elf2β is critical for elf5-mediated GDP-dissociation inhibitor activity and translational control

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ABSTRACT

In protein synthesis initiation factor elf2 binds initiator tRNA to ribosomes and facilitates start codon selection. elf2 GDP/GTP status is regulated by elf5 (GAP and GDI functions) and elf2B (GEF and GDF activities), while elf2α phosphorylation in response to diverse signals is a major point of translational control. Here we characterize a growth suppressor mutation in elf2β that prevents elf5 GDI and alters cellular responses to reduced elf2B activity, including control of GCN4 translation. By monitoring the binding of fluorescent nucleotides and initiator tRNA to purified elf2 we show that the elf2β mutation does not affect intrinsic elf2 affinities for these ligands, neither does it interfere with elf2 binding to 43S pre-initiation complex components. Instead we show that the elf2β mutation prevents elf5 GDI stabilizing nucleotide binding to elf2, thereby altering the off-rate of GDP from elf2•GDP/elf5 complexes. This enables cells to grow with reduced elf2B GEF activity but impairs activation of GCN4 targets in response to amino acid starvation. These findings provide support for the importance of elf5 GDI activity in vivo and demonstrate that elf2β acts in concert with elf5 to prevent premature release of GDP from elf2γ and thereby ensure tight control of protein synthesis initiation.

INTRODUCTION

In eukaryotic protein synthesis initiation the guanosine-5′-triphosphate (GTP) binding protein elf2 delivers initiator tRNA (Met−tRNAi) to the small ribosomal subunit as part of the 43S pre-initiation complex (PIC). Following mRNA selection elf2 also assists with AUG codon selection during scanning to ensure accurate protein synthesis (1,2). The affinity of elf2 for Met−tRNAi is dictated by its nucleotide status as elf2•GTP has high affinity for Met−tRNAi, while elf2•GDP does not (3). Upon GTP hydrolysis and phosphate release elf2•GDP loses affinity for tRNA and leaves the PIC (4). Hence elf2 must reacquire GTP to facilitate continued rounds of translation initiation. The release of GDP from elf2 is promoted by the guanine nucleotide exchange factor (GEF) elf2B; an important mediator of protein synthesis control (5).

elf2B GEF activity is specifically inhibited to modulate protein synthesis initiation rates in response to diverse cues across all eukaryotes studied. Signals include nutritional imbalances (6), endoplasmic reticulum stress and modulation of learning and memory (7–11), which activate one of four protein kinases that each phosphorylate elf2α at serine 51 elf2α(pS). This converts elf2 into a competitive inhibitor of elf2B GEF (12–14), by promoting tight binding of elf2α to the αβ subunits of elf2B (15), presumably in a manner such that the GEF domain at the elf2B C-terminus (16) cannot access elf2γ-bound GDP for nucleotide exchange. Elevated levels of elf2α(pS) lowers overall translation and activates translation of a subset of translationally controlled genes such as those bearing upstream open reading frames (uORFs) including GCN4 in yeast and ATF4 in mammals (17,18). Glia appear particularly sensitive to reductions in elf2B activity because missense mutations in elf2B subunits cause the fatal neurological disorder Leukoencephalopathy with vanishing white matter (VWM; OMIM #603896) (19).

Hydrolysis of elf2•GTP within the PIC is promoted by the GTPase activating protein (GAP) elf5 (4), and upon AUG-codon recognition, elf5/elf2•GDP complexes dissociate prior to 60S joining (20). elf5/elf2•GDP complexes represent an abundant complex in yeast cells (21,22). elf5 has a second function within this complex to prevent spontaneous release of GDP from elf2, termed GDP disso-

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cation inhibitor (GDI) activity (23). eIF5 GDI antagonizes eIF2B GEF and is necessary for tight control of translation by eIF2αP by ensuring that there is minimal eIF2B independent GDP exchange. This GDI function requires the eIF5 C-terminal domain (CTD) and the adjacent upstream linker region (LR). Mutations in either domain that weaken eIF5–eIF2B interactions eliminate GDI activity (23). When eIF2B-free active eIF2B is present, it can efficiently displace eIF5 from eIF2αGDP. Consistent with other G protein systems, this additional role for eIF2B is termed a GDI-displacement factor (GDF) function (14). Thus eIF2B GDF activity displaces eIF5 GDI from eIF2αGDP to enable eIF2B GEF and continued rounds of protein synthesis (2,24).

Here we present a molecular analysis of a growth suppressor mutation that spontaneously arose during our studies of eIF2B GEF mutants. We show that it is a novel missense allele within the β subunit of eIF2 (E189K). Biochemical analyses reveal that this mutation does not affect the intrinsic nucleotide, Met–tRNAi, or 43S PIC-binding affinities of eIF2. Instead we find that it prevents eIF5 GDI activity. Our genetic studies show that this alters the sensitivity of cells to eIF2B activity and eIF2αP, thereby affecting the derepression of GCN4 translation. These results uncover an important role for eIF2β–eIF5 interactions for controlling eIF5 GDI activity, suggesting that eIF2β E189 makes important contributions to this function. They also demonstrate that eIF2β is important for determining the cellular responses to the eukaryote-wide eIF2 translation control mechanism.

**MATERIALS AND METHODS**

**Yeast genetics and cell culture**

Strains used and generated in this work are derivatives of S288c and are described in Supplementary Table S1. Strains used and generated in this work are derivatives of S288c and are described in Supplementary Table S1. Yeast genetics and cell culture

**MATERIALS AND METHODS**

**Genomic DNA sequencing**

GP3755 and GP3763 genomic DNA was sequenced using a SOLiD4 sequencer (Life Technologies) at the University of Manchester genomics technologies facility using standard kits and protocols. To generate paired-end reads (F3 [start of DNA fragment] = 50 bp and F5 [end of DNA fragment] = 35 bp) the CSFASTA and QUAL files for each sample were converted to FASTQ format using the solid2fastq script (BFAST software) (27,28). The reads were quality filtered using SOLID_preprocess_filter_v2.pl (29) (settings used -x y -p 3 -q 22 -y y -e 10 -d -9). The reads were mapped to the *Saccharomyces cerevisiae* reference genome ‘sacCer3’ obtained from UCSC (30) (http://genome.ucsc.edu), which includes the assembled chromosomes, the mitochondrial genome and the two micron plasmid. The F3 reads were mapped using BFAST ‘match’ and the F5 reads were mapped using BFAST + BWA 0.7.0.a ‘bwaaln’. Mapping was completed as single-ended using the F3 reads only, or the individually mapped F3 and F5 reads were both used for paired-end mapping. In both cases the ‘-a 2’ flag of the post-process step was used to obtain uniquely mapping alignments. The F3 reads of properly paired reads were obtained using ‘samtools view -f 66’ (31). SRMA v0.1.15 was used to realign reads to correct for erroneous mapping caused by the presence of INDELS at the end of reads. Variant detection was performed using samtools v0.1.18, using mpileup and bcftools without BAQ computation (32,33). Candidate variants where filtered using vcfutil.pl requiring at least 20 high-quality read matches and a maximum coverage calculated as three or five times the median coverage (including 0 counts). Variants were also filtered to remove any with QUAL (variant quality) <40.

**Verification sif285-1 is sui3-E189K**

Genomic DNA was isolated from yeast strains GP3771-4 using standard procedures (25) and the *SUI3 ORF am-
plified by PCR using primers eIF2betaF1 (ATGTCCTC- CGATTAGCTGC) and eIF2betaR286 (TCACATTCTC- CTTCTTTACC). PCR products were screened by BspE1 digestion and verified by Sanger DNA sequence analysis.

Sequence alignment
Alignment generated using Clustal X (34) and the following eIF2B sequences (termed EIF2S2, SU13 or TIF212) from Genbank: S. cerevisiae (CA97959.1), Candida albicans (XP_715210.1), Candida glabrata (XP_446841.1), Cryptococcus neoformans (AAW43820.1), Schizosaccharomyces pombe (CAB11076.2), Caenorhabditis elegans (CAB00049.1), Homo sapiens (CAG33015.1), Mus musculus (AA113767.1), Arabidopsis thaliana (AAK29672.1), Oryza sativa (NP_001060649.1), Glycine max (XP_006575455.1).

Protein synthesis
The rate of protein synthesis was measured in exponential phase cells in SC-uracil-methionine at 30°C in a water bath. Once cells (10 ml cultures in 50 ml flasks) reached A600 = 0.1, 160 μl methionine mix was added (1 μl [35S] methionine (1175 Ci/μmol Tran35S-label, MP Biomedicals) in 50 μM methionine). 100 μl samples were removed immediately and every 20 min over a 2 h period and were immediately precipitated with ice-cold 20% trichloroacetic acid (TCA). Total protein incorporation was measured by scintillation counting of TCA-precipitates absorbed onto GF/C filters (Whatman) washed with ice-cold acetone.

For Polysome profile analysis yeast strains were grown to an A600 = 0.7 in 100 ml YPD medium. For glucose starvation, cells were collected by rapid centrifugation at room temperature each time. Each measurement was blanked against a control without nucleotide to account for any affect of eIF2 and data were corrected for dilution effects caused by volume addition and normalized to starting values before being fitted to a single site binding model: y = 1 + [(Fmax - 1)*x/(x + Kd)].

Steady state fluorescence
To assay nucleotide affinity, fluorescence intensity of 100 nM of BODIPY-FL-GDP or BODIPY-FL-GTP (Thermo Fisher Scientific) in 180 μl of assay buffer (30 mM HEPES, 100 mM KCl, 1 mM DTT, 1 mM EDTA, 10% glycerol, pH 7.4) was measured using a Fluoromax-4 spectrophotometer (Horiba) (490 nm excitation, 509 nm emission). Yeast Met-tRNAi was assayed in a similar manner using 20 nM BOP-N-Met-tRNAi (tRNA probes) but with the addition of 1 mM GTP. Change in fluorescence intensity was measured upon addition of increasing amounts of apo-eIF2, incubating for 5 min at room temperature each time. Each measurement was blanked against a control without nucleotide to account for any affect of eIF2 and data were corrected for dilution effects caused by volume addition and normalized to starting values before being fitted to a single site binding model: y = 1 + [(Fmax - 1)*x/(x + Kd)].

GDP dissociation assay
Fluorescent eIF2•BODIPY-GDP binary complex was formed by incubating apo-eIF2 with a 2× excess of BODIPY-FL-GDP (Thermo Fisher Scientific) for 20 min at room temperature. Excess nucleotide was removed by passing through a G-50 sephadex column (GE Healthcare). Labelling efficiency was calculated to exceed 90%. To measure GDP release, 20 nM eIF2•BODIPY-GDP was quickly mixed with 1 mM of unlabelled GDP (±eIF2B and ±GST-eIF5) in 180 μl of assay buffer (30 mM HEPES, 100 mM KCl, 10 mM MgCl2, pH 7.4) and fluorescence intensity was continuously measured using a Fluoromax-4 spectrophotometer (Horiba) (490 nm excitation, 509 nm emission, 0.1 s integration time). Experimental data were fitted to exponential dissociation curves to determine the rate constants (Koff) at each eIF2B concentration. K1/2 and Kmax values

GST pull-down assay
Purified GST-eIF5 or GST alone (200 pmol) and eIF2 (200 pmol) were incubated in 500 μl of interaction buffer (30 mM HEPES pH 7.5, 100 mM KCl, 0.1 mM ethylenediaminetetraacetic acid (EDTA), 1 mM 1,4-dithiothreitol (DTT), 2.5 mM MgCl2) with 25 μl of glutathione-Sepharose beads for 1 h at 4°C. Beads were washed three times with 500 μl of interaction buffer and bound proteins were eluted by boiling in protein-sample buffer. Western blotting was performed as above.

Protein purification
eIF2 was purified from yeast strains GP7145 and GP7146 expressing V5-su13-E189K or the V5-SU13 derivatives of eIF2B by sequential nickel affinity, HiTrap Heparin and HiTrap Q sepharose chromatography as described (14). To obtain apo–eIF2 free from nucleotide, eIF2B was dialysed with EDTA (30 mM HEPES, 100 mM KCl, 1 mM DTT, 1 mM EDTA, 10% glucose, pH 7.4) then with magnesium (30 mM HEPES, 100 mM KCl, 1 mM DTT, 0.1 mM MgCl2; 10% glucose, pH 7.4). GST-eIF5 was purified from Escherichia coli as described (23). eIF2B was purified from yeast strain GP5949, as described (39).

β-galactosidase assay
Measurement of β-galactosidase levels from a HIS4-LacZ fusion genomically-integrated at ura3-52 was performed exactly as described (36) using whole protein extracts from cells grown in SC medium lacking histidine and uracil or following addition of 10 mM 3AT for 6 h. Units are nmol o-nitrophenyl-β-D-galacto-pyranoside cleaved/min/μg total protein ± SE, n = 3.

FLAG immune precipitation and western blotting
FLAG (Sigma-Aldrich) immune precipitation and western blotting were performed as described recently (23,37,38) using whole cell extracts to capture FLAG-eIF2B and associated proteins. Bound proteins were probed using specific antibodies and quantitative IR western blot detection was performed using IRDye® 800CW goat anti-rabbit IgG or IRDye® 680RD goat anti-mouse IgG with an Odyssey Fc imager (Li-Cor).
were determined from curve fitting $y = (K_{\text{max}} \times x)/(K_{\text{1/2}} + x) + c$. Yeast eIF2B was recently shown to be a functional dimer (40), so likely interacts with eIF2 as a 2:2 complex. For consistency with prior studies, eIF2B and eIF2 concentrations stated and equations used refer to 5-subunit and 3-subunit monomeric complexes respectively.

RESULTS

$sif285-1$ is a spontaneous growth suppressor of eIF2B mutations

A spontaneous suppressor of the slow-growth of one yeast eIF2Be mutated strain (gcd6-F250L) was noted following routine sub-culture on solid medium (Supplementary Figure S1A). The suppressor-bearing strain carries a deletion of the essential GCD6 gene and is complemented with a plasmid borne gcd6-F250L allele. Following transformation and plasmid shuffling the growth-suppressor strain showed a clear difference in growth rate compared to wild-type (WT) cells when both were combined with the gcd6-F250L mutation which grow slowly in the WT background and grow well in the growth-suppressor strain. The unknown mutation was hence initially named $sif285-1$ for suppressor of eIF2B5 and the WT version termed $SIF285$ (Figure 1A and Supplementary Figure S1B). This name is used in the description of our analyses here prior to the identification the molecular defect causing the mutation.

Measurements of growth-rates in liquid cultures, total protein synthesis by $^{35}$S-methionine incorporation and polysome profile analyses showed the defect in eIF2B activity conferred by the F250L mutation affected general protein synthesis in an otherwise WT strain, as expected from previous studies (41). In contrast, these defects were largely suppressed by the presence of the $sif285-1$ mutation (Figure 1A–C). $sif285-1$ was also able to restore faster growth to all slow-growing gcd6 mutations tested (Supplementary Figure S1C), including suppressing the lethality of gcd6-N249K, a Gcd6 mutation which retains residual GEF activity that is insufficient to permit growth of otherwise WT cells (41). However, $sif285-1$ could not rescue a complete deletion of GCD6. Mating to create a/a diploid cells showed that the $sif285-1$ phenotype was semi-dominant (Supplementary Figure S1D). Sporulation and tetrad dissection showed the fast-growing phenotype segregated 2:2 (Supplementary Figure S1E), indicating that $sif285-1$ was likely a single gene alteration. Mutations in multiple factors can cause a loss of translational control, including mRNA-decay pathway mutants (42). To assess if $sif285-1$ is generally resistant to translational control responses, we monitored the initial cell responses to glucose starvation by polysome profile analysis following 10 min glucose withdrawal, as this response is independent of eIF2B inhibition (43). $sif285-1$ mutant cell translation remains glucose-withdrawal sensitive, suggesting $sif285-1$ suppression has at least some eIF2B specificity (Figure 1D). Because yeast eIF2B mutations constitutively derepress GCN4 translation and the Gcn4-mediated general amino acid control response (GAAC) independent of Gcn2 (termed a Gcd+ phenotype), we assessed resistance of $sif285-1$ cells to 3AT, a competitive inhibitor of histidine biosynthesis that activates GAAC via the signalling pathway (box in Figure 1F) (17). $sif285-1$ could not prevent the Gcn2 independent 3AT resistance of gcd6Δ gcd6-F250L cells in a replica plating assay (Figure 1E). To examine if $sif285-1$ cells alter the Gcn2-dependent GAAC response, we transformed these cells with a WT Gcn2 plasmid and repeated our growth analyses. $sif285-1$ Gcn2 cell growth was impaired by 3AT, indicating that $sif285-1$ was a GACN4 non-derepressible mutant (a Gcn+ phenotype, Figure 1F).

This observation was confirmed by analysing the levels of β-galactosidase expressed from a genomically integrated HIS4-lacZ for strains shown in (A), but following transformation with a Gcn2 plasmid (pAV1198). Right box summarizes the signalling pathway leading to derepression of HIS4 expression by amino acid starvation (+3AT).

Figure 1. An extragenic spontaneous slow-growth suppressor of eIF2B mutations. (A) $sif285-1$ suppresses the slow-growth of the gcd6-F250L mutation. Growth of strains GP3771-3774 on YPD. Relevant genotypes and doubling times from liquid cultures are shown. (B) Total protein synthesis rates ($^{35}$S-met incorporation) ± SE (n = 3). Student’s t-test indicates only F250L differs significantly from the other three strains (P < 0.05). (C and D) Polysome profile analysis for strains shown in (A) grown in YPD or following 10 min incubation in YP lacking glucose (D). Traces show A$_{254}$ recorded by continuous fractionation of 15–50% sucrose gradients. (E) Replica printing of the strains shown in panel A on minimal SD ±3AT. (F) Growth on SD ±3AT media and β-galactosidase assay of HIS4-lacZ for strains shown in (A), but following transformation with a Gcn2 plasmid (pAV1198). Right box summarizes the signalling pathway leading to derepression of HIS4 expression by amino acid starvation (+3AT).
that the *sif285-1* mutation permits cells to grow with reduced requirement for eIF2B that alters the threshold for induction of the GAAC response, but does not confer complete insensitivity to eIF2B. Because mutations in human eIF2B that lead to VWM usually impair eIF2B GEF activity and can cause heightened ATF4 expression (44,45), we decided it would be of interest to characterize the *sif285-1* mutation further. Standard complementation experiments showed the *sif285-1* mutation was not in any of the five eIF2B subunits.

**sif285-1 is E189K in eIF2β (SU3)**

To identify the *sif285-1* defect we isolated genomic DNA from the original *sif285-1* strain (GP3563) and its parent (strain GP3755) and sequenced it by SOLiD4 next generation sequencing (ABI). Following mapping to the reference *S. cerevisiae* full genome (UCSC version sacCer3), the reads were screened for single nucleotide variants (SNV) and deletions using standard informatics approaches. SNV analysis revealed only seven clear differences between the WT and *sif285-1* genomes that had at least 20 high-quality mapped reads (Supplementary Table S3). Only two SNVs corresponded to missense mutations; the first change *gcd6-F250L* was known and expected, while the second caused an *E189K* change in *SU3*. *SU3* encodes for the eIF2β subunit: a known interacting partner of Gcd6/eIF2Be (26), and was therefore the likely *sif285-1* candidate. The missense mutation was predicted to eliminate a BspE1 restriction site in *SU3*. The *SU3* ORF was amplified by PCR from the strains and both BspE1 digestion (Supplementary Figure S2) and Sanger DNA sequencing (Figure 2A) verified that only strains bearing *sif285-1* carry the *sui3-E189K* mutation.

We performed a series of experiments to confirm that the slow-growth-suppression and Gen− phenotypes of *sif285-1* were allelic with an *E189K* mutation in *SU3*. Firstly, both *CEN* and two micron plasmids bearing WT *SU3* were transformed into WT and *sif285-1* strains. The fast growth rate of *sif285-1* *gcd6-F250L* double mutant suppressor strains is complemented by additional copies of *SU3*. Thus slower growth is restored to *sif285-1* cells bearing *gcd6-F250L* and *SU3* plasmids (Figure 2B; compare rows 8 and 12 with row 4), as expected if *sif285-1* is a mutation in *SU3*. Secondly, site-directed mutagenesis introduced *sui3-E189K* onto a plasmid that was shuffled into a ‘clean’ *sui3Δ* strain as the sole source of eIF2B. The resulting strain grew as WT, indicating that like *sif285-1*, *sui3-E189K* does not confer a significant growth defect (Figure 2C). Thirdly, examining GAAC in these strains revealed that *sui3-E189K* confers a Gen− phenotype and fails to derepress HIS4-lacZ expression in response to 3AT treatment (Figure 2C) analogous to the *sif285-1* results shown in Figure 1F. Finally we created a *gcd6Δ sui3Δ* double knock out strain and complemented each deletion with either WT or mutant versions of each gene. These strains re-created the original slow-growth phenotype of *gcd6-F250L* and its suppression by *sui3-E189K* (compare Figure 2D with Figure 1A). Taken together, these data are fully consistent with *sui3-E189K* being the cause of the *sif285-1* suppressor phenotype.

**sui3-E189K conservation**

The region of eIF2β around E189 is highly conserved across diverse eukaryotes (Figure 3A). E189 itself is acidic (E or D) in plants and lower eukaryotes, while the equivalent residue is already a lysine in animal sequences including humans. The significance of this evolutionary difference is not clear. E189 is located away from mutations studied previously that exhibit different phenotypes and interaction defects (Figure 3B). The eIF2β amino-terminal poly-lysine blocks (termed ‘K-boxes’) are important for stable interactions with both eIF5 and eIF2Be (26). Nuclear magnetic resonance studies showed K-box 2 binds directly to the eIF5 carboxy-terminal domain (CTD) (46). The eIF2β-CTD residue S264 has been implicated in stabilizing GTP binding and regulating GTP hydrolysis as S264Y (SU3-2) permits enhanced initiation at a non-AUG start codons (47).

Recent structural analysis of the yeast PIC included structural determination for eIF2β (26) and its complex bound with other initiation factors and a short AUG containing mRNA to the 40S ribosome (48). In this model the eIF2β E189 side chain projects from the surface of eIF2B, permitting potential ligand modulation. In the structural model, conserved residues close to eIF2β E189 appear to stabilize Met-tRNAi binding to eIF2GTP, while residues 218-220 contact eIF1A and ribosomal protein uS19 interacts with
residues 166-169 at a different eIF2β surface. These data indicate that eIF2β makes multiple important contacts within the PIC (48) and the relative positioning of E189 suggests that E189K could alter ligand binding to eIF2 (Figure 3C). We therefore decided to perform a series of experiments to determine the biochemical impact of E189K on eIF2 ligand interactions.

E189K does not affect GDP, GTP or tRNA affinity for eIF2

WT and mutant eIF2 (eIF2βE189K) were purified from an overexpression yeast strain engineered so that the introduced plasmid was the sole source of eIF2β (Supplementary Table S1 and Figure S3A). During purification incubation with EDTA was used to remove residual bound magnesium and nucleotide, as described previously (3). To assess nucleotide and Met–tRNAi binding affinities we monitored the change in fluorescence upon binding of eIF2 to ligands labelled with BODIPY derivatives at room temperature (Figure 4A). These experiments show that BODIPY-GDP and BOP-N-Met–tRNAi affinities for WT eIF2 are within the ranges measured previously by other techniques (K_GDP = 19.1 nM and K_Met–tRNAi = 2.8 nM) (3,49). Measured BODIPY-GTP affinity is tighter than when GTP-binding was measured by filter-binding assays (3,49), so that the difference between GDP and GTP affinities is significantly less than noted previously. Perhaps surprisingly, the affinities for each ligand are unchanged by the E189K mutation (Figure 4A). Next, we pre-bound BODIPY-GDP to eIF2 or eIF2βE189K, and purified these complexes away from unbound BODIPY-GDP. By incubating each complex with excess unlabelled GDP, we monitored the release of BODIPY-GDP and determined its off-rate. The K_off BODIPY-GDP was identical for both WT and eIF2βE189K (WT K_off = 0.292 ± 0.025 min⁻¹ and eIF2βE189K K_off = 0.293 ± 0.015 min⁻¹; Figure 4B, 0 nM eIF2B). These results demonstrate that eIF2β-E189K mutation does not simply alter the binding of either nucleotide or Met–tRNAi. Hence eIF2βE189K does not share defects in ligand interactions described previously for other eIF2 mutations (47,49).

E189K modestly reduces eIF2B GEF activity, but not PIC interactions

Because the E189K mutation suppresses the slow-growth of eIF2B mutants with reduced GEF activity, one possible mechanism of mutant action was the enhancement of eIF2B activity. We tested this idea by adding increasing concentrations of purified eIF2B (39) to purified eIF2βBODIPY-GDP to determine K₄ and K₃₅ values for eIF2B GEF.

The measured K_max for WT eIF2 was ~14-fold faster than the rate determined previously with a filter-binding assay (14). This likely reflects that these fluorescent assays were performed at room temperature rather than 0°C used previously. Surprisingly, the rate of nucleotide exchange from eIF2βE189K complexes was modestly impaired relative to that with WT eIF2, increasing K₄ from 51.7 to 92.6 nM eIF2B (Figure 4B). This experiment identified the first clear difference between WT and eIF2βE189K complexes, but because impairment of eIF2B function by E189K should exacerbate rather than suppress the growth defect of eIF2B mutants, this observation cannot readily account for the growth suppression phenotype uncovered.

As outlined above, eIF2B contacts other ligands within the PIC including eIF5, 40S ribosomes, eIF3 and eIF1. To examine eIF2 protein interactions more widely, we performed anti-FLAG immunoprecipitation of FLAG-eIF2B from WT and E189K cells and assessed the ability of known eIF2-interacting partners to co-associate by quantitative western blotting. The only significant defect observed was a small, but statistically significant, 20% reduction in association of eIF2B subunits with eIF2β-E189K observed upon quantification of the signals (Figure 4C). Together with the modest reduction in activity seen in the GEF assays, these results suggest that one consequence of the eIF2B mutation is impaired eIF2B binding and activity. In contrast, binding of eIF2 to PIC factors eIF5, eIF3, eIF1 and 40S ribosomes appears unchanged (Figure 4C), consistent with the growth characteristics of eIF2β-E189K cells. These data support the idea that the E189K mutation does not significantly, if at all, interfere with eIF2 functions in PIC formation, and scanning/AUG recognition, but does modestly reduce eIF2B interaction and activity.
eIF2β-E189K eliminates eIF5 GDI activity

We found previously that eIF5 mutants that impair its GDI activity can partially suppress slow-growth phenotypes of eIF2B mutants and confer a Gcn- phenotype (14,23) in a manner similar to that seen with eIF2β-E189K and eIF2B alleles, including rescue of the lethality of the gcd6-249K mutation (Supplementary Figure S1C). This suggests a link between eIF2β-E189K and eIF5 GDI mutants. However, the eIF5 GDI mutants characterized act, at least in part, by reducing the affinity of eIF5 for eIF2 (23), whereas the eIF2β-E189K mutation does not reduce eIF5 interaction in vivo (Figure 4C) or in vitro (Supplementary Figure S3B). These observations imply that reduced eIF5 binding to eIF2 does not explain the suppressor phenotype. We therefore decided to assess eIF5 GDI activity directly. We monitored eIF5 GDI activity by incubating eIF2β•BODIPY-GDP (20 nM) with purified eIF5 (0–80 nM) and recorded the fluorescence change upon dissociation of BODIPY-GDP from eIF2. GDP-binding to WT eIF2 was stabilized by eIF5 in line with previous observations implying that a 1:1 eIF2β•BODIPY-GDP/eIF5 complex forms that stabilizes GDP binding to eIF2 (23). Importantly, GDP binding to eIF2β-E189K was not stabilized by eIF5 even at excess eIF5 levels (Figure 5A). Hence the experiment shows that eIF2β-E189K eliminates eIF5 GDI activity and this finding represents the likely cause contributing to the original gcd6-250L slow-growth suppression phenotype observed in sif285-1/sui3-E189K mutant cells (Figure 1).

To examine the impact of eIF5 on eIF2B GDF and GEF activities, we monitored BODIPY-GDP release from eIF2 pre-bound to eIF5 over a wide range of eIF2B concentrations. In line with previous observations (14) GDP release from WT eIF2 was inhibited at low eIF2B concentrations by eIF5 GDI. At higher eIF2B levels GDP release is unaffected by eIF5 because excess eIF2B can fully displace eIF5, nullifying GDI (compare blue [+eIF5] with black [–eIF5] in Figure 5B, left panel). In contrast, even at low eIF2B concentrations the kinetics of GDP-release from eIF2β-E189K are completely unaffected by the presence of eIF5 (compare green symbol with red symbols in Figure 5B, zoomed right panel). Our assays contain 20 nM each of eIF2β•GDP and eIF5, so eIF2B concentrations below 8 nM likely reflect factor ratios in vivo.
Figure 5. E189K antagonizes eIF5 GDI activity. (A) Kinetics of BODIPY-FL-GDP release from preformed purified WT and mutant (β E189K) eIF2α-GDP complexes (20 nM) with varying concentrations of GST-eIF5 ± SD (n = 3). Molar eIF2α-GST-eIF5 protein ratios are shown. Asterisks (*) mark points with statistically significant difference to WT ($P < 0.01$, unpaired Student’s $t$-test). (B) Kinetics of BODIPY-FL-GDP release from preformed eIF2 complexes (20 nM) in the presence of different eIF2B concentrations and eIF5 (20 nM). Curves shown in Figure 3B are repeated for direct comparison. Right panel shows a zoomed image focussing on eIF2B 0–12.5 nM ± SD (n = 3). Statistical significant increase in the rate of GDP release from eIF5/eIF2 WT relative to eIF5/eIF2 βE189K is indicated (∗ = $P < 0.01$, † = $P < 0.05$, unpaired Student’s $t$-test). (C) Growth and HIS4-LacZ expression of E189K and WT eIF2 cells (strains GP3773 and GP3771) transformed with plasmids carrying the indicated constitutively active GCN2 alleles.

The increased rate of GDP release from eIF2$^{\beta E189K}$ observed with 5 nM or less eIF2B is statistically significant (points marked * or † in Figure 5B, right panel) and is fully consistent with the slow-growth suppression phenotype observed in cells where low eIF2B GEF activity normally limits growth-rate (Figures 1 and 2). To conclude, our data indicate that eIF2$^{\beta}$-E189K eliminates eIF5 GDI activity and thereby facilitates faster rates of nucleotide exchange when eIF2B activity is limiting. This is consistent with our observations of enhanced protein synthesis rates and enhanced rates of cell growth in eIF2$^{\beta}$-E189K/sif285-1 cells bearing defective eIF2B mutants with lower eIF2B GEF activity.

Physiologically eIF2B activity is reduced in vivo by phosphorylation of eIF2α at serine 51. A prediction of our genetic (Gcn− phenotype) and biochemical results (loss of GDI activity) is that eIF2$^{\beta}$-E189K should reduce the sensitivity of cells to very high levels of phosphorylated eIF2.

We tested this hypothesis by transforming cells with plasmids expressing one of two constitutively-active alleles of the Gcn2 kinase (GCN2C) that each cause high levels of eIF2(αP) and result in severe slow-growth (51). In accord with our predictions, eIF2$^{\beta}$-E189K reverted the slow-growth of both GCN2C mutants tested and lowered the aberrantly high expression of the HIS4-LacZ reporter seen in SUI3 GCN2C cells (Figure 5C). Our data demonstrate that eIF2$^{\beta}$-E189 and GDI are critical for the normal tight control of translation and GAAC by eIF2(αP).

**DISCUSSION**

In this study, we have described the serendipitous discovery and subsequent genetic identification and molecular analysis of a growth suppressor mutation that spontaneously arose during our studies of eIF2B GEF mutants. sif285-
J is a novel missense allele within the β subunit of eIF2 (E189K) that does not affect the intrinsic nucleotide, Met–tRNA, or 43S PIC-binding affinities of eIF2. Instead we find that it prevents eIF5 GDI activity. This single amino acid change alters the sensitivity of cells to reduced eIF2B activity thereby repressing the cellular GAAC response to elevated eIF2α. Taken together our data show that eIF2β is critically important for guiding the GDI activity of eIF5. They also demonstrate that eIF2β is important for determining the normal cellular responses to eIF2α involving activation of GCC4 target genes during activation.

sui3-E189K is the first eIF2 mutation described that prevents stabilization of GDP binding to eIF2γ by eIF5 GDI. Previously described mutations in eIF2 subunits typically have different defects in protein synthesis. For example mutations in eIF2γ that directly alter its affinity for GTP/GDP or Met–tRNA, (49); Sui− mutations in each of the three eIF2 subunits that enhance initiation at non-AUG codons (47,52–54); mutations that affect subunit interactions and impair eIF2 complex integrity (55). All these mutations impair cell growth and protein synthesis. In addition, the three repeated lysine blocks (K-boxes) within the amino terminus of eIF2β have been found critical for binding to both eIF5 and eIF2B and their complete deletion is lethal in vivo (26,46,56). sui3-E189K shares growth properties with Gcn− mutations in eIF2α that alter the ability of eIF2 to respond to stress/GAAC signalling. These alleles were shown to overcome the inhibition of translation by phosphorylated eIF2 by either impairing interactions between eIF2α and eIF2 kinases or by weakening the tight binding of phosphorylated eIF2 to eIF2B (57,58). Gcn− mutations have also been found in other PIC factors including eIF1A (59), eIF3 subunits (60) and eIF5 (61). GCN4 reporter gene analyses of these Gcn− mutations revealed defects in uORF1 AUG codon recognition (leaky-scanning) or scanning contributions to the Gcn− phenotypes. However these mutants also cause general defects in protein synthesis leading to aberrant polysome profiles and/or poor rates of growth, presumably caused by widespread initiation defects in one or more of the steps in protein synthesis initiation on many mRNAs. sui3-E189K does not share these global translation defects (Figures 1 and 2) and in contrast improves growth of otherwise sick eIF2B mutant strains. In addition our biochemical analyses (Figure 4) indicate that the E189K mutation does not significantly, if at all, interfere with eIF2 interactions with PIC factors. This is again in contrast to similar analyses with mutants that do interfere with function in PIC formation, scanning or AUG recognition (59–61). Hence sui3-E189K is the first Gcn− mutant in eIF2β and defines a novel class of eIF2 mutation that impairs eIF5 GDI function (Figure 5A).

Previously we studied features of eIF5 necessary for GDI activity. We found that GDI requires the eIF5-CTD that interacts with eIF2β, as well as the eIF5 LR containing a conserved DWEAR motif that binds to eIF2γ (23). Mutations within the eIF5-CTD and DWEAR motif were found to weaken eIF5 binding to eIF2 and to eliminate GDI (23). Those data suggested that eIF5 GDI was functioning primarily via protein–protein interactions with eIF2β and γ. As eIF2γ binds GDP we suggested that it was eIF5–eIF2γ interactions that were most critical for GDI. However, our new data imply that eIF2β has a more critical role in mediating eIF5 GDI, and does more than simply providing an anchoring point for eIF5 through eIF5-CTD–eIF2β K-box interactions, because eIF5 binds to eIF2βE189K as well as it does to WT eIF2 (Figure 4C and Supplementary Figure S3B), but has no GDI activity (Figure 5A).

Recent cryo-electron microscopy of the yeast PIC of eIF2 within the PIC has provided significant insight into the interactions between eIF2 subunits, GTP, Met–tRNA, and other PIC proteins and 40S ribosomes. However, density associated with eIF5 was not defined (48). As shown in Figure 3C, the structure indicates that eIF2β–E189 sits at the surface of eIF2β with its side-chain projecting into the solvent and is close to residues that interact with Met–tRNA. In contrast eIF2β–S264, a residue implicated in controlling GTP-hydrolysis is within the CTD (pale blue in Figure 3C). The eIF2β-S264Y mutation (sui3-2) promotes premature hydrolysis of eIF2-bound GTP, permitting aberrant initiation at non-AUG codons (47). The EM structure shows that eIF2β-S264 binds both the tRNA, acceptor arm and the eIF2γ GTP-binding domain including the conserved switch 1 region that is important for controlling nucleotide binding (Sw1, orange in Figure 3C). S264 itself is juxtaposed to eIF2γ-bound GTP and is therefore ideally placed to limit nucleotide release from eIF2γ (48). There may be mechanistic similarities between eIF2β’s role in the PIC and its role in the eIF2αGDP/eIF5 complex. Within the PIC in WT cells eIF2β–S264 is critical to prevent premature hydrolysis of GTP and P1 release by eIF5 at non-AUG codons. Here we now show that in eIF2αGDP/eIF5 complexes, eIF2β is critical for eIF5 GDI to prevent premature eIF2β-independent release of GDP (47). Thus in both reactions eIF2β is important for ensuring tight regulation of eIF2 nucleotide status.

eIF5 is composed of two structured domains joined by an LR (62). There are separate structures for the amino terminal GAP domain and a CTD. The CTD is α-helical forming HEAT repeats that are shared with the GEF domain of eIF2Be (63,64) and region of the CTD is implicated in binding to the K-boxes of eIF2B (65) (46). The DWEAR motif implicated in eIF5 GDI lies just upstream of the crystallized CTD (63), so was placed in the LR (23). However, structure predictions indicate it is likely also α-helical and is probably an extended part of the CTD, similar to the longer α-helical CTD of related eIF5 mimic proteins (66). Combining the known structural and new functional information provides a framework for developing a model for eIF5 GDI function. This is shown in cartoon form in Figure 6, with elements of eIF2 depicted to mimic the structure view shown in Figure 3C. From the PIC structure, eIF2 has a core gamma subunit with eIF2α and β acting as ‘arms’ constrained by interactions with Met–tRNA, (Figure 6A). Upon GTP hydrolysis and release of Met–tRNA, eIF2GDP is released from the initiation complex and likely retains a similar overall form, but because the subunits are no longer constrained by Met–tRNA, interactions, the α and β arms may enjoy enhanced conformational flexibility. eIF2β movement relative to eIF2γ could permit the GDP release we observe in the absence of other ligands. This idea is shown in Figure 6B. We propose that eIF5 binding to eIF2β and γ acts as a ‘molecular clamp’ to stabilize an
Figure 6. Model for role of βE189 ensuring tight control of GDP release from eIF2. (A) eIF2 heterotrimer is shown in cartoon form based on the structure model shown in Figure 3C with eIF2y core bound to GTP and with α and β ‘arms’ ‘grasping’ Met–tRNA (purple). The structure of the N-terminal region of eIF2β bearing the lysine repeat ‘K-boxes’ is not known and is shown here as a white oval with black K-box stripes and is sequentially positioned over eIF2γ. βS264 is depicted as a yellow triangle adjacent to GTP bound to eIF2y G domain and Sw1 and BE189 is shown as a magenta triangle. (B) Following GTP hydrolysis and eIF2 release from the PIC and in the absence of other ligands, eIF2β ‘arm’ position is flexible and its movement contributes to the relatively high off-rate of GDP. (C) Binding of eIF5 GDI acts as a molecular clamp constraining eIF2β and stabilizing GDP binding to eIF2y. (D) E189K alters eIF5-eIF2 interactions so that GDP is no longer stabilized despite eIF5 interaction, shown by eIF2β ‘arm’ flexibility.

eIF5 conformation that prohibits GDP release (Figure 6C). Hence eIF5 GDI ensures tight eIF2β-dependent regulation of eIF2αGDP/GTP cycling. In contrast, when eIF5 binds to eIF2E189K, it cannot stabilize the conformation needed to prevent low-level GDP release (Figure 6D) and thereby enables faster growth to eIF2β mutants with low GEF activity (Figures 1 and 2) and prevents tight regulation of translation and GAAC by eIF2α (Figures 1D, 2 and 5C). To test this hypothetical model will likely require significant progress toward understanding structures and dynamics of the interactions between eIF5 and the eIF2 complex, which is beyond the scope of this study.

SUPPLEMENTARY DATA
Supplementary Data are available at NAR Online.

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REFERENCES
1. Hinnebusch, A.G. (2014) The scanning mechanism of eukaryotic translation initiation. Annu. Rev. Biochem., 83, 792–812.
2. Dever, T.E., Kinzy, T.G. and Pavitt, G.D. (2016) Mechanism and regulation of protein synthesis in Saccharomyces cerevisiae. Genetics, 203, 65–107.
3. Kapp, L.D. and Lorsch, J.R. (2004) GTP-dependent recognition of the methionine moiety on initiator tRNA by translation factor eIF2. J. Mol. Biol., 335, 923–936.
4. Algire, M.A., Maag, D. and Lorsch, J.R. (2005) Pi release from eIF2, not GTP hydrolysis, is the step controlled by start-site selection during eukaryotic translation initiation. Mol. Cell, 20, 251–262.
5. Pavitt, G.D. (2005) eIF2B, a mediator of general and gene-specific translational control. Biochem. Soc. Trans., 33, 1487–1492.
6. Wek, R.C., Jiang, H.Y. and Anthony, T.G. (2006) Coping with stress: eIF2 kinases and translational control. Biochem. Soc. Trans., 34, 7–11.
7. Costa-Mattioli, M., Gorbalenya, E., Stoffel, M., Harding, H., Herdy, B., Azzi, M., Brustle, O., Bidinosti, M., Ben Mamou, C., Marcinkiewicz, E., Yoshida, M. et al. (2005) Translational control of hippocampal synaptic plasticity and memory by the eIF2alpha kinase GCN2. Nature, 436, 1166–1173.
8. Pavitt, G.D. (2013) Less translational control, more memory. Elife, 2, e00895.
9. Sidrauski, C., Acosta-Alvear, D., Khoutorsky, A., Vedantham, P., Hearn, B.R., Li, H., Gamache, K., Gallagher, C.M., Ang, K.K., Wilson, C. et al. (2013) Pharmacological brake-release of mRNA translation enhances cognitive memory. Elife, 2, e00498.
10. Sidrauski, C., McGeachy, A.M., Ingolia, N.T. and Walter, P. (2015) The small molecule ISRIB reverses eIF2α phosphorylation on translation and stress granule assembly. Elife, 4, e05033.
11. Sekine, Y., Zvyazanova, A., Crespo-Casado, A., Fischer, P.M., Harding, H.P. and Ron, D. (2015) Stress responses. Mutations in a translation initiation factor identify the target of a memory-enhancing compound. Science, 348, 1027–1030.
12. Rowlands, A.G., Panners, R. and Henshaw, E.C. (1988) The catalytic mechanism of guanine nucleotide exchange factor action and competitive inhibition by phosphorylated eukaryotic initiation factor 2. J. Biol. Chem., 263, 5526–5533.
13. Pavitt, G.D., Ramaiya, K.V., Kimball, S.R. and Hinnebusch, A.G. (1998) eIF2 independently binds two distinct eIF2B subcomplexes that catalyze and regulate guanine-nucleotide exchange. Genes Dev., 12, 514–526.
14. Jennings, M.D., Zhou, Y., Mohammad-Qureshi, S.S., Bennett, D. and Pavitt, G.D. (2013) eIF2B promotes eIF5 dissociation from eIF2αGDP to facilitate guanine nucleotide exchange for translation initiation. Genes Dev., 27, 2696–2707.
15. Krishnamoorthy, T., Pavitt, G.D., Zhang, F., Dever, T.E. and Hinnebusch, A.G. (2001) Tight binding of the phosphorylated alpha subunit of initiation factor 2 (eIF2alpha) to the regulatory subunits of guanine nucleotide exchange factor eIF2B is required for inhibition of translation initiation. Mol. Cell. Biol., 21, 5018–5030.
16. Gomez, E., Mohammad, S.S. and Pavitt, G.D. (2002) Characterization of the minimal catalytic domain within eIF2B: the guanine-nucleotide exchange factor for translation initiation. EMBO J., 21, 5292–5301.
17. Hinnebusch, A.G. (2005) Translational regulation of GCN4 and the general amino acid control of yeast. Annu. Rev. Microbiol., 59, 407–450.
18. Jackson,R.J., Hellen,C.U. and Pestova,T.V. (2010) The mechanism of eukaryotic translation initiation and principles of its regulation. Nat. Rev. Mol. Cell Biol., 11, 113–127.
19. Pavitt,G.D. and Proud,C.G. (2009) Protein synthesis and its control in neuronal cells with a focus on vanishing white matter disease. Biochem. Soc. Trans., 37, 1289–1301.
20. Unbehaun,A., Borukhov,S.I., Hellen,C.U. and Pestova,T.V. (2004) Release of initiation factors from 43S complexes during ribosomal subunit joining and the link between establishment of codon-anticodon base-pairing and hydrolysis of eIF2-bound GTP. Genes Dev., 18, 3078–3093.
21. Singh,C.R., Lee,B., Udagawa,T., Yamamoto,Y., Pavitt,G.D. and Asano,K. (2006) An eIF5/eIF2 complex antagonizes guanine nucleotide exchange by eIF2B during translation initiation. EMBO J., 25, 4537–4546.
22. Singh,C.R., Udagawa,T., Lee,B., Wassink,S., He,H., Yamamoto,Y., Anderson,J.T., Pavitt,G.D. and Asano,K. (2007) Change in nutritional status modulates the abundance of critical pre-initiation intermediate complexes during translation initiation in vivo. J. Mol. Biol., 370, 315–330.
23. Jennings,M.D. and Pavitt,G.D. (2010) eIF5 has GDI activity necessary for translational control by eIF2 phosphorylation. Nature, 465, 378–381.
24. Jennings,M.D. and Pavitt,G.D. (2014) A new function and complexity for protein translation initiation factor eIF2B. Cell Cycle, 13, 2660–2669.
25. Amberg,D.C., Burke,D.J. and Strathern,J.N. (2005) Methods In Yeast Genetics. Cold Spring Harbor Laboratory Press, NY.
26. Asano,K., Krishnamoorthy,T., Phan,L., Pavitt,G.D. and Hinnebusch,A.G. (1999) Conserved bipartite motifs in yeast eIF2 and eIF2B, GTPase-activating and GDP-GTP exchange factors in translation initiation, mediate binding to their common substrate eIF2. EMBO J., 18, 1673–1688.
27. Homer,N., Merriman,B. and Nelson,S.F. (2009) BFAST: an alignment tool for large scale genome resequencing. PLoS One, 4, e7767.
28. Homer,N., Merriman,B. and Nelson,S.F. (2009) Local alignment of two-base encoded DNA sequence. BMC Bioinformatics, 10, 175.
29. Sassel,A. and Michael,T.P. (2010) Filtering error from SOLID Output. Bioinformatics, 26, 849–850.
30. Kent,W.J., Sugnet,C.W., Furey,T.S., Roskin,K.M., Pringle,T.H., Zahler,A.M. and Haussler,D. (2002) The human genome browser at UCSC. Genome Res., 12, 996–1006.
31. Li,H., Handsaker,B., Wysoker,A., Fennell,T., Ruan,J., Homer,N., Marth,G., Abecasis,G., Durbin,R. and Genome Project Data Processing, S. (2009) The sequence alignment/map format and SAMtools. Bioinformatics, 25, 2078–2079.
32. Homer,N. and Nelson,S.F. (2010) Improved variant discovery through local re-alignment of short-read next-generation sequencing data using SRMA. Genome Biol., 11, R70.
33. Li,H. (2011) A statistical method for SNP calling, mutation discovery, association mapping and population genetic parameter estimation from sequencing data. Bioinformatics, 27, 2987–2993.
34. Chenna,R., Sugawara,H., Koike,T., Lopez,R., Gibson,T.J., Higgins,D.G. and Thompson,J.D. (2003) Multiple sequence alignment with the Clustal series of programs. Nucleic Acids Res., 31, 3947–3950.
35. Richardson,J.P., Mohammad,S.S. and Pavitt,G.D. (2004) Mutations causing childhood ataxia with central nervous system hypomyelination reduce eukaryotic initiation factor 2B complex formation and activity. Mol. Cell. Biol., 24, 2352–2363.
36. Dever,T.E. (1997) Using GCN4 as a reporter of eIF2alpha phosphorylation and translational regulation in yeast. Methods, 11, 403–417.
37. Mohammad-Qureshi,S.S., Haddad,R., Hemingway,E.J., Richardson,J.P. and Pavitt,G.D. (2007) Critical contacts between the eukaryotic initiation factor 2B (eIF2B) catalytic domain and both eIF2beta and -2gamma mediate guanine nucleotide exchange. Mol. Cell. Biol., 27, 5225–5234.
38. Castelli,L.M., Talavera,D., Kershaw,C.J., Mohammad-Qureshi,S.S., Costello,J.L., Rowe,W., Sims,P.F., Grant,C.M., Hubbard,S.J., Ashe,M.P. et al. (2015) The 4E-BP Cdf20p mediates both eIF4E-dependent and independent repression of translation. PLoS Genet., 11, e1005233.
39. Mohammad-Qureshi,S.S., Haddad,R., Palmer,K.S., Richardson,J.P., Gomez,E. and Pavitt,G.D. (2007) Purification of FLAG-tagged eukaryotic initiation factor 2B complexes, subcomplexes, and fragments from Saccharomyces cerevisiae. Methods Enzymol., 431, 1–13.
40. Gordon,Y., Schmidt,C., Jennings,M.D., Matak-Vinkovic,D., Pavitt,G.D. and Robinson,C.V. (2014) eIF2B is a decameric guanine nucleotide exchange factor with a gamma2epsilon2 tetrameric core. Nat. Commun., 5, 3902.
41. Gomez,E. and Pavitt,G.D. (2000) Identification of domains and residues within the epsilon subunit of eukaryotic translation initiation factor 2B (eIF2B epsilon) required for guanine nucleotide exchange reveals a novel activation function promoted by eIF2B complex formation. Mol. Cell. Biol., 20, 3965–3976.
42. Holmes,L.E., Campbell,S.G., De Long,S.K., Sachs,A.B. and Ashe,M.P. (2004) Loss of translational control in yeast compromised for the major mRNA decay pathway. Mol. Cell. Biol., 24, 2998–3010.
43. Ashe,M.P., De Long,S.K. and Sachs,A.B. (2000) Glucose depletion rapidly inhibits translation initiation in yeast. Mol. Biol. Cell, 11, 633–648.
44. Horzinski,L., Hughe,A., Cardoso,M.C., Gontchier,C., Ouchchane,L., Schiffmann,R., Blanc,P., Boespflug-Tanguy,O. and Fogli,A. (2009) Eukaryotic initiation factor 2B (eIF2B) GEF activity as a diagnostic tool for eIF2B-related disorders. PLoS One, 4, e8318.
45. de Almeida,R.A., Fogli,A., Gaillard,M., Scheper,G.C., Boespflug-Tanguy,O. and Pavitt,G.D. (2013) A yeast purification system for human translation initiation factors eIF2 and eIF2Bpsilon and their use in the diagnosis of CACH/VWD disease. PLoS One, 8, e53958.
46. Luna,R.E., Arthani,H., Hiraishi,H., Nanda,J., Martin-Marcos,P., Markus,M.A., Akabayov,B., Milbradt,A.G., Luna,L.E., Sce,H.C. et al. (2012) The C-terminal domain of eukaryotic initiation factor 5 promotes start codon recognition by its dynamic interplay with eIF1 and eIF2beta. Cell Rep., 1, 689–702.
47. Huang,H.K., Yoon,H., Hannig,E.M. and Donahue,T.F. (1997) GTP hydrolysis controls stringent selection of the AUG start codon during translation initiation inSaccharomyces cerevisiae. Genes Dev., 11, 2396–2413.
48. Llacer,J.L., Hussain,T., Marler,L., Aitken,C.E., Thakur,A., Lorsch,J.R., Hinnebusch,A.G. and Ramakrishnan,V. (2015) Conformational differences between open and closed states of the eukaryotic translation initiation complex. Mol. Cell, 59, 399–412.
49. Erickson,F.L. and Hannig,E.M. (1996) Ligand interactions with eukaryotic translation initiation factor 2: role of the gamma-subunit. EMBO J., 15, 6311–6320.
50. von der Haar,T. and McCarthy,J.E. (2002) Intracellular translation initiation factor levels in Saccharomyces cerevisiae and their role in cap-complex function. Mol. Microbiol., 46, 531–544.
51. Ramirez,M., Wek,R.C., Vazquez de Aldana,C.R., Jackson,B.M., Freeman,B. and Hinnebusch,A.G. (1992) Mutations activating the yeast eIF-2 alpha kinase GCN2- isolation of alleles altering the domain related to histidyl- tRNA synthetases. Mol. Cell. Biol., 12, 5801–5815.
52. Donahue,T.F., Cigan,A.M., Pabich,E.K. and Valavicus,B.C. (1988) Mutations at a Zinc(H) finger motif in the yeast eIF-2 beta gene alter ribosomal start-site selection during the scanning process. Cell, 54, 895–906.
53. Cigan,A.M., Pabich,E.K., Feng,L. and Donahue,T.F. (1989) Yeast translation initiation suppressor sui2 encodes the alpha subunit of eukaryotic initiation factor 2 and shares sequence identity with the human alpha subunit. Proc. Natl. Acad. Sci. U.S.A., 86, 2784–2788.
54. Along,P.V., Cao,C. and Dever,T.E. (2008) Translation initiation factor 2gamma mutation alters start codon selection independent of Met-tRNA binding. Mol. Cell. Biol., 28, 6877–6888.
55. Borck,G., Shin,B.S., Stiller,B., Mimouni-Bloch,A., Thiele,H., Kim,J.R., Thakur,M., Skinner,C., Aschenbach,L., Smirin-Yosef,P. et al. (2012) eIF2gamma mutation that disrupts eIF2 complex integrity links intellectual disability to impaired translation initiation. Mol. Cell, 48, 641–646.
56. Das,S., Maiti,T., Das,K. and Maitra,U. (1997) Specific interaction of eukaryotic translation initiation factor 5 (eIF5) with the beta-subunit of eIF2. J. Biol. Chem., 272, 31712–31718.
57. Vazquez de Aldana,C.R., Dever,T.E. and Hinnebusch,A.G. (1993) The mechanism of eukaryotic translation initiation
factor 2 (eIF-2 alpha) that overcome the inhibitory effect of eIF-2 alpha phosphorylation on translation initiation. *Proc. Natl. Acad. Sci. U.S.A.*, 90, 7215–7219.

58. Dey, M., Trieselmann, B., Locke, E.G., Lu, J., Cao, C., Dar, A.C., Krishnamoorthy, T., Dong, J., Sicheri, F. and Dever, T.E. (2005) PKR and GCN2 kinases and guanine nucleotide exchange factor eukaryotic translation initiation factor 2B (eIF2B) recognize overlapping surfaces on eIF2alpha. *Mol. Cell. Biol.*, 25, 3063–3075.

59. Fekete, C.A., Mitchell, S.F., Cherkasova, V.A., Applefield, D., Algire, M.A., Maag, D., Saini, A.K., Lorsch, J.R. and Hinnebusch, A.G. (2007) N- and C-terminal residues of eIF1A have opposing effects on the fidelity of start codon selection. *EMBO J.*, 26, 1602–1614.

60. Chiu, W.L., Wagner, S., Herrmannova, A., Burela, L., Zhang, F., Saini, A.K., Valasek, L. and Hinnebusch, A.G. (2010) The C-terminal region of eukaryotic translation initiation factor 3a (eIF3a) promotes mRNA recruitment, scanning, and, together with eIF3j and the eIF3b RNA recognition motif, selection of AUG start codons. *Mol. Cell. Biol.*, 30, 4415–4434.

61. Singh, C.R., Curtis, C., Yamamoto, Y., Hall, N.S., Kruse, D.S., He, H., Hannig, E.M. and Asano, K. (2005) Eukaryotic translation initiation factor 5 is critical for integrity of the scanning preinitiation complex and accurate control of GCN4 translation. *Mol. Cell. Biol.*, 25, 5480–5491.

62. Jennings, M.D. and Pavitt, G.D. (2010) eIF5 is a dual function GAP and GDI for eukaryotic translational control. *Small GTPases*, 1, 118–123.

63. Wei, Z., Xue, Y., Xu, H. and Gong, W. (2006) Crystal structure of the C-terminal domain of S. cerevisiae eIF5. *J. Mol. Biol.*, 359, 1–9.

64. Boesen, T., Mohammad, S.S., Pavitt, G.D. and Andersen, G.R. (2004) Structure of the catalytic fragment of translation initiation factor 2B and identification of a critically important catalytic residue. *J. Biol. Chem.*, 279, 10584–10592.

65. Yamamoto, Y., Singh, C.R., Marintchev, A., Hall, N.S., Hannig, E.M., Wagner, G. and Asano, K. (2005) The eukaryotic initiation factor (eIF) 5 HEAT domain mediates multifactor assembly and scanning with distinct interfaces to eIF1, eIF2, eIF3, and eIF4G. *Proc. Natl. Acad. Sci. U.S.A.*, 102, 16164–16169.

66. Singh, C.R., Watanabe, R., Zhou, D., Jennings, M.D., Fukao, A., Lee, B., Ikeda, Y., Chiorini, J.A., Campbell, S.G., Ashe, M.P. et al. (2011) Mechanisms of translational regulation by a human eIF5-mimic protein. *Nucleic Acids Res.*, 39, 8314–8328.