprotein Ibβ gene (Figs. 3 and 4). Utilizing a 3' probe, transcripts of 3.5 and 1.0 kb were observed, each presumably the product of different gene promot-
ers, yet utilizing the same polyadenylation recognition sequence (Figs. 3 and 4).

Sequence analysis of transcripts polyadenylated from the imperfect recognition sequence. A common cDNA sequence representing the 2.3-kb transcript is presented in Fig. 5. The presented sequence is a composite of two cDNA clones, designated λ12F and λ6C, isolated from fetal brain and endothelial cell cDNA libraries, respectively. Clone λ6C contained the sequence from nucleotide 83 to 2,032 (Fig. 5). Nucleotide sequence 1–82 was derived from clone λ12F. The sequence contains an open reading frame encoding 369 amino acids with an initiating methionine codon (nucleotide 88–90) immediately preceded by a consensus Kozak sequence (AUG) (Fig. 5).

The most 5’ sequence illustrated in Fig. 5 differs from the 5’ sequence of λ6C (Fig. 3). Specifically, the sequences are identical in the 3’ direction starting at nucleotide 104 of the λ6C sequence (Fig. 2) and nucleotide 142 of the composite sequence presented in Fig. 5. Concerns that the more 5’ sequence of λ6C (nucleotides 1–103 of Fig. 2) might be a cloning artifact were eased by the observation of multiple bacteriophage λ clones, designated λ12F and λ6C, containing two open reading frames (see also Fig. 4). Nucleotide sequence 83–2,032 (Fig. 5) represents a composite of two cDNA clones, designated λ12F and λ6C. The sequences are identical with 3.5- and 1.0-kb species resulting from polyadenylation via the recognition sequence within the platelet GP Ibβ gene. A photograph of the autoradiograph is shown along with the migrating positions of RNA molecular weight standards.

Discussion

Our results provide evidence that a gene residing in close 5’ proximity to the platelet GP Ibβ gene directly influences the expression of the platelet GP Ibβ gene. The utilization of the platelet GP Ibβ polyadenylation sequence by the more 5’ gene results from an imperfect polyadenylation recognition sequence (AACAAT) in the more 5’ gene. Indeed, in vitro studies have confirmed that mutation of a consensus polyadenylation sequence (AATAAA → AACAAT) dramatically reduces the efficiency of polyadenylation (15, 16). In the present study, an imperfect polyadenylation sequence results in the alternative usage of a polyadenylation recognition site within a nearby 3’ gene. The net result is a longer transcript, approximating 3.5 kb, and containing two open reading frames (Fig. 3).

In the course of our analysis, we isolated multiple clones from the CHRF-288–11, endothelial cell and fetal brain cDNA libraries. In addition, we isolated and characterized more than 40 clones identified from the EST database and available through the cDNA consortium. It is clear from this analysis that the 5’ sequence varies with a number of alternatively processed variants. In fact, a shorter exposure of the Northern blots presented in Figs. 1 and 4 suggests the 3.5 kb mRNA transcript is composed of several transcripts of slightly varying length. The sequences presented in Figs. 2 and 5 represent two of the more common processing variants identified. Whether these processing variants are specific for a particular cell type was not obvious from the characterization of our clones. Additional length heterogeneity within the longer transcript was provided by our isolation of a single cDNA clone containing the complete intron within the GP Ibβ gene (data not shown). The presence or absence of the GP Ib intron would produce a 274 ribonucleotide length difference among the mature
transcripts. Future studies characterizing genomic clones of this locus may aid in clarifying the identity of each processing variant.

A search of the current GenBank database with the polypeptide sequence derived from \textit{A8EN} (Fig. 5) suggests the encoded polypeptide would be an additional member of a growing family of proteins related to the cell division cycle (CDC) proteins of yeast (17). We have designated this polypeptide as human cell division cycle-related (hCDCrel) protein 1, anticipating additional variants and related human sequences will be identified. Homologous proteins have been found in mouse and \textit{Drosophila} and an alignment with the deduced amino acid sequence of hCDCrel-1 is presented (18–20; Fig. 6). Experimental evidence for the \textit{Drosophila} homologue confirms its role in cytokinesis suggesting that a unifying functional attribute of these proteins may be a role in some aspect of cell division (20, 21). Structurally, the unifying feature of these proteins is the presence of GTP- or phosphate-binding motifs (22, 23; Fig. 6). Whether the 5' gene and the platelet GP Ib\(\beta\) gene are related remains unknown. The imperfect polyadenylation sequence within the more 5' gene is less than 250 nucleotides upstream of the megakaryocytic transcription initiation site within the platelet GP Ib\(\beta\) gene (Fig. 2). Indeed, in our own preliminary studies a fragment extending 262 base pairs upstream of the platelet glycoprotein Ib\(\beta\) transcription initiation site can support reporter gene expression in human erythroleukemia (HEL) cells suggesting it may represent a megakaryocytic promoter sequence.

Our analysis of \textit{A6C} (Fig. 2) suggests some unidentified 5' sequence approximating 300 nucleotides remains to be identified based on a mature transcript size of 3.5 kb. This analysis will be necessary to determine if the longer transcript contains the potential to encode a polypeptide differing at the amino terminus from hCDCrel-1. Certainly, a 3.5-kb transcript containing the potential to code for two proteins would be unusual. Polycistronic units and internal ribosome binding sites in prokaryotes are common, but the translation of eukaryotic mRNA is most commonly initiated by ribosome binding to a 5' terminal cap structure and scanning for an initiation codon (24). According to this mechanism, we might predict for the 3.5-kb transcript illustrated in Fig. 3 efficient protein synthesis from the more 5' coding sequence, and likewise, no protein synthesis from the more 3' open reading frame encoding platelet glycoprotein Ib\(\beta\). However, the initiation of translation in eukaryotic cells via a cap-independent method, or via an internal ribosome entry site, is an efficient process for the synthesis of viral proteins dependent upon cellular not viral gene products (25–27). Indeed, it has been suggested that eukaryotic internal ribosome binding sites may be more widespread than previously recognized but lacking, to date, the description of a consensus motif (27). The synthesis of platelet glycoprotein Ib\(\beta\) from the longer transcript may represent an example of internal ribosome binding and will be examined in future studies to determine if synthesis of this protein can occur via a cap-independent mechanism.

Kelly et al. reported the cDNA cloning of an endothelial cell-expressed form of the 3.5-kb GP Ib\(\beta\) transcript (8). However, their cloning did not identify the 3' or 5' boundaries of the endothelial cDNA fragments. In fact, their reported sequence contained \(\sim 600\) nucleotides 5' to the platelet polyadenylation site and was a composite of three overlapping cDNA fragments. They concluded that the longer transcript potentially expresses a 45-kD protein that is essentially a fusion protein composed of a unique amino terminus coupled to the platelet GP Ib\(\beta\) polypeptide sequence. This conclusion was based on immunoreactivity of a 45-kD protein present in endothelial cell extracts and recognized by a polyclonal anti-GP Ib\(\beta\) antibody. Their open reading frame for this protein assumed translation through the single intron within the platelet GP Ib\(\beta\) gene (9). Unlike the reported sequence of Kelly et al., our derived sequence does not contain an open reading frame capable of synthesizing the 45-kD polypeptide due to a number of insertions and deletions differing from their reported sequence. Including the present study, three different reports contain DNA sequence through the regions encoding the potential 45-kD protein. A comparison of the reported genomic sequence of Yagi et al. (9) to the cDNA sequence reported by Kelly et al. (8) reveals four discrepancies. Using the sequence of Kelly et al. as a reference, the differences are an inserted cytosine base at position 675, the absence of a cytosine/guanine dinucleotide pair at position 844 and a silent transversion at position 1,275. Three of these discrepancies are critical since they

Figure 6. Comparison of the conceptual translation product, hCDCrel-1, to similar GenBank entries. The polypeptide, designated hCDCrel-1, shares homology with a family of GTP-binding proteins. Two murine proteins, H5 and D6 (GenBank accession numbers X61452 and D28540, respectively) and a \textit{Drosophila} peanut homologue (accession number U08103) are aligned illustrating the conservation of sequence through more than 300 amino acids of each sequence. hCDCrel-1 is 75, 56, and 46% identical to H5, D6, and peanut polypeptides, respectively. Each of the aligned proteins contains motifs involved in GDP/GTP exchange (\textit{GX}_{4,5} \textit{GK[S/T]} or in GTP hydrolysis (\text{DX},\text{G}), residues 51–59 and 108–111 of hCDCrel-1.
result in frame-shifts within the proposed open reading frame encoding the 45-kD polypeptide (8). Our sequence analysis agrees with that of Yagi et al. at these positions further supporting the conclusion that the synthesis of this 45-kD polypeptide is not possible. Thus, the immunoreactivity of a 45-kD endothelial cell protein most likely represents cross-reacting material distinct from GP Ibβ peptide. However, based on this previous study there are ~50 EST database entries containing gene sequence identical to that presented in Figs. 2 and 5, yet designated platelet GP Ibβ precursor polypeptide. Our data would support a renaming of these sequences to hCDCrel-1 protein.

The current studies provide two directions for future research. First, we have identified a new gene within the 22q11.2 chromosomal locus. Initiatives to identify novel genes from this locus have been fueled by studies linking DiGeorge syndrome and other cardio-facial abnormalities to this region of chromosome 22 (28, 29). The close relationship of DiGeorge syndrome and the platelet GP Ibβ gene has been verified by chromosomal deletion of 22q11.2 in an individual diagnosed with both the Bernard-Soulier syndrome (congenital absence of the platelet GP Ib-IX receptor) and the DiGeorge syndrome (30). Indeed, the sequence similarities between the Drosophila peanut protein and hCDCrel-1 suggests a function for hCDCrel-1 in development since the expression of the peanut protein is linked to proliferation of the imaginal disc tissues of Drosophila (20). Thus, the identity of the 5’ gene and its product should be characterized as another candidate gene for normal development. Finally, the identification of platelet GP Ibβ coding sequence in cells of nonhematopoietic origin suggests the presence of this protein in cells other than megakaryocytes and platelets. Thus, a previously unrecognized functional consequence for the expression of this protein may exist and should be examined.

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