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# Structural basis of eIF2B-catalyzed nucleotide exchange and phosphoregulation by the integrated stress response

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## Abstract

The integrated stress response (ISR) tunes the rate of protein synthesis. Control is exerted by phosphorylation of the general translation initiation factor eIF2. eIF2 is a GTPase, that becomes activated by eIF2B, a two-fold symmetric and heterodecameric complex that functions as eIF2's dedicated nucleotide exchange factor. Phosphorylation converts eIF2 from a substrate into an inhibitor of eIF2B. We report cryoEM structures of eIF2 bound to eIF2B in the dephosphorylated state. The structures reveal that the eIF2B decamer is a static platform upon which one or two flexible eIF2 trimers bind and align with eIF2B's bipartite catalytic centers to catalyze nucleotide exchange. Phosphorylation refolds eIF2a, allowing it to contact eIF2B at a different interface and, we surmise, thereby sequesters it into a non-productive complex.

# One Sentence Summary:

Supplementary Materials: Materials and Methods Figures S1-S7 Tables S1-S3 Movie S1 References (38-46)

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Competing interests: P.W. is an inventor on U.S. Patent 9708247 held by the Regents of the University of California that describes ISRIB and its analogs. Rights to the invention have been licensed by UCSF to Calico.

Accession numbers: Accession numbers for the human eIF2•eIF2B structures are as follows: EMD-0649, EMD-0651, EMD-0664 (density maps; Electron Microscopy Data Bank) and PDB-6081, PDB-6085, PDB-609Z (coordinates of atomic models; Protein Data Bank).

Data availability: All data needed to evaluate the conclusions in the paper are present in the paper and/or the supplementary materials, and the structural data is available in public databases. All of the raw cryo-EM data is also available upon request. The GP6452 yeast strain is available under a material transfer agreement with the University of Manchester.

Numerous factors regulate translation of the genetic code into proteins, including eukaryotic translation initiation factor 2 (eIF2), a GTPase composed of  $\alpha$ ,  $\beta$ , and  $\gamma$  subunits. During initiation, eIF2 binds tRNA<sup>Met</sup> and GTP to form a ternary complex that scans mRNAs for start codons. Following start codon detection, eIF2 $\gamma$  hydrolyzes its GTP and translation initiates. For eIF2 reactivation, GDP is replaced by GTP upon catalysis by a dedicated guanine nucleotide exchange factor (GEF), eIF2B.

eIF2 and eIF2B control translation initiation. Stress-responsive kinases phosphorylate eIF2a at conserved Ser51, transforming eIF2 from substrate into a competitive GEF inhibitor. Phosphoregulation of eIF2 is known as the integrated stress response (ISR)(1). Once activated, the ISR reduces overall protein synthesis, while enhancing translation of a small subset of mRNAs in response to cellular threats, including protein misfolding, infection, inflammation, and starvation(1–3).

eIF2B comprises two copies each of an  $\alpha$ ,  $\beta$ ,  $\gamma$ ,  $\delta$ , and  $\epsilon$  subunit that assemble into a twofold symmetric heterodecamer(4,5). The eIF2B $\epsilon$  subunit contains the enzyme's catalytic center and associates closely with eIF2B $\gamma$ . Two copies each of the eIF2B $\beta$  and  $\delta$  subunits form the complex's core, bridged by two eIF2B $\alpha$  subunits across the symmetry interface(4,6). Genetic and biochemical studies identified residues responsible for eIF2B's catalytic activity and suggested how eIF2 binding to eIF2B may differ following eIF2 $\alpha$ -S51 phosphorylation(4,7–10). Yet, how eIF2 recognizes eIF2B and how eIF2B catalyzes nucleotide exchange remained unknown, as did the transformation of eIF2 from a substrate to high-affinity inhibitor of eIF2B following its phosphorylation.

A potent small-molecule, drug-like inhibitor of the integrated stress response, ISRIB, allays the effects of eIF2a phosphorylation by activating eIF2B(11–13). Upon adding ISRIB, cells undergoing the ISR resume translation(12,13). When administered to rodents, ISRIB enhances cognition and ameliorates cognitive deficits caused by traumatic brain injury(14) and prion-induced neurodegeneration(15). Furthermore, eIF2B activation rescues cognitive and motor function in mouse models of leukoencephalopathy with vanishing white matter disease (VWMD), a fatal familial disorder associated with mutations spread over all eIF2B subunits(16).

ISRIB bridges the symmetric interface of two eIF2B subcomplexes to enhance the formation of the decameric eIF2B holoenzyme(17,18), enhancing available GEF activity by promoting higher-order assembly of the eIF2B decamer. However, it has remained enigmatic why decameric eIF2B would be more active than its unassembled subcomplexes. To explore this question, we determined structures of eIF2B bound with both its substrate, eIF2 $\alpha$ , $\beta$ , $\gamma$ , and inhibitor, eIF2 $\alpha$ •P.

We co-expressed all five subunits of human eIF2B in *E. coli* and all three subunits of human eIF2 in *S. cerevisiae* (Fig.S1A–B). The yeast expression strain lacked *GCN2*, the eIF2 kinase, to ensure expression of homogenously non-phosphorylated eIF2(19). We incubated

ISRIB and purified eIF2 at concentrations near the Michaelis constant of the nucleotide exchange reaction ( $K_m = \sim 1.5 \mu M$ ,(17)) and added an inter-amine crosslinker to stabilize complexes before sample vitrification and cryoEM analysis (Fig.S2A–C). We resolved two structures: eIF2B bound asymmetrically to a single eIF2 trimer and eIF2B bound symmetrically to two eIF2 trimers (Figs.1A–F,S3A,S4–5,TableS1–S3).

Snaking across the surface of eIF2B, we observed density consistent in size and shape with eIF2 subunits and the previously unresolved eIF2Be HEAT domain. Comparison with homologous structures of eIF2a and eIF2 $\gamma$  revealed that the assembled eIF2•eIF2B complex retained similarity to the structures of these individually analyzed domains (20, 21)(Fig. 2,Fig.S6). We only resolved a single helix of eIF2 $\beta$  (Fig. 1A,D,Fig. 2A), consistent with other studies(20,21). In both reconstructions, all five subunits of eIF2B can be superimposed on previously determined structures lacking eIF2 (RMSD of ~0.6Å)(17). Thus, eIF2B retained its overall arrangement when bound to one or two eIF2s (Fig. 1A–F), indicating that eIF2 binds via equivalent modes to both sides of a static eIF2B scaffold with no allostery in eIF2B upon eIF2 engagement. This is consistent with non-cooperative kinetics reported for nucleotide exchange by eIF2B decamers(17).

Bound to eIF2B, eIF2 adopted an extended 150Å conformation (Figs.1–2) with eIF2's central nucleotide-binding  $\gamma$ -subunit flanked by its  $\alpha$ - and  $\beta$ -subunits at its opposing ends. eIF2 $\gamma$  contains classical GTP-binding motifs, including the nucleobase-binding G4 motif, the phosphate-binding P-loop, and switch helices 1 and 2. eIF2B recognizes eIF2 via coincident binding of both eIF2 $\alpha$  and eIF2 $\gamma$ . Binding to both eIF2 subunits involves bipartite elements of eIF2B(Figs. 1,2A–C).

First, bipartite recognition of eIF2 $\gamma$  involves two domains of eIF2Be that function together to splay open the nucleotide-binding site. Our nucleotide-free cryoEM model is similar to the  $\gamma$ -subunit of GTP-bound aIF2 from *S.solfataricus*(24) (Fig.2D–E, average RMSD ~2.3Å). Yet, surrounding the GTP binding pocket, the structures diverged considerably with the P-loop in eIF2B•eIF2 partially occluding the nucleotide-binding site (RMSD of ~12Å). Prior work implicated the HEAT domain in catalysis(23,24). In agreement, eIF2 $\gamma$  interacts with the HEAT domain, including a partially hydrophobic surface that includes eIF2Be Y583(Fig. 2C). On the opposing side of the nucleotide-binding pocket, the central core of eIF2Be engaged with an open loop conformation of Switch 1. This change appears due to electrostatic interactions between eIF2 $\gamma$  R75 in Switch 1 and Q258 and D262 in eIF2Be. Thus, both eIF2Be's HEAT domain and core collaborate to open the nucleotide-binding site (Fig. 2B–D).

The second example of bipartite recognition concerns eIF2a binding in the cleft between eIF2B $\beta$  and eIF2B $\delta$ ' ( $\delta$ ' indicates the  $\delta$ -subunit from the opposing tetramer, Figs.1–3). Notably, this binding site only exists when two tetramers of eIF2B( $\beta\gamma\delta\epsilon$ ) associate to form the symmetry interface in octameric eIF2B( $\beta\gamma\delta\epsilon$ )<sub>2</sub>. eIF2a contains two structured domains separated by a flexible linker(Fig. 1–2,S6). The N-terminus consists of an OB-fold, common in tRNA-binding proteins(20). The OB-fold is further elaborated with a positively-charged loop (the S-loop), while the C-terminal  $\alpha\beta$ -fold connects eIF2a to eIF2 $\gamma$ . The S-loop harbors S51 and is responsible for all of the resolvable contacts between eIF2a and eIF2B's

β subunit (Fig. 3A). Prior work implicated a conserved 'K<sup>79</sup>GYID<sup>83</sup>' motif in eIF2a as being important for eIF2B binding(10). Of note, an interaction between Y81 was wellresolved adjacent to the equally prominent R250 on eIF2Bδ' (Fig. 3B). When we mutated R250 to either alanine or glutamate, neither mutation affected the residual GEF activity displayed by dissociated tetramers (Fig. 3D;R250A k<sub>obs</sub>=0.013min<sup>-1</sup>,R250E k<sub>obs</sub>=0.023min <sup>-1</sup>,wild-type k<sub>obs</sub>=0.016min<sup>-1</sup>), while both mutants diminished the GEF activity of the ISRIB-stabilized eIF2B octamer when compared to wild-type (Fig. 3E;R250A k<sub>obs</sub>=0.012min<sup>-1</sup>,R250E k<sub>obs</sub>=0.017min<sup>-1</sup>,wild-type k<sub>obs</sub>=0.063min<sup>-1</sup>). This is consistent with the notion that unphosphorylated eIF2a interacts with the *trans*-tetramer only upon assembly of octameric or decameric eIF2B across its symmetry interface.

On the *cis*-tetramer, eIF2a's positively-charged S-loop binds negatively charged and polar residues along the exposed surface of eIF2B $\beta$ . This binding site is consistent with yeast studies suggesting that mutations in this site compromise eIF2 binding(7). Examination of the structure identified a potential hydrogen bond between eIF2B $\beta$  N132 and eIF2a R52 (Fig. 3C). We substituted N132 with aspartate, anticipating that the introduced charge complementarity would enhance binding to eIF2a R52. When compared to wild-type eIF2B tetramers, eIF2B- $\beta$ N132D tetramers and ISRIB-stabilized octamers indeed proved to be gain-of-function mutations, exhibiting ~two-fold enhanced GEF activity (Figs.3F–G,S1D, eIF2B( $\beta\gamma\delta\epsilon$ )  $\beta$ N132D k<sub>obs</sub>=0.044 min<sup>-1</sup>, eIF2B( $\beta\gamma\delta\epsilon$ )<sub>2</sub>  $\beta$ N132D k<sub>obs</sub>=0.169 min<sup>-1</sup>). This is consistent with eIF2B tetramers possessing reduced activity when compared to assembled octameric or decameric holo-eIF2B. eIF2a binding in the cleft between tetramers further supports the notion that ISRIB enhances eIF2B's GEF activity by promoting higher-order assembly.

To understand how eIF2a phosphorylation on S51 transforms eIF2 from substrate to inhibitor, we co-expressed the isolated eIF2a subunit in *E.coli* with the kinase domain of PERK (Fig.S1C). We incubated pre-assembled eIF2B decamers with an excess of eIF2a-P, followed by crosslinking and vitrification. Reconstruction of the eIF2B decamer adorned with a two copies of eIF2a-P (Figs. 4A,S7–S8,Tables S1–S3), revealed eIF2a-P bridging the interface between eIF2Bδ and eIF2Ba(Fig. 4A). Intriguingly, we observed no overlap between the binding sites of non-phosphorylated eIF2a described above and eIF2a-P (Fig. 4B–C).

Density for both eIF2a S51-P and two arginines positioned ~4Å away, eIF2a R53 and R63, were well-resolved and suggestive of an electrostatic coordination responsible for phosphorylation-induced refolding of the S-loop (MovieS1,Fig. 4F–G,S8), as initially observed by Kashiwagi, et al (this issue). The phosphorylation-induced rearrangement also positions hydrophobic residues on eIF2a for potential interactions with hydrophobic residues on eIF2B (including eIF2a I55, I58, and L61 and eIF2B8 L314, A315, A318, and F322).

This structural model agrees with analyses in yeast and mammalian systems. First, eIF2Ba is dispensable for viability in yeast, yet eIF2Ba deletion impairs phospho-inhibition of eIF2B, consistent with the subunit's role in binding eIF2a-P(27). Point mutations with identical phenotypes cluster at the interface between eIF2Ba and eIF2B\delta, e.g., eIF2Ba

F239 and eIF2B& M506 and P508(28, 29). Importantly, eIF2B& L314 complements the hydrophobic surface of the eIF2a S-loop that is exposed upon refolding, and mutation of the equivalent position in *S. cerevisiae*, L381Q, impairs the ISR in yeast(28). These data validate the phosphorylation-induced refolding and relocation of eIF2a-P observed here.

Our analyses reveal the mechanistic basis of eIF2B's nucleotide exchange activity and suggest how phosphorylation converts eIF2 from substrate to inhibitor. The non-phosphorylated form of eIF2 binds to a composite surface created only in the assembled decamer, allowing both the core and the flexibly attached HEAT domain of eIF2Be to engage its target in concert for enhanced GEF activity.

By contrast, eIF2a-P adopts a new conformation and suggests how the S-loop may become incompatible for binding to the site where nonphosphorylated eIF2a binds as a substrate (Movie S1). Phosphorylation thus enables a distinct binding mode on the opposite side of eIF2B where eIF2a-P lies exiled at the interface of eIF2Ba and eIF2B8. In eIF2•P, the rearrangement of the S-loop derives from an intramolecular electrostatic interaction between R63 and R53 and the phosphate, which also exposes a hydrophobic surface upon phosphorylation-induced refolding. We surmise that this new binding mode is nonproductive for nucleotide exchange on eIF2-P and sequesters the catalytic domains into an inhibited state that prevents the catalytic moieties of eIF2Be from properly engaging in productive nucleotide exchange.

## Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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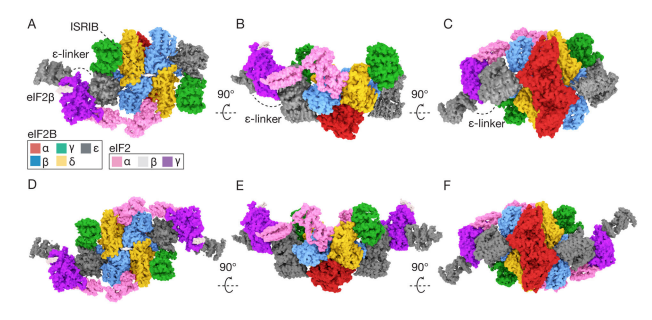
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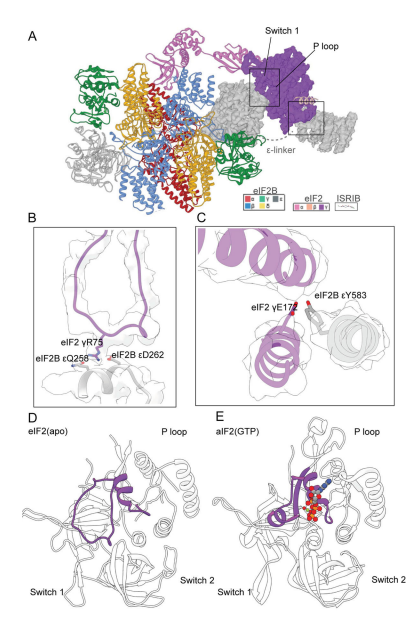
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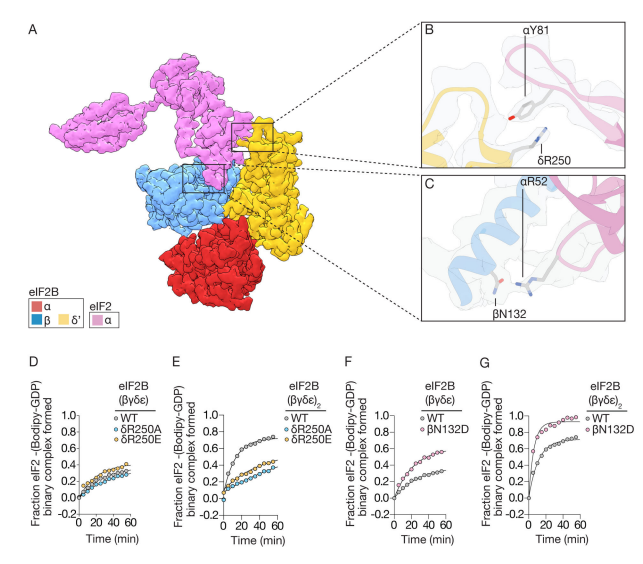
**Fig. 1. eIF2B heterodecamer bound to one or two eIF2 heterotrimers.** (A-C) Orthogonal views of a single versus (D-E) a pair of elongated eIF2 heterotrimers bound to ISRIB-stabilized eIF2B decamers. ISRIB density is rendered in white.



## Fig. 2. The bipartite basis of guanine nucleotide exchange by eIF2B.

(A) Structural model of a single eIF2 heterotrimer bound to eIF2B decamer, emphasizing the cryoEM density for eIF2 $\gamma$  and its interactions with eIF2B $\epsilon$ . (B) Comparison of an aIF2 structure bound to GTP (PDB: 4RCY) and (C) GDP (PDB: 4RD6) with the open, nucleotide-free state of eIF2 (D) reported here.

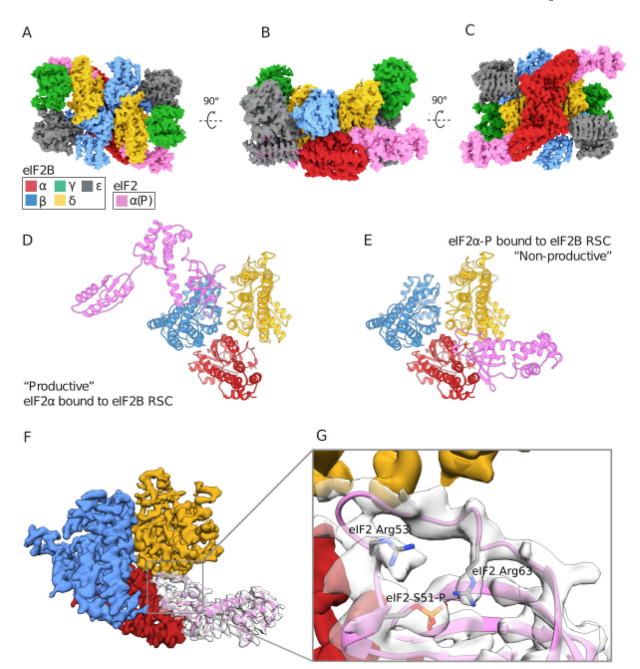
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### Fig. 3. The bipartite basis of eIF2a recognition and assembly-stimulated activity.

(A) cryoEM density for eIF2a bound to the regulatory subcomplex  $(\alpha,\beta,\delta \text{ or RSC})$  of eIF2B. (B) Density and zoom-in detail of a cation- $\pi$  interaction between eIF2B $\delta$  and eIF2a. (C) polar interactions between eIF2B $\beta$  and the S-loop of eIF2a. (D,F) GEF activity of wildtype versus mutated eIF2B $(\beta\gamma\delta\epsilon)$  tetramers, and (E,G) ISRIB-stabilized eIF2B $(\beta\gamma\delta\epsilon)_2$  octamers measured by loading of fluorescent GDP onto eIF2.

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### Fig. 4. The structural basis of phosphoregulation by the ISR.

(A-C) Orthogonal views of a pair of S51-phosphorylated eIF2a subunits bound to the eIF2B decamer. (D) Comparison of the productive binding mode of non-phosphorylated eIF2a, versus (E) the non-productive and non-overlapping binding mode of phosphorylated eIF2a. (F-G) CryoEM density and interpretation of the phosphorylated eIF2a binding mode and refolded conformation of the S-loop, placing the S51 phosphate moiety near eIF2 a R53 and R63.