Novel RNA-binding Protein P311 Binds Eukaryotic Translation Initiation Factor 3 Subunit b (eIF3b) to Promote Translation of Transforming Growth Factor β1-3 (TGF-β1-3)*

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Background: P311 is a stimulator of TGF-β1-3 translation, but its mechanism of action is unknown.

Results: P311 binds eIF3b and the 5’UTRs of TGF-β1-3 mRNAs. Thereby, P311 recruits TGF-β1-3 mRNAs to the translation machinery.

Conclusion: P311 is an RNA-binding protein that binds to eIF3b to stimulate TGF-β1-3 translation.

Significance: Our studies add a new level of complexity to TGF-β signaling regulation.

P311, a conserved 8-kDa intracellular protein expressed in brain, smooth muscle, regenerating tissues, and malignant glioblastomas, represents the first documented stimulator of TGF-β1-3 translation in vitro and in vivo. Here we initiated efforts to define the mechanism underlying P311 function. POND character (Predictor Of Naturally Disordered Regions) analysis suggested that P311 binds directly to TGF-β1-3 translation, but its mechanism of action is unknown. Previously, little had been known regarding the processes underlying the translation of TGF-β1-3 (1, 2). Therefore, requiring an interacting partner to acquire tertiary structure and function. Immunoprecipitation coupled with mass spectrometry identified eIF3 subunit b (eIF3b) as a novel P311 binding partner. Immunohistochemical colocalization, GST pulldown, and surface plasmon resonance studies revealed that P311 binds directly to TGF-β1-3 translation. Because TGF-β1-3 translation involves three separate steps: initiation, elongation, and termination (22), with initiation serving generally as the rate-limiting step in this process (23, 24). In eukaryotes, translation initiation begins with the binding of the small (40 S) ribosome subunit to a series of initiation factors, including eIF1, eIF1A, eIF3, and eIF5, and the eIF2-GTP-Met-tRNA_Met ternary complex (TC), to form the 43 S preinitiation complex (PIC). The 43 S PIC then binds to the capped 5’ end of mRNA through eIF4G and the eIF3 complex to scan for the start codon. Following the eIF5-dependent release of eIF2-GDP

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The abbreviations used are: TC, ternary complex; PIC, preinitiation complex; IC, initiation complex; SPR, surface plasmon resonance; IP, immunoprecipitation; Ab, antibody; VSMC, vascular smooth muscle cell; RIP, RNA immunoprecipitation; EBM, eIF3b binding motif, aa, amino acid; HSLMC, human lung smooth muscle cell; EV, empty vector; RRM, RNA recognition motif.

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and the other eIFs present in the 43 S PIC, the 43 S ribosome subsequently recruits the large (60 S) ribosomal subunit to produce an 80 S initiation complex (IC) containing Met-tRNA<sub>Met</sub> base-paired to AUG in the P site of the ribosome, which is the site where the first aminoacyl tRNA (fMet-tRNA<sub>Met</sub>) enters to begin the elongation phase of protein synthesis (23–26).

Of all the initiation factors involved in mRNA translation, eIF3 is the largest and most complex. With a molecular mass of 800 kDa, eIF3 is composed of 13 protein subunits (elF3a-elF3m), which are present in stoichiometrically equal amounts (27, 28), with six of the subunits (i.e. elF3a, b, c, e, f, and h) representing core subunits sufficient to support translation initiation (29, 30). eIF3 stimulates most of the reactions in the initiation pathway, including assembly of the elF2-GTP-Met-tRNA<sub>Met</sub>TC, binding of TC and other components of the 43 S PIC to the 40 S subunit, mRNA recruitment and scanning for AUG recognition and prevention of premature association of 43 S binds to 60 S ribosomal subunits (23, 25, 31, 32). The role of each eIF3 subunit in normal and pathological translation is currently the subject of intense study.

Here we demonstrate that P311 is an intrinsically disordered protein that binds elF3b and the 5′UTRs of TGF-β1, 2, and 3 mRNAs concomitantly, with little or no binding to other tested 5′UTRs. By directly promoting the enrichment of mRNAs in the translation machinery, P311 enhances TGF-β protein levels. Our findings define P311 as a novel RNA-binding protein as well as one of the few documented elF3b partners to lie outside of the general translation initiation machinery.

EXPERIMENTAL PROCEDURES

**Cell Cultures and Animal Tissues**—Mouse NIH-3T3 cells were obtained from the ATCC. NIH-3T3 cells were cultured in DMEM supplemented with 10% (v/v) fetal bovine serum, 100 units/ml penicillin-streptomycin, and 2.5 μg/ml Fungizone. Aortas were dissected from 4-month-old C57BL/6 male mice, and the attached fat and soft tissues were trimmed under a microdissecting microscope. The dissected blood vessels were immediately used for experiments or frozen at −80 °C until use.

**Transfection, Gene Silencing, and Reporter Assay**—For transient transfections, NIH-3T3 cells at ~70% confluence were transfected with a P311 expression construct, pCMV-MYC-p311, and/or the elF3b expression construct pCMV-HA-elF3b (Clontech) using Lipofectamine 2000 reagent in Opti-MEM (Life Technologies). For elF3b and elF3a gene silencing, FlexiTube GeneSolution siRNAs (targeting four different regions of the Elf3b or Elf3a gene) (Qiagen) were transfected into cells using Lipofectamine RNAiMAX reagent (Life Technologies). For the reporter assay, TGF-β1-3 5′ and 3′UTRs were cloned into the pGL3-promoter firefly luciferase vector (Promega) to generate reporter constructs (1). NIH-3T3 cells were plated on 6-well plates and transfected with the P311 expression vector, elF3b siRNA, and the reporter constructs and Renilla luciferase internal control vector pRL-SV40. Forty-eight hours after cotransfection, the Dual-Luciferase reporter assay system (Promega) was used to determine the ratio of firefly to Renilla luciferase activity, as indicated by the manufacturer.

**Protein Purification**—To purify full-length elF3b and fragments from BL21(DE3) Escherichia coli (New England Biolabs), a GST tag was fused at the N terminus and expressed in the pGEX-6P-1 vector. The proteins were induced to express in BL21(DE3) cells by 0.4 mM isopropyl 1-thio-β-D-galactopyranoside at 37 °C for 4 h. The E. coli cells were then lysed by sonication in 20 mM phosphate buffer (pH 7.5) containing 500 mM NaCl, 1 mM DTT, 0.2% Tween 20, 0.2% Nonidet P-40, and protease inhibitors. Cell lysates were clarified by centrifugation at 4 °C and 13,000 rpm for 30 min. The supernatant was loaded onto a GSTrap column (GE Healthcare). After extensive washes with the lysis buffer, the proteins were eluted in the same buffer containing 10 mM glutathione. The eluted proteins were dialyzed overnight against the binding buffer (25 mM Tris-HCl, 150 mM NaCl, 1 mM DTT, 0.02% Tween 20, 0.02% Nonidet P-40, and 10% glycerol (pH 7.5)). To purify full-length P311 from E. coli, a thioredoxin tag and a His<sub>6</sub> tag were fused at the N terminus. The protein was purified with a similar method as mentioned above using a HisTrap column (GE Healthcare). While dialyzing overnight against the binding buffer, the thioredoxin tag and the His<sub>6</sub> tag were cleaved off with a His<sub>6</sub>-c-tagged 3C protease. The cleaved product was further purified by passing through a HisTrap column to remove the tags and the 3C protease.

**Surface Plasmon Resonance (SPR)**—For analysis of the binding of P311 and elF3b by SPR, recombinant P311 was immobilized onto the surface of a CM5 (GE Healthcare) sensor chip to ~2000 RU (response units). Purified elF3b (the GST tag was cleaved) at various concentrations was allowed to flow over the chip, and binding to the P311 was measured by SPR (Biacore 3000, GE Healthcare) with a negative control surface without immobilized P311. The concentrations of elF3b in the fluid phase were 5, 2, 1, 0.5, and 0 μM in HBS-EP buffer (0.01 M HEPES, pH 7.4, 0.15 mM NaCl, 3 mM EDTA, 0.005% v/v Surfactant P20) (GE Healthcare) at 4 °C, respectively.

**Co-IP and GST-pulldown**—For co-IP, MYC-P311/HA-elF3b-transfected NIH-3T3 cells were lysed in lysis buffer (25 mM Tris-HCl, 150 mM NaCl, 1 mM DTT, 0.02% Tween 20, 1% Nonidet P-40, and 10% glycerol (pH 7.5)) and clarified by centrifugation at 4 °C and 13,000 rpm for 10 min. The supernatants (1 mg of total protein in a 600-μl volume) were incubated with 20 μl of anti-MYC beads (50% slurry) (Sigma-Aldrich) or 30 μl of anti-HA beads (25% slurry) (Santa Cruz Biotechnology) by rotating at 4 °C for 2 h. After extensive washing three times with 800 μl of lysis buffer, the immunoprecipitated proteins were eluted by 20 μl of SDS loading buffer for 10 min at 95 °C and then analyzed by Western blotting. For mass spectrometry, the co-IP material was washed extensively four times with 800 μl of high stringent radioimmune precipitation assay buffer (Pierce). For GST pulldown, the purified GST-elF3b fragments were incubated with purified P311 in binding buffer on a rocker at 4 °C for 2 h, and then 20 μl of GST beads (25% slurry) (Pierce) was added to the sample. Two hours later, the beads were washed three times with 800 μl of binding buffer, and the proteins on the beads were eluted by SDS loading buffer and analyzed by Western blotting.

**Mass Spectroscopy**—Proteins were immunoprecipitated from NIH-3T3 cells transfected with MYC-P311 or empty vector using anti-MYC Ab. The immunoprecipitated samples were resolved by SDS-PAGE and digested by trypsin. The trypsinized
samples were analyzed using the Thermo Scientific LTQ Orbitrap LC/MS system. Peptide spectral matching was performed using the Mascot searching algorithm. The results from mass spectrometry analysis are shown in Fig. 1B.

**Western Blot Analysis**—Cell lysates, co-IP/GST pulldown samples, or tissue lysates were resolved on SDS-PAGE gels and transferred onto nitrocellulose membranes (Bio-Rad) for immunoblot analysis. Antibodies against P311 were generated in rabbits by Cocalico Biological, Inc. using bacterially purified P311 as the antigen. Anti-MYC (Sigma–Aldrich), anti-HA Ab, anti-GST Ab, and anti-GAPDH (Santa Cruz Biotechnology) were also used to detect target proteins. An ECL plus Western blotting detection system (GE Healthcare) was used for detecting the chemiluminescence signal.

**Immunofluorescence Staining**—To detect the colocalization of P311 and eIF3b in VSMCs isolated from aortas (1), cells were cultured on four-chamber slides (Thermo Scientific) until 50% confluence. The VSMCs were then washed three times with 400 μl of PBS and fixed with 4% PFA (paraformaldehyde) for 10 min at room temperature. The fixed cells were permeabilized with 0.2% Triton X-100 for 10 min. After washing extensively with PBS in a 100-ml container, cells were blocked with 3% BSA for 30 min, followed by incubation with primary anti-P311 Ab overnight at 4 °C. After washing, cells were incubated with Alexa Fluor 488 goat anti-rat secondary Ab (Life Technologies) and incubated sequentially with anti-eIF3b primary Ab (Abcam), followed by incubation with Alexa Fluor 594 secondary Ab (Life Technologies). Slides were mounted with 50% glycerol containing DAPI for nuclear staining and observed under a Zeiss MicroImaging

**Peptide Synthesis and Competition Assays**—P311 peptides (N, M, and C) were chemically synthesized using an ABI 433A peptide synthesizer using Fmoc (N-[9-fluorenylethoxycarbonyl]-protected aas (Midwest Bio-Tech). Peptides were side chain-unprotected and removed from the resin by TFA treatment in the presence of 2.5% trisopropylsilane, 2.0% anisole, and 0.5% water and then precipitated by addition of diethyl ether. Crude peptides were purified by preparative reversed-phase HPLC using a C18 column. Molecular weights of the final products were determined by mass spectrometry (Bruker UltraflexTreme MALDI-TOF/TOF). Purified P311 and the GST-eIF3b F4 fragment (10 μM final concentration) were incubated with increasing amounts of synthesized peptides (0, 10, 50, and 100 μM final concentrations) in a final volume of 100 μl of binding buffer. The samples were incubated at 4 °C with gentle rocking, and then GST beads ( Pierce) were added to the sample. After washing three times with 800 μl of binding buffer, the supernatants were removed by centrifugation at 3000 rpm for 30 s, and then the beads were boiled for 10 min in 15 μl of SDS loading buffer. Eluted proteins were resolved by SDS-PAGE and then analyzed by Western blotting.

**RNA precipitation (RIP)**—Forty-eight hours after transfection, cells were preincubated with cycloheximide (100 μg/ml for 15 min) and lysed with polysome extraction buffer containing 20 mM Tris-HCl (pH 7.5), 100 mM KCl, 5 mM MgCl2, and 0.5% Nonidet P-40. Cytoplasmic lysates were fractionated by ultracentrifugation through 10–50% linear sucrose gradients and divided into 12 fractions. Proteins were analyzed by Western blotting, and RNA was extracted from each fraction using TRIzol (Life Technologies) and used for RT-qPCR analysis.

**Statistics**—Quantitative data are presented as the mean ± S.D. and were compared statistically by Student’s *t* test, *p < 0.05* was considered statistically significant.

**RESULTS**

**P311 Interacts with eIF3b in an RNA-independent Manner**—POND® (predictor of naturally disordered regions) analysis (34) suggested that P311 is an intrinsically unstructured protein. CD spectroscopy confirmed the lack of structure in the isolated protein in solution (Fig. 1). Specific ellipticity at all wavelengths measured (260–200 nm) showed a completely linear relationship to protein concentration, indicating that the protein is monomeric under these conditions. As an intrinsically unstructured protein, P311 could acquire tertiary structure by binding to a ligand, e.g., another protein (35). To screen for potential binding partners, we performed co-IPs using cell lysates prepared from NIH-3T3 cells expressing MYC-tagged P311 and X-MYC Ab. Mass spectrometric analysis of the co-IP material consistently identified eIF3b as well as several cytoskeletal proteins (Fig. 1B). P311-eIF3b interaction was confirmed by co-IP of MYC-P311 and HA-eIF3b, followed by
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A

![Graph showing ellipsometry against wavelength (nm).]

B

| Protein          | Number of peptides |
|------------------|--------------------|
| P311 (NREP)      | 38                 |
| MYH9             | 15                 |
| ACTB             | 8                  |
| EIF3B            | 8                  |
| FLNA*            | 6                  |
| GDI2             | 3                  |
| RAB7             | 3                  |
| EIF3G            | 2                  |
| RPS13            | 2                  |
| RBMX             | 1                  |

C

**NIH-3T3**

- **KDa**
  - 110
  - 10
  - 3.5

- **IP: HA**
  - HA-elf3b
  - MYC-P311

- **WB: HA**
  - 110
  - 10
  - 3.5

- **WB: MYC**
  - 110
  - 10
  - 3.5

- **WB: GAPDH**
  - 110
  - 10
  - 40

D

**Aorta**

- **IP: elf3b**
  - 110
  - 15
  - 10
  - 3.5

- **IP: P311**
  - 110
  - 15
  - 10
  - 3.5

- **WB: elf3b**
  - 110
  - 15
  - 10

- **WB: P311**
  - 110
  - 15
  - 10

E

- **P311**
- **elf3b**
- **DAPI**
- **Merge**

F

- **Control siRNA**
  - elf3b
  - MYC-P311
  - GAPDH

- **elf3b siRNA**
  - (200 nM + EV)
  - (200 nM + P311)

G

- **Control siRNA**
  - elf3a
  - MYC-P311
  - GAPDH

- **elf3a siRNA**
  - (200 nM + EV)
  - (200 nM + P311)

**Bar graphs**

- **Fold increase in luciferase activity**
  - TGF-β1 5'/3'-UTR
  - TGF-β2 5'/3'-UTR
  - TGF-β3 5'/3'-UTR
  - β-Actin 5'/3'-UTR

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  - TGF-β1 5'/3'-UTR
  - TGF-β2 5'/3'-UTR
  - TGF-β3 5'/3'-UTR
  - β-Actin 5'/3'-UTR
immunoblot analysis using X-MYC and X-HA Abs (Fig. 1C).

Interaction of endogenous P311 and eIF3b was demonstrated by immunoblot identification of P311 in co-IPs performed with X-eIF3b Ab on lysates of mouse aortic smooth muscle (Fig. 1D) and by confocal microscopy, revealing their colocalization in VSMCs (Fig. 1E).

Because the elf3 complex binds different types of RNA (36, 37), we tested whether the P311-elf3b interaction might be RNA-mediated by treating the samples with RNase prior to co-IP. Samples without RNase treatment were used as controls. The co-IP material was washed and resolved by SDS-PAGE. No change was observed in P311-elf3b binding in the absence of RNA (data not shown).

**elf3b-P311 Interaction Does Not Affect the Level and Composition of elf3b-associated Subunits—**Immunoblot analysis of elf3b IP material with the appropriate Abs showed no significant difference in the levels and composition of elf3b and elf3b-associated subunits (elf3a, g, i, and j) in the presence or absence of P311, indicating that P311 does not serve as an intermediary to recruit elf3b-associated subunits to the elf3 complex (data not shown).

**elf3b-P311 Interaction Is Required for P311-induced Translation of TGF-β.** elf3a were silenced by RNAi (Fig. 1, F and G) in the presence or absence of P311, and then RNA reporter assays were performed using TGF-β5/3’UTR expression vectors (1). Upon elf3b down-regulation, the absence of P311 decreased TGF-β translation more dramatically than when P311 was present (Fig. 1F). Upon elf3a down-regulation (negative control), no significant difference was observed between the absence and presence of P311 on TGF-β translation (Fig. 1G). Together, these results indicate that elf3b is required for P311-controlled TGF-β translation, whereas elf3a is not.

**P311-elf3b Interaction Is through Direct Binding—**GST pulldown assays were performed using purified recombinant P311 and GST-elf3b with GST as a negative control. As shown in Fig. 2A, GST-elf3b, but not GST, bound to P311. SPR was used to examine whether there is direct P311-elf3b binding. After removing the GST tag from the recombinant elf3b (Fig. 2B), P311 was immobilized on a BIAcore sensor chip, and various concentrations of elf3b were allowed to flow over the chip while the interaction of P311 with elf3b in the mobile fluid phase was monitored. A BIAcore sensor chip without immobilized P311 served as a negative control. The SPR study confirmed direct P311-elf3b interaction with a Kd of 1.26 μM and 1:1 (mol:mol) stoichiometry. (Fig. 2B).

**P311-elf3b Binding Is Mediated by the elf3b RRM and the Conserved P311 elf3b Binding Motif (EBM).** A series of GST-elf3b fragments was generated for GST pulldown studies (Fig. 2C). Only the full-length elf3b (FL) and an N-terminal fragment encompassing the elf3b RRM fragment (F2) were able to bind to P311 (Fig. 2D). By testing each half of the F2 fragment (F3 and F4), F4, representing the elf3b RRM, was identified as the binding site for P311 (Fig. 2E).

P311 was divided into three segments of similar length: N-terminal (amino acid (aa) residues 1–22), referred to as N segment; middle (aa residues 23–46), referred to as M segment; and C-terminal (aa residues 47–68), referred to as C segment (Fig. 3A). Peptides representing each of these three segments (referred to as N peptide, M peptide, and C peptide) were chemically synthesized and used to perform GST-F4 gel motility shift assays with GST as a negative control. A shift in F4 motility was observed when incubated with the M peptide but not with the other two (Fig. 3B). Competition assays showed that addition of M peptide reduced P311 binding to F4 in a concentration-dependent manner, whereas N and C peptides had no effect (Fig. 3C).

To further demarcate the elf3b binding site within the P311 M segment, we searched for conserved elf31 aa sequences among higher eukaryotic species. This analysis revealed the presence of a highly conserved 11-aa-long stretch (from leucine 30 to lysine 40) within the middle region of P311 (Fig. 3D, stars). A series of tandem mutations in the conserved region were generated by alanine substitutions (Fig. 3D). The P311 mutants were coexpressed together with HA-elf3b, followed by co-IPs with X-HA Ab. The mutant in which alanine residues were substituted for all 11 aa inhibited P311-elf3b binding by 95% (Fig. 3E). Therefore, this 11-aa sequence was designated as an EBM and its mutant as P311(MutEBM) (Fig. 3E). In vitro binding assays were performed to confirm the involvement of this region. Recombinant GST-F4 was incubated with MYC-P311 or MYC-P311(MutEBM), each of them bound to X-MYC Ab-conjugated beads. After extensive washing, the material bound to the beads was examined by Western blotting using X-GST Ab. As shown in Fig. 3F, the elf3b F4 fragment bound to P311, whereas no binding was observed between F4 and P311(MutEBM).

**The EBM Is Required for P311 Induction of TGF-β Translation and for a P311-mediated Increase in the Levels of TGF-β in Human Lung Smooth Muscle Cells (HLSMCs).** Polysome fractionation analysis of NIH-3T3 cells expressing P311 or P311(MutEBM) was performed. The majority of P311, but not P311(MutEBM), β-actin, or GAPDH, cosedimented with elf3b

**FIGURE 1. P311 Interacts with elf3b. A, P311 is an unstructured protein. Purified recombinant P311 at different concentrations was analyzed by CD. mdeg, millidegrees. B, summary of the mass spectrometry analysis. MYC-P311 and potentially interacting proteins were communoprecipitated from NIH-3T3 cells transfected with MYC-P311 or empty vector using anti-MYC Ab. The communoprecipitated samples were analyzed by mass spectrometry using the Thermo Scientific LTQ Orbitrap LC/MS system. Peptide spectral matching was performed using the Mascot searching algorithm. C, Co-IP of HA-elf3b and MYC-P311 from NIH-3T3 cells transfected with either of them alone or in combination. After 48 h, the cells were lysed, and the lysates were used for IP with the indicated antibodies. IP material was resolved by SDS-PAGE, transferred to membranes, and immunoblotted with the indicated antibodies. WB, Western blot. D, co-IP of endogenous elf3b and P311 from mouse aortic smooth muscle lysate using the indicated antibodies. E, confocal microscopic colocalization of immunostained P311 and elf3b in VSMCs. F, luciferase reporter assays for TGF-β5’UTRs in NIH-3T3 cells transfected with P311 or EV while elf3b is concomitantly down-regulated by siRNA or exposed to scrambled siRNA control. G, luciferase reporter assays for TGF-β5’UTRs in NIH-3T3 cells transfected with EV or P311 while elf3a is concomitantly silenced by siRNA or exposed to scrambled siRNA control. In F and G, elf3b and elf3a were silenced by siRNAs targeting four different regions of the elf3b and elf3a genes, respectively. A final concentration of 200 nm was used. *p < 0.05; **p < 0.01.
FIGURE 2. P311 directly binds to the eIF3b RRM. A, GST pulldown assays using bacterially expressed recombinant GST-eIF3b and P311. GST-eIF3b was incubated with or without P311, followed by addition of GST beads. After extensive washing, the proteins bound to the GST beads were separated by SDS-PAGE and detected with anti-GST or anti-P311 Ab. The input recombinant proteins were separated by SDS-PAGE and visualized with Coomassie Blue staining. BSA was added as a nonspecific competitor. The asterisks indicate the main bands of input of full-length GST-eIF3b. WB, Western blot. B, analysis of P311-eIF3b binding by SPR. Recombinant P311 was immobilized onto the surface of a CMS sensor chip to 2000 RU. Various concentrations of purified eIF3b (after removal of the GST tag) were allowed to flow over the chip, and binding to the P311 was measured by SPR with a negative control surface devoid of P311. The concentrations of eIF3b in the fluid phase were 5, 2, 1, 0.5, and 0 M, respectively. C, eIF3b and its fragments used in the following studies. Each fragment was fused with a GST tag at the N terminus for protein purification and GST pulldown using GST beads. FL, full-length. D and E, binding of eIF3b RRM to P311. Recombinant GST tag alone, GST-full-length eIF3b (FL), and GST-eIF3b fragments (F1, GST-eIF3b 1–500 aa; F2, GST-eIF3b 1–300 aa; F3, GST-eIF3b 1–160 aa; F4, GST-eIF3b 161–264 aa) were incubated with purified P311, followed by addition of GST beads. After washing, the proteins were released from the beads in SDS gel loading buffer and analyzed by Western blotting. An aliquot of the binding reaction was analyzed separately by SDS-PAGE and stained with Coomassie Blue as input (left panels). BSA was added to all reactions as a nonspecific competitor. The asterisks indicate the main bands of input GST-eIF3b fragments.
and 40 S ribosome (Fig. 4A). A portion of P311 was found to be cosedimenting with small and large polysomes (fractions 6–10), suggesting that P311 might associate with mRNA present in polysomes. To determine whether the mRNAs encoding TGF-β were present in the same fraction as P311, we analyzed their relative distribution in the gradients (Fig. 4B). Compared

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with the distribution of TGF-β1 mRNA in empty vector (EV)-
transfected control cells (Fig. 4B, *green curve*, peaking at frac-
tion 8–9), overexpression of P311 shifted the TGF-β1 mRNA to
higher gradient fractions (mainly fraction 10), consistent
with the increase in TGF-β1 mRNA translation in P311-trans-
feected cells. In P311(MutER or)–transfected cells, the shift of
the peak at fraction 10 was reduced significantly (Fig. 4B, *blue
curve*, peaking at fractions 8 and 10). Similar distributions were
seen for TGF-β2 and 3 (Fig. 4B). The distribution of the β-actin
mRNA control largely overlapped among EV-, P311-, and
P311(MutER)-transfected cells (Fig. 4B). The recruitment of
TGF-β RNAs to the eIF3 complex by either P311 or
P311(MutER) was determined by RIP assays. HA-eIF3b and
MYC-P311 or MYC-P311(MutER) were cotransfected to
NIH-3T3 cells. eIF3 subunits were then cross-linked by
dimethyl 3.3’-dithiobispropionimidate · 2 HCl, a cleavable
cross-linker that stabilizes adjacent protein-protein interac-
tions (38), prior to IP of the eIF3 complex with an X-HA Ab.
These studies showed greater enrichment in TGF-β RNAs in
the eIF3 complex from cells expressing P311 than in cells
expressing P311(MutER) (Fig. 4C). Finally, reporter assays
using the 5’/3’UTR constructs of the three TGF-β isoforms
showed decreased luciferase activity in NIH-3T3 cells trans-
feected with P311(MutER) compared with P311 (Fig. 4D).

As expected from studies presented in Fig. 4, ELISA demon-
strated that P311(MutER) failed to increase TGF-β3 levels in
HLSMCs (Fig. 5A). Consistent with the role of P311 in TGF-
β1-3 translation, the levels of the corresponding mRNAs were
unchanged (Fig. 5B).

**P311 Binds to the 5’UTRs of TGF-β mRNAs through a Con-
served P311 RRM-like Signature Motif**—Although Blast and
searches in the database of European Molecular Biology Lab-
atory—The European Bioinformatics Institute (EMBL-EBI)
of the P311 aa sequence failed to identify functional motifs, a
recent analysis using BindN with 90% specificity (39) pre-
dicted a conserved RRM-like signature consisting of 13 pos-
tively charged aas (Fig. 6A, *arrowheads*) partially overlap-
ping with the EBM (Fig. 6A). To determine whether P311
directly binds to TGF-β mRNAs, NIH-3T3 cells transfected
with MYC-P311 or EV were exposed to UV light to specifi-
cally stabilize protein-RNA interaction (40). This was fol-
lowed by RIP using X-MYC Ab. These studies showed that
P311 directly binds to TGF-β mRNAs (Fig. 6B). The binding
to TGF-β mRNA was largely decreased when expressing the
P311(MutER) (all 13 potential RNA-binding sites mutated
to alanine), with no significant effect on other control
mRNAs. In vitro RNA-protein binding assays followed by
RNA-protein EMSA were performed using recombinant
P311, P311(MutER) and in vitro–produced 5’UTRs from
several mRNAs, including those that encode TGF-βs, GADPH, β-actin, 18 S RNA, and the protein product of the
TGF-β1 neighboring gene B9d2. All RNAs were confirmed
to be effective in protein binding using as a control elf4EAn,
an established RNA-binding protein (41, 42) (data not shown).
The interaction of P311 with increasing concentrations of
these various RNAs was observed by RNA-protein EMSA
(Fig. 6C). The relative density of bound P311 signal was ana-
lyzed using ImageJ software, which indicated that P311 binds
to the 5’UTR of TGF-β mRNAs with a significantly higher
affinity than to all the controls RNAs tested (Fig. 6D).

**DISCUSSION**

P311 was discovered more than two decades ago (2), but its
biological importance has only been revealed recently in studies
demonstrating its stimulatory effects on TGF-β1-3 translation
and, therefore, on blood pressure regulation (1). Because
TGF-βs play key roles in multiple pathophysiological pro-
cesses, we directed further efforts toward understanding the
molecular mechanisms by which P311 regulates TGF-β mRNA
translation.

We found that P311 is intrinsically unstructured. Therefore,
it could require a binding partner to acquire tertiary structure
and function (35). Hence, we employed transfected NIH-3T3
cells, which do not express P311 (3), as well as P311-expressing
VSMCs (1) to search for P311 binding proteins with potential
involvement in mRNA translation. Our studies led to the
identification of elf3b, a subunit of the multicomplex translation
initiation factor elf3 (43), as a direct binding partner of P311.

In co-IP mass spectrometry studies, P311 also showed an
association with several cytoskeletal proteins, including fil-
amin, previously reported to interact with P311 (44), myosin
heavy chain, and β-actin, suggesting that, besides the stimula-
tion of translation of TGF-β1-3, another P311 function may be
to link TGF-β transcripts to molecular motors for active trans-
port (45). The previously observed binding of P311 to the
TGF-β1 latent associated protein by two-yeast hybridization
and co-IP (46) was not seen in this study. One of the reasons for
this discrepancy could be that, in the previous study, latent
associated protein was introduced into NIH-3T3 cells by
digoxigenin permeabilization of the cellular membrane before
performing co-IP, a method which may have led to a large
increase of intracellular latent associated protein, facilitating its
interaction with P311.

elf3, the largest elf complex, is composed of 13 subunits,
named elf3a to elf3m (30). Several of them have been identi-
fied as core subunits (47). elf3b is one of them, whereas
the others are elf3j, elf3a, elf3g, elf3i, and elf3e (25, 48). These

![FIGURE 3. A conserved P311 aa sequence is required for elf3b binding. The sequence is referred to as EBM. A, aa sequence alignment of P311 from different species using ClustalW. Identical aas, similar aas, and nearly similar aas are shown underneath the sequences as stars, single dots, or double dots, respectively. N, N-terminal (1–22 aa); M, middle (23–46 aa); C, C-terminal (47–68 aa). B, gel mobility shift assay performed with purified GST-elf3b F4 (shown in Fig. 2C) in the absence or presence of increasing amounts of synthesized N, M, and C P311 peptides. After incubation, the proteins were resolved on a 4–20% Tris-glycine native gel, followed by Coomassie Blue staining. C, competition assay performed by GST pulldown using equal amounts of purified GST-F4 incubated with 10 μM purified P311 in the presence or absence of increasing concentrations of synthesized P311 peptides (10, 50, and 100 μM). The resulting Western blots (WB) were probed with the indicated antibodies. D, a conserved P311 region among different species. Tandem mutations were made for the following experiments. E, lysates from NIH-3T3 cells expressing either MYC-P311 with or without the indicated mutations and HA-elf3b were subjected to IP and Western blotting. F, equal amounts of the recombinant GST-elf3b F4 were incubated with the MYC-beads bound with MYC-P311 or MYC-P311(MutER). Proteins were released from the beads in SDS-PAGE loading buffer and analyzed by Western blotting.](ASBMBV289N49F3.png)
core subunits are necessary and sufficient to initiate translation (47, 49–51).

Although eIF3b is highly expressed in several forms of cancer (52–55) and its down-regulation has been shown to inhibit the proliferation and metastatic potential of colon, bladder, and prostate cancers (52, 53), little is known about eIF3b function(s) at the molecular level. Previous research indicated that eIF3b serves as a binding scaffold for the other core subunits and for eIF2 (47, 49–51), whereas its interaction with eIF3a, eIF3j, and eIF1a ensures proper formation of the scanning-arrested conformation required for stringent AUG recognition (56).

FIGURE 4. The EBM is required for P311 induction of translation of TGF-β1-3. A, cytoplasmic extracts from EV-, P311-, and P311(MutEBM)-expressing cells were fractionated through sucrose gradients, with the lightest components sedimenting at the top (fraction 1); small (40S) and large (60S) ribosomal subunits and monosomes (80S) in fractions 2–6; and progressively larger polysomes, ranging from low to high molecular weight, in fractions 6–12. Proteins on polysome gradients were detected by Western blotting. --, no ribosomal components; OD, optical density. B, polysome analysis of TGF-β1-3 mRNAs. The relative distribution of TGF-β1-3 mRNAs and β-actin mRNA on polysome gradients was studied by RT-qPCR analysis of the RNA present in each of 12 gradient fractions and is represented as percent of total mRNA. C, RIP assay showing enrichment in the 5’ UTRs of TGF-β mRNAs in the eIF3 IP from cells transfected with EV, P311, or P311(MutEBM). eIF3 complexes were subjected to IP through HA-eIF3b, and the subunits were detected using the corresponding antibodies. RNA enrichment was detected by RT-qPCR. D, luciferase reporter assays for TGF-β 5’/3’ UTRs in NIH-3T3 cells transfected with EV, P311, or P311(MutEBM). *, p < 0.05; **, p < 0.01.
It has been reported recently that eIF3b interacts with ribosomal proteins S9e, S3 and S14 (57, 58). In this regard, different ribosomal proteins, as well as several translation initiation factors, were inconsistently detected in P311 co-IP mass spectrometry studies performed under low stringent conditions, but none was identified when the co-IP stringency was increased, rendering highly unlikely the possibility that ribosomal proteins and/or translation initiation factors other than eIF3b are direct P311 binding partners. Outside of the translation machinery, eIF3b interacts with the eukaryotic cytosolic chaperonin CCP (chaperonin-containing TCP-1 (tailless complex polypeptide 1)), which promotes the correct folding of the eIF3b, eIF3h, and eIF3i subunits (59). eIF3b interaction with the RNA-binding proteins AU-rich element RNA binding protein (AUF1) and ceramide kinase-like (CERKL) has also been observed recently (58, 60). None of these proteins were, however, identified in our co-IP mass spectroscopy studies.

SPR analysis indicated that the interaction between eIF3b and P311 has a higher affinity ($K_d$ 1.26 μM) than the reported interaction between eIF3b and eIF3j ($K_d$ 20 μM (36)). Therefore, P311 could, theoretically, displace eIF3j from the eIF3 complex. However, we observed no differences in eIF3b-eIF3j binding in the presence or absence of P311, and neither the composition nor the stoichiometry of the eIF3b-associated subcomplex (eIF3a, eIF3g, and eIF3i) (30) were affected. A new study determined the affinity of eIF3j to eIF3 at $K_d$ 100 nm (61). This tight affinity supports our finding. Therefore, we ruled out the possibility that P311 may stimulate translation of TGF-β by producing qualitative and/or quantitative changes in the eIF3b-eIF3a-eIF3g-eIF3i subcomplex.

Structural studies have demonstrated previously that eIF3b folds into three independent domains (62). Its N-terminal domain contains an RRM (30, 36, 63), which, in mammals, is non-canonical (63) but rather serves as the binding site for eIF3a (30), eIF3j (36), and P311 (our study). The N-terminal domain is connected with the central WD40 domain, which contains short, ~40-aa motifs often terminating in a WD, known as WD or β-transducin repeats. Proteins with WD40 repeats are known to serve as a platform for the assembly of protein complexes (64, 65). As such, the eIF3b WD40 domain interacts with the 40 S ribosomal subunit (57). This domain is followed by the C-terminal domain, necessary for the interaction with subunits eIF3i and eIF3g (47, 66). Mutational analysis suggests that P311 binding to eIF3b involves the RRM. This finding supports the possibility that eIF3b-bound P311 is located in close proximity to eIF3a and eIF3j. Our studies, however, revealed no P311 interaction or competition with either of them, indicating that the RRM region of eIF3b has multiple binding partners. RRM-containing genes represent about 0.5–1% of all human genes (67). Therefore, if the RRM represents the only P311 binding motif in the context of mRNA translation, the number of transcripts regulated by P311 should be relatively small. On the other hand, we identified the eIF3b binding region in P311 as a centrally located, 11-aa-long sequence (from leucine 30 to lysine 40), highly conserved among eukaryotic species, which we named EBM.

Systematic biochemical and functional studies demonstrated that EBM-mediated binding to eIF3b is required for P311 stimulation of TGF-β1-3 translation. The requirement of EBM-mediated P311-eIF3b interaction was confirmed by observing that expression of P311, but not P311(MutEBM), increased TGF-β1-3 levels in HLSMCs without increasing their corresponding mRNAs.
The specific requirement for elf3b in P311-mediated translation of mRNAs encoding TGF-β1-3 has been demonstrated in studies in which elf3b or elf3a control were down-regulated by RNAi in the presence or absence of P311. In such studies, P311 stimulated TGF-β translation when this was inhibited as a result of elf3b down-regulation, but P311 had no effect on TGF-β translation when inhibited by down-regulation of elf3a. Although Blast and EMBL-EBI searches failed to identify distinct functional motifs in P311, a BindN search with 90% specificity (39) detected 13 potential RNA-binding residues distributed throughout its length, opening the possibility that P311 may bind RNA. Our in vivo and in vitro studies demonstrated that, indeed, P311 directly binds to TGF-β1-3 mRNAs but not to other mRNA controls. The binding was reduced significantly when substituting P311 for P311(MutRRM).

During the process of translation initiation, several elf3 subunits interact with RNA. The subunit elf3j directly binds to the 40 S ribosomal RNA (68), and the subunits elf3a, elf3c, and elf3d physically interact with the 5′UTRs of mRNAs (69–72). Therefore, one or more of these subunits is likely to interact with TGF-β 5′UTRs, explaining why translation of TGF-β is not completely shut down in the absence of P311 (1).

As mentioned above, two mRNA-binding proteins, AUf1 and CERKL, also interact with elf3b. AUf1 binding with elf3b, along with S6 and S14, promotes the circadian translation of cryptochrome (58), and although the normal CERKL binds to elf3b, the C125W CERKL, a pathological mutant that causes retinitis pigmentosa and cone-rod dystrophy, loses its interaction with elf3b (60). To date, however, no other mRNA binding protein has been shown to display activities similar to that we now ascribe to P311.

In conclusion, our findings indicate that, by concomitantly binding to elf3b through its EBM and to the 5′UTR of TGF-β1-3 mRNAs through its RRM, P311 recruits TGF-β1-3 mRNAs to the translation machinery, which, in turn, leads to the co-IP-mass spectrometry studies because, in such a case, the effect on translation would be global.
increased translation of TGF-βs and to the up-regulation of TGF-β1-3 levels.

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