Mutations Linked to Leukoencephalopathy with Vanishing White Matter Impair the Function of the Eukaryotic Initiation Factor 2B Complex in Diverse Ways

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Leukoencephalopathy with vanishing white matter (VWM) is a severe inherited human neurodegenerative disorder that is caused by mutations in the genes for the subunits of eukaryotic initiation factor 2B (eIF2B), a heteropentameric guanine nucleotide exchange factor that regulates both global and mRNA-specific translation. Marked variability is evident in the clinical severity and time course of VWM in patients. Here we have studied the effects of VWM mutations on the function of human eIF2B. All the mutations tested cause partial loss of activity. Frameshift mutations in genes for eIF2Bε or eIF2Bβ lead to truncated polypeptides that fail to form complexes with the other subunits and are effectively null mutations. Certain point mutations also impair the ability of eIF2Bβ or eIF2Bε to form eIF2B holocomplexes and also diminish the intrinsic nucleotide exchange activity of eIF2B. A point mutation in the catalytic domain of eIF2Bε impairs its ability to bind the substrate, while two mutations in eIF2Bβ actually enhance eIF2 binding. We provide evidence that expression of VWM mutant eIF2B may enhance the translation of specific mRNAs. The variability of the clinical phenotype in VWM may reflect the multiple ways in which VWM mutations affect eIF2B function.

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Leukoencephalopathy with vanishing white matter (VWM) is a severe inherited human neurodegenerative disease (35). This condition is also known as childhood ataxia with diffuse central nervous system hypomyelination (34). Clinically, this condition is characterized by progressive ataxia, spasticity, and variable optic atrophy, as well as seizures. Mental decline, if seen, is usually mild. More rapid deterioration is associated with minor head injury or fever, and the latter often results in coma. Death occurs after a variable time period, from several years up to decades. In magnetic resonance imaging, a diagnostic pattern with diffuse abnormality of cerebral white matter is observed (35, 36). Over time, the abnormal white matter vanishes to be replaced by cerebrospinal fluid or tissue water. There is considerable phenotypic variation. For example, the so-called “Cree” leukoencephalopathy seen in North American Indians is especially severe, leading to very early onset of neurological deterioration (3 to 9 months) and death before 21 months (8). Recent work has revealed additional serious forms of VWM that involve very early onset and early death (7, 8). In some patients, the ovaries are also affected (35).

VWM shows autosomal recessive inheritance. Linkage analysis initially assigned a VWM locus to a 5-centimorgan interval on chromosome 3q27 (21). Subsequent work showed that that mutations in the gene for the ε subunit of the translation factor eukaryotic initiation factor 2B (eIF2Bε) were associated with VWM. These included several point or frameshift mutations within this gene. In some patients, linkage analysis did not point to a role for the eIF2Bε (or EIF2B5) gene. In several of them, VWM was found to be associated with mutations in the β subunit of eIF2B, a heteropentamer (22). Subsequent work showed that mutations in any eIF2B gene can cause VWM (37). All VWM patients have now been found to have mutations in one or other of the genes for the subunits of eIF2B. Affected individuals carry two mutated copies of a given eIF2B gene; i.e., inheritance shows an autosomal recessive pattern. The particularly severe Cree form of this disease is linked to a point mutation (R195H) in eIF2Bε (8), and a second severe form is linked to a different point mutation in the same gene (7).

eIF2B is a guanine nucleotide exchange factor that plays a key role in the initiation of mRNA translation and in its control (28). It promotes GDP/GTP exchange on initiation factor eIF2. In its GTP-bound state, eIF2 recruits the initiator Met-tRNA to the 40S subunit during translation initiation. Since it is the anticodon of this tRNA that recognizes the start codon in the mRNA, active eIF2 is required for all normal translation initiation events. eIF2B activity can be regulated in a number of ways. The best documented of these is the phosphorylation of the α subunit of eIF2 at Ser51 (4), which converts it from a substrate of eIF2B into a potent competitive inhibitor (30). Phosphorylation of eIF2α is increased under a wide range of cellular stress conditions and can be catalyzed by several different protein kinases (3). Since active eIF2.GTP is required for all translation initiation events, inhibition of eIF2B activity causes a decrease in general protein synthesis. However, reduced eIF2B activity actually enhances the translation of certain mRNAs by virtue of the presence of upstream open reading frames (uORFs) in their 5’ untranslated regions (UTRs). Examples include the mRNAs for the transcription regulators GCN4 in (16) and ATF4 in mammals (13). Since these pro-
teins modulate transcription of multiple genes, decreased eIF2B activity may also affect the expression of many genes. eIF2B is composed of five nonidentical subunits, α through ε (16). Of these, eIF2Be displays catalytic activity when expressed alone, and the minimal catalytic region of the yeast ortholog was recently delineated (9). The corresponding region of the mammalian protein contains the site that is phosphorylated by glycogen synthase kinase 3 and that inactivates eIF2B (39). The other subunits do not possess a region homologous to this catalytic segment and appear not to be involved in catalysis. However, eIF2Bε shows similarity to the N-termina1 part of eIF2Be and forms a binary complex with eIF2Be in yeast, which is referred to as the catalytic subcomplex (25). The other three subunits, α, β, and δ, also show mutual sequence similarity and form a trimeric subcomplex in yeast. Evidence from careful genetic studies with yeast indicates that these subunits play a key role in sensitizing the eIF2B holocomplex to inhibition by eIF2[α]P (20, 24, 40). Indeed, recombinant eIF2α can bind to the εβ subcomplex, which has been termed the regulatory subcomplex (25). Although eIF2Be itself can mediate guanine nucleotide exchange, its activity is greatly enhanced by its association with the other subunits (6, 10).

VWM is the first example of an inherited human disease caused by mutations in a component of the basal translational machinery. A clear priority is now to establish how mutations in eIF2B lead to VWM and the other lesions associated with this condition. It will also be important to understand why they give rise primarily to a neurological disease and why its severity and age of onset are so variable. Here, as an essential first step, we have studied how they affect the integrity and function of the eIF2B complex. Our data show that all the VWM mutations tested cause partial loss of activity of human eIF2B. However, the magnitude of the effect differs between mutations. Most interestingly, the basis of the decreased activity differs between mutations, prompting the possibility that the variable phenotype of the condition may reflect the distinct ways in which different mutations affect the eIF2B complex.

MATERIALS AND METHODS

Materials. Chemicals and biochemicals were obtained as described previously (40). Antibodies to eIF2B subunits were obtained from Santa Cruz, apart from the anti-eIF2Be antibody which was described in reference 38. Anti-myc was from Sigma. Antihemagglutinin (anti-HA) was from Boehringer Mannheim, and anti-eIF2Bε has been described previously (23). Antibodies to phosphorylated eIF2Bε were from Cell Signaling Technology, and anti-eIF2ε was a monoclonal kindly provided by the late E. C. Henshaw.

Construction of expression vectors. Full-length eIF2B cDNAs were obtained from the IMAGE consortium or were cloned into an expression vector with N-terminal hexahistidine (His6) and myc tags. As the C terminus of eIF2B cDNA were cloned into an expression vector with N-terminal hexahistidine further information, see Table 2). These mutant cDNAs or the wild-type (WT) (RT-PCR) (Table 1). VWM mutations were made by PCR-based technology (for kindy provided by the late E. C. Henshaw.

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TABLE 1. Sources of cDNAs and WT human eIF2B constructs

<table>
<thead>
<tr>
<th>Gene</th>
<th>Source(s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>eIF2B1</td>
<td>IMAGE clone 3534548</td>
</tr>
<tr>
<td>eIF2B3</td>
<td>IMAGE clone 455592, RT-PCR for C terminus</td>
</tr>
<tr>
<td>eIF2B4</td>
<td>IMAGE clone 3534267, RT-PCR for C terminus</td>
</tr>
</tbody>
</table>

a IMAGE clones were bought from MRC gene service (London, United Kingdom) or Invitrogen (Paisley, United Kingdom). RT-PCR was carried out with human HEK 293 cells RNA to generate full-length cDNA for eIF2B1, eIF2B3, and eIF2B4 cDNAs. All products were confirmed by DNA sequencing. WT eIF2B and eIF2B5 were cloned into pHis with His and myc tags, at the N terminus, myc-tag-only constructs of five individual subunits were made by cloning into pCMV-tag3 (Stratagene).

Cell culture, transfection, treatment, and lysis. Human embryonic kidney 293 (HEK 293) cells were grown and lysed exactly as described earlier (40). Cells were transfected by using the calcium phosphate precipitation method as described in reference 11. Unless otherwise stated, cells were lysed about 48 h after transfection. For studies on the ATF-linked report, HEK 293 cells (in 10-cm-diameter dishes) were transfected with pEGFP or pATF4-EGFP plus the construct under study (or empty vector as a control) as indicated. Twenty-four hours after transfection, the media were changed to Dulbecco’s modified Eagle’s medium without methionine. Four-and-a-half microliters of [35S]methionine was then added. Sixteen hours later, cells were lysed and lysates were used for immunoprecipitation (IP) with anti-GFP antibody (Roche). One quarter of the sample was loaded in each lane. For Northern blot analyses, 48 h after transfection, one dish of cells for each combination of vectors was collected for RNA isolation with Trizion (Invitrogen). One quarter of the total RNA isolated was loaded per lane in a 1.4% formaldehyde-agarose gel. Northern blot hybridization with probes for glyceraldehyde-3-phosphate dehydrogenase (GAPDH) and GAPDH were done as described by Sambrook et al. (31). Films were developed after 6 h (GFP) or 2 days (GAPDH) exposure at −80°C.

For thermal treatment of cells, dishes were transferred from the incubator (37°C) to a water bath at the desired temperature (usually 41°C) for 2 h. A water bath was used to ensure that cells quickly reached the new temperature.

RESULTS

Choice of VWM mutants for investigation in this study. Our aim was to gain insight into the effects of VWM mutations on the function of human eIF2B complexes. We therefore overexpressed WT or mutant human eIF2B subunits in HEK 293 cells, as His6- and/or myc-tagged proteins. We chose HEK 293 cells because they are highly transfectable human cells and because it has already been shown that eIF2Be expressed in them is incorporated into functional eIF2B complexes (39). Many VWM mutations occur in eIF2Be, and we chose were purified on Ni-nitrolotriacetic acid (NTA)-agarose (Qiagen). Typically 300 μg of protein was used for this type of analysis. The bound material was analyzed by SDS-PAGE–Western blotting as described above. To measure eIF2B activity, complexes were isolated in the same way and GDP/GTP exchange activity was determined by using our standard assay with complexes containing purified human eIF2 and [3H]GDP as substrate. Since the levels of expression of WT or mutant eIF2Bε could vary significantly within a given experiment, we first assessed the amount of the purified complexes by SDS-PAGE–immunoblotting for myc and then adjusted the amounts of material for the assay accordingly to ensure that similar amounts of recombinant eIF2B were used in all assays.

For analysis of eIF2B expression, analysis of eIF2B complexes and activity.

For thermal treatment of cells, dishes were transferred from the incubator (37°C) to a water bath at the desired temperature (usually 41°C) for 2 h.
TABLE 3. Mutations of eIF2B studied in this report*

<table>
<thead>
<tr>
<th>Subunit</th>
<th>DNA mutation</th>
<th>Effect on protein sequence</th>
<th>Comment on mutation</th>
<th>Other remark(s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>eIF2B</td>
<td>792delTinsACA</td>
<td>F264fs</td>
<td>Compound heterozygous</td>
<td>Gives truncated protein</td>
</tr>
<tr>
<td>eIF2B</td>
<td>A271G</td>
<td>T91A</td>
<td>Homozygous</td>
<td>T conserved in mammals</td>
</tr>
<tr>
<td>eIF2B</td>
<td>G338A</td>
<td>R113H</td>
<td>Homozygous</td>
<td>R in rabbit; H in rodents; varies in rest</td>
</tr>
<tr>
<td>eIF2B</td>
<td>G984A</td>
<td>R195H</td>
<td>Homozygous</td>
<td>R in mammals</td>
</tr>
<tr>
<td>eIF2B</td>
<td>G943G</td>
<td>R315G</td>
<td>Homozygous</td>
<td>Varies</td>
</tr>
<tr>
<td>eIF2B</td>
<td>G1016C</td>
<td>R339P</td>
<td>Compound heterozygous with T91A</td>
<td>R in mammals</td>
</tr>
<tr>
<td>eIF2B</td>
<td>T1882C</td>
<td>M203fs</td>
<td>Compound heterozygous with T91A</td>
<td>W in mammals; W, Y, or L in rest; Gives truncated protein</td>
</tr>
<tr>
<td>eIF2B</td>
<td>607 612delinsTG</td>
<td>E213G</td>
<td>Homozygous</td>
<td>E in all spp. except <em>A. thaliana</em></td>
</tr>
<tr>
<td>eIF2B</td>
<td>A638G</td>
<td>V316D</td>
<td>Compound heterozygous with E213G</td>
<td>V in all known sequences</td>
</tr>
</tbody>
</table>

*Conservation refers to the known sequences: mammalian sequences are from rat, rabbit, and mouse; others are from *Caenorhabditis elegans*, *A. thaliana*, *D. melanogaster*, and *S. cerevisiae.*
eIF2Bε (Fig. 1B). Without eIF2Bγ, eIF2Bε was unable to bind even the combination of eIF2Bα, -β, and -δ (Fig. 1B). Indeed, the only four-subunit combination that forms stable complexes is that of β, γ, δ, and ε (Fig. 1B), consistent with the findings that eIF2B lacking eIF2Bε can be generated in insect cells (6) or isolated from mammalian cells (5).

His-myc-eIF2Bβ was unable to associate with any of the other individual subunits (Fig. 1A). We did not detect an eIF2Bαβεδ complex, suggesting that association with eIF2Bε may be needed for stable interaction between the mammalian α, β, and δ subunits (in contrast with the situation for yeast eIF2B (reference 40 and Fig. 1C). These data are the first detailed information on the assembly of mammalian eIF2B. Figure 1D is a summary cartoon showing which complexes can form.

Certain VWM mutations affect the ability of eIF2B subunits to form holocomplexes. To study whether VWM mutations affected assembly of eIF2B complexes, HEK 293 cells were transfected with vectors for VWM/WT eIF2B subunits, either with the cDNA for a single subunit (Fig. 2) or with the cDNAs for all five subunits (Fig. 3). The missense mutants expressed as polypeptides of sizes similar to that of the WT protein (Fig. 2A). As expected, the constructs mimicking the frameshift mutants gave small polypeptides (Fig. 2B).

Initially, we expressed the VWM mutant His-myc-tagged eIF2B subunits alone and used Ni-NTA–agarose to isolate them and study their ability to interact with the other, endogenous subunits. A limitation here is that we are unable to detect eIF2Bε by using the available antisera. However, as shown above, eIF2Bε is not required for complex formation between the other four subunits. The β, γ, and δ subunits were clearly detected in pull-downs from cells expressing WT His-myc-eIF2Bε and for each of the five point mutants of eIF2Bε tested (Fig. 2C). In contrast, the other subunits were not ob-
served in pull-downs from cells expressing the eIF2Bε frame-shift mutant, presumably because the truncated polypeptide lacks regions required for interaction with them. It also lacks the catalytic domain (9) and is thus functionally null. Lastly, the pull-downs for the T91A and H9004ε/CAT mutants consistently showed less of the other four subunits than were seen in pull-downs from WT eIF2Bε, suggesting that they have reduced ability to form eIF2B holocomplexes (Fig. 2C).

eIF2Bβ(E213G) formed complexes with the other eIF2B subunits to almost the same extent as WT eIF2Bβ (Fig. 2D). In contrast, only very weak signals were observed for the other subunits following affinity isolation of eIF2Bβ(V316D) on NTA-agarose (Fig. 2D), indicating that it cannot efficiently form holocomplexes. The polypeptide that mimics the frameshift M203fs failed to interact with the other subunits, suggesting that the C terminus of eIF2Bβ is needed for this. Thus, certain VWM mutations affect eIF2B function by impairing holocomplex formation. As complex formation greatly enhances eIF2B activity (10), this is likely to reduce markedly eIF2B activity in patients’ cells.

To assess what proportion of total eIF2B complexes in the transiently transfected cells contain the recombinant subunit, we immunoprecipitated complexes with anti-eIF2Bε and analyzed the precipitates through SDS-PAGE–immunoblotting by using anti-eIF2Bε or -H9252. As seen in Fig. 2E, >50% and almost 80% of the complexes contain the recombinant eIF2Bε and -β proteins, respectively.

The potencies of antisera for eIF2Bβ, -γ, -δ, and -ε differ considerably, and there is no suitable antibody for eIF2Bα. We therefore also used an additional approach in which we coexpressed the eIF2B subunit under study with the other four polypeptides, each with a myc tag. All five subunits were fairly evenly expressed as assessed by immunoblotting of lysates with anti-myc (Fig. 3A). However, eIF2Bε(ΔCAT) and the truncation mutants of eIF2Bε and -β were expressed at lower levels, perhaps because they are less able to form complexes. When His-myc-tagged eIF2Bε missense mutant polypeptides were isolated on Ni-NTA–agarose, the other four myc-tagged subunits were readily detected in every case (Fig. 3B, top), showing that eIF2Bε is incorporated into these complexes.

FIG. 2. Formation of complexes containing a VWM mutant version of eIF2Bβ or ε. HEK 293 cells were transfected with vectors for WT or selected VWM mutants of the β or ε subunits of eIF2B, as fusions with His and myc tags, or with the empty vector (Vec). (A and B) Samples of cell lysate were analyzed by SDS-PAGE and Western blotting by using anti-myc. (C and D) Samples of cell lysate were subjected to chromatography on Ni-NTA–agarose prior to immunoblot analysis of the bound material with anti-myc. -eIF2Bβ, and -eIF2Bε as well as anti-eIF2Bβε (C) or -eIF2Bε (D). Δε and Δβ in panels B and D indicate the frameshift mutants, the positions of the truncated polypeptides that they encode being indicated by diagonal arrows. Positions of the other, endogenous eIF2B subunits are also shown. Weak signals for the anti-eIF2Bβ and ε antisera in panels C and D, respectively, mean that their signals are shown for longer exposures as insert panels. The asterisk indicates a nonspecific detection by anti-myc of a protein that binds nonspecifically to Ni-NTA–agarose. (E) Cells were transfected with vectors for myc/His-eIF2Bε or ε. Lysates were subjected to IP with anti-eIF2Bε, and precipitates were analyzed by SDS-PAGE and immunoblotting with anti-eIF2Bε (left side) or anti-eIF2Bβε (right side).
FIG. 3. Formation of complexes containing VWM variants of eIF2Bβ or ε. HEK 293 cells were transfected with vectors for WT or selected VWM mutants of the β or ε subunits of eIF2β, or the ΔCAT mutant of eIF2βε, as fusions with His and myc tags, and with vectors encoding myc-tagged version of the other four subunits. As a negative control, cells were transfected with empty vectors (indicated). Samples of cell lysate were analyzed directly by SDS-PAGE and Western blotting (A) or first subjected to chromatography on Ni-NTA–agarose prior to analysis by SDS-PAGE–immunoblotting (B). Blots were developed with anti-myc (A and B) and anti-eIF2α (B). Positions of the myc-tagged eIF2B subunits and of eIF2α are shown. Δε and Δβ indicate the positions of the truncated polypeptides arising from the frameshift mutations. In the lower part of panel B, only the vectors for WT or mutant eIF2βε were used (or empty vector as negative control). Samples of cell lysate were subjected to chromatography on Ni-NTA–agarose prior to analysis by SDS-PAGE and Western blotting with anti-eIF2(Ser51[β]). Equal expression of eIF2B subunits was confirmed by probing with anti-myc (data not shown). Where indicated, cells were pretreated with sodium arsenite (0.5 mM, 30 min) prior to lysis. Cells were also transfected with the empty vector, and similar pull-downs were performed to confirm that very little, if any, eIF2 was retained nonspecifically on the resin (B, upper and lower sections). (C and D) As done for panel B, vectors for the indicated His-myc-tagged eIF2B subunits were used together with myc-tagged versions of the other four WT subunits. Where indicated, cells also received DNA for the myc-tagged versions of eIF2Bε or β. Data are immunoblots of material retained on Ni-NTA–agarose (except where indicated, “lysate”). In panel D, the two arrows for eIF2βε indicate that the broad band seen for this subunit actually consists of a poorly resolved doublet of species, the lower one being the myc-tagged eIF2βε and the upper one being the myc/His-tagged subunit. (E) HEK 293 cells were transfected with HA- and myc-tagged versions of eIF2βε; those shown in lane 2 also received the vector for myc-eIF2βγ. Samples of lysates were subjected to IP with anti-HA followed by SDS-PAGE and immunoblotting, development being with anti-HA (upper section) or anti-myc (lower section).
Similar data were obtained for WT and VWM mutant eIF2Bβ (Fig. 3B). The data for the eIF2Bε(T91A) and eIF2Bβ(V316D) mutants obtained here differ from those of Fig. 2, in which only single subunits were expressed and where both showed decreased ability to form eIF2B complexes. In the earlier experiments, the overexpressed subunit must compete with the corresponding endogenous polypeptide for binding to a small pool of the other four eIF2B polypeptides. It thus appears that eIF2Bε(T91A) and eIF2Bβ(V316D) can indeed form complexes but fail to compete efficiently with the endogenous WT eIF2Bβ.

To test this, we examined whether the mutant eIF2B proteins could compete with WT myc-tagged eIF2Bβ and -ε to form holocomplexes. As expected, coexpression of WT myc-eIF2Bβ with His-myc WT eIF2Bβ or the E213G mutant reduced somewhat the amounts of the other subunits that bound to the His-myc-tagged polypeptide (Fig. 3C). However, coexpression of WT eIF2Bβ almost completely blocked the association of the eIF2Bβ(V316D) mutant with the other subunits.

In the case of eIF2Bε(T91A), we saw only a modest effect from expressing WT-myc eIF2Bε alongside the T91A-His-myc protein (Fig. 3D). Surprisingly, we also saw that, along with the His-myc-eIF2B polypeptide, the slightly smaller myc-tagged eIF2Bε polypeptide was also present in the pull-down, suggesting that eIF2Bε might dimerize, which could account for the small decrease in recovery of the T91A mutant. To confirm this, we coexpressed HA-tagged eIF2Bε with His-myc-eIF2Bε. After IP with anti-HA, His-myc-eIF2Bε was indeed clearly seen (Fig. 3E, lower part, lane 1). As described above, eIF2Bε interacts with eIF2Bβ in eIF2B holocomplexes (2, 25). Parts of their sequences are homologous (2) and could be involved in the γ-ε interaction. Thus, one explanation is that, when eIF2Bε is expressed at high levels, it self dimerizes via the region resembling eIF2Bβ. To test this, we overexpressed myc-eIF2Bβ with both HA- and myc-tagged eIF2Bε: strikingly, only myc-eIF2Bε was now seen in the anti-HA IP (Fig. 3E, lower part, lane 2). Dimerization of eIF2Bε is thus likely an artifact of overexpression.

VWM mutations decrease the activity of eIF2. The major aim of this study was to assess whether, and how, VWM mutations affect the function of eIF2B. The above data show that most of the mutated subunits can efficiently form eIF2B complexes, at least when overexpressed with the four other subunits. We therefore assayed the activity of complexes containing recombinant WT or VWM mutant subunits after isolation from extracts of transfected cells by using NTA-resin and our standard assay with preformed eIF2-[3H]GDP complexes as substrate. Initially, we expressed a single WT and/or mutant eIF2B subunit and isolated complexes containing this polypeptide and endogenous subunits. Although eIF2Bε displays some activity on its own (6, 10), this is greatly enhanced by association with the other subunits (10), as verified by earlier studies with HEK 293 cells (39). Thus, although the pull-down material contains both free His-myc-eIF2Bε and complexes containing this polypeptide, the activity detected overwhelmingly reflects that of complexes. In all cases, complexes containing VWM mutants of eIF2Bε showed lower activity than WT complexes (Fig. 4A). The defect varied from mild for, e.g., eIF2Bε(R315G) to severe for the W628R mutation. No exchange activity was detected for NTA-pull-downs of the eIF2Bβ or eIF2Bε truncation mutants, consistent with their inability to form eIF2B holocomplexes and with the absence from the eIF2Bε frameshift mutant of the C-terminal catalytic domain (9).

By using the above approach, we could not obtain accurate data for the eIF2Bε(T91A) and eIF2Bβ(V316D) mutants as they do not efficiently form eIF2B holocomplexes. Thus, we tested the VWM mutant under study expressed together with the other four WT subunits, since they can form holocomplexes under this set of conditions (Fig. 3B, upper part). The eIF2Bε(T91A) and eIF2Bβ(V316D) mutants still displayed low intrinsic GDP/GTP exchange activity, in addition to their defective abilities to form complexes (Fig. 4B). Since Western analysis of these pull-downs shows similar amounts of all five recombinant subunits (Fig. 3B, upper part), these data also accurately reflect the activities of other VWM mutant eIF2B complexes. The data in Fig. 1 rule out the occurrence of diverse complexes of differing subunit compositions.
Some VWM mutations affect the binding of eIF2B to eIF2.
Since neither of the truncation mutants forms complexes and
both are thus effectively null mutations, we focused our studies
on the point mutations, which all reduce eIF2B activity. One
way in which they might do this is by altering the ability of
eIF2B to bind eIF2. This involves multiple interactions. A
lysine block in eIF2\(\beta\)/H9252 interacts with the extreme C terminus of
eIF2B\(\epsilon\) (1, 18, 19). The eIF2B\(\epsilon\)/H9251/H9252/H9254 subcomplex interacts with
eIF2\(\beta\)/H9251, especially when eIF2\(\beta\)/H9251 is phosphorylated (20, 25). To
study binding of eIF2B to eIF2, we analyzed Ni-NTA–
agarose pull-downs from cells transfected with vectors for the His-myc-
tagged WT-VWM mutant eIF2B\(\epsilon\) or -/H9252- and myc-tagged ver-
sions of the other four subunits for the presence of eIF2\(\beta\). In
most cases, complexes containing VWM mutants showed binding
to eIF2 similar to that shown by the WT complexes (Fig.
3B, upper part). However, in three cases marked differences
were seen. Firstly, complexes containing eIF2B\(\epsilon\)(W628R)
bound markedly less eIF2. Secondly, both the eIF2B\(\beta\) point
mutations tested here actually increase the binding of eIF2. In
other experiments, NTA–agarose pull-downs from cells transfected with vectors for the His-myc-
tagged WT-VWM mutant eIF2B\(\epsilon\) or -/H9252- and myc-tagged ver-
sions of the other four subunits for the presence of eIF2\(\epsilon\).

The VWM mutations do not confer a temperature-sensitive
phenotype. Clinically, the onset or deterioration of VWM
symptoms in children is associated with febrile infection (12,
34–36). It has therefore been suggested that VWM mutant
eIF2B complexes might show a temperature-sensitive phen-
type. We used a range of approaches to test this. WT or VWM
eIF2B polypeptides were expressed in HEK 293 cells, as de-
tailed above. In some experiments, we subjected cells to ther-
mal stress by incubating them at an elevated temperature,
usually 41°C. Cells were lysed and samples were processed for
analysis of formation of eIF2B complexes and an activity assay.
Elevated temperatures did not generally affect the integrity of
eIF2B complexes containing WT or VWM mutant eIF2B\(\epsilon\) or
-\(\beta\) (Fig. 5A and B): amounts of other subunits copurifying with
the myc-His-tagged eIF2B\(\epsilon\) subunit under study were the same
from lysates of cells kept at 37°C or were transferred for 2 h to
a higher temperature. In some cases, the total amount of the
tagged subunit fell at the higher temperature but not the
amount of the other subunits copurifying with it. Effects were
similar for the WT and VWM mutant polypeptides and likely
reflect greater instability of the excess single subunits than of
those that incorporated into complexes. In some experiments,
eIF2B complexes containing eIF2B\(\epsilon\)(T91A) did appear less
stable than those containing WT eIF2B\(\epsilon\) (Fig. 5C). It was also
possible that the intrinsic activity of VWM eIF2B is thermo-
sensitive. However, in multiple experiments, we saw no ther-
mosensitivity of WT or VWM eIF2B at temperatures that
ranged up to suprapathological values (44°C). This was true
whether the elevated temperature was applied to the cells prior to isolation of eIF2B complexes, following their isolation but prior to assay or during the assay.

Functional consequences of VWM mutations for specific protein synthesis in vivo. It was important to assess the effects of VWM mutations on the function of eIF2B in vivo. However, even though free eIF2Bε has only low activity, overall eIF2B activity in cells expressing it will increase, obscuring any effects due to impaired activity due to VWM mutations. We therefore focused on eIF2Bβ as it lacks intrinsic activity and 80% of eIF2B complexes in cells overexpressing eIF2Bβ contain the recombinant subunit (Fig. 2E). The translation of certain mRNAs whose 5′ UTRs contain multiple uORFs provides a sensitive readout for small decreases in eIF2B activity, e.g., the ATF4 mRNA is regulated in this way (13). We therefore used a vector encoding enhanced green fluorescent protein (EGFP) downstream of the ATF4 5′ UTR to assess alterations in eIF2B activity in vivo. The EGFP was cloned behind the ATF4 5′ UTR with an HA tag to facilitate detection (Fig. 6A). The control vector pEGFP yields an mRNA with a very short 5′ UTR devoid of uORFs.

As shown in Fig. 6B, treatment of HEK 293 cells with tunicamycin A (TnA) led to increased phosphorylation of eIF2α due to activation of the eIF2α kinase PERK (14). In cells transfected with empty vector or with vector for WT eIF2Bβ, the level of expression of HA-tagged EGFP was low under control conditions and increased after treatment with TnA (Fig. 6B). This is as expected, since the TnA-induced increase in eIF2α phosphorylation inhibits eIF2B, promoting translation of the ATF4-HA-EGFP mRNA. WT eIF2Bβ had little effect on the expression of HA-EGFP. In cells expressing either of the eIF2Bβ mutants, increased basal expression of HA-EGFP was observed even without TnA treatment (Fig. 6B and C), consistent with the data showing that both mutations impair the activity of eIF2B. These findings also imply that, in VWM patients, these mutations may increase translation of mRNAs such as that for ATF4, potentially leading (as ATF4 is a transcription factor) to changes in the expression of multiple genes. No change in expression of EGFP from the control vector was seen (Fig. 6C), implying that there are no gross changes in protein synthesis (Fig. 2E). This is consistent with data from direct measurements of protein synthesis (our un-

![FIG. 6. VWM mutants of eIF2Bβ enhance expression of EGFP from a reporter construct containing the ATF4 5′ UTR. (A) Diagram showing the reporter construct based on the 5′ UTR of the ATF4 mRNA. The broad arrows indicate that the reporter vector gives rise to two translation products, one of which lacks the HA tag (lower, broken arrow) (see panels B, C, and E). (B to E) HEK 293 cells were transfected with vectors for WT eIF2Bβ or the indicated VWM mutants or the empty vector (Vec), along with the reporter construct containing the 5′ UTR of mouse ATF4 ahead of the coding region for HA-tagged EGFP. (C) The control reporter (pEGFP) was used in place of the ATF4-based vector where indicated. In panels B, C, and E, samples were analyzed by SDS-PAGE and Western blotting by using the indicated antisera. (Anti-eIF4A was used as a loading control.) (D) Samples of lysate were processed for Northern blot analysis. Shown are the autoradiography data obtained with probes for GAPDH (top) and GFP (bottom) and the stained gel showing the positions of the 18S and 28S rRNAs (middle: the top and middle panels serve as loading controls). The ATF4-EGFP mRNA is, as expected, larger than the EGFP one. (E) Cells were labeled with [35S]methionine (see Materials and Methods), and samples of cell lysate were analyzed by IP with anti-GFP, SDS-PAGE, and fluorography to detect the labeled HA-EGFP polypeptides (positions indicated by arrowheads). Again, two bands are seen here, for the reason described above.

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published findings) and the fact that no growth defects have been reported for VWM patients with these mutations.

When cells expressing VWM mutants of eIF2Bβ were treated with TnA, a further increase in HA-EGFP expression was seen. Given that a high proportion of eIF2B complexes in cells transfected with eIF2Bβ cDNAs contain the recombinant subunit (Fig. 2E), the data suggest that all the mutant complexes probably remain sensitive to inhibition by eIF2α (eIF2α/P). Northern blot analysis revealed that mRNA expression was not altered by VWM eIF2B (Fig. 6D) and that increased rates of synthesis of EGFP were directly confirmed by [35S]methionine labeling (Fig. 6E).

Further analysis revealed that expression of the VWM mutant eIF2Bβ subunits, but not WT eIF2Bβ, by itself increased the phosphorylation of eIF2α (Fig. 6B, see also Fig. 3B, lower part). Thus, the elevated level of expression of EGFP from the ATF4-based reporter may stem both from increased eIF2α phosphorylation and the lower intrinsic activity of the eIF2B complexes containing the mutant eIF2Bβ. The overall point remains the same, that expression of the VWM mutants activates translation of mRNAs containing uORFs. Since eIF2B binds phosphorylated eIF2 more tightly than it binds dephosphorylated eIF2 (30), this finding may explain why more eIF2 is found associated with eIF2B complexes containing the eIF2Bβ mutants (Fig. 3B).

**DISCUSSION**

The aim of this study was to examine the effects of VWM mutations in eIF2B genes on the properties of human eIF2B. This is the first key step to understanding the molecular and cellular pathology of this devastating condition. The data described here provide the first information on the effects on the human eIF2B complex of mutations that give rise to VWM in humans and reveal that all the VWM mutations tested in either the β or ε subunit of eIF2B result in a partial loss of function. This is probably as expected, as VWM shows a recessive pattern of inheritance, and is in agreement with the data in reference 29 on the effects of VWM mutations on yeast eIF2B. Our data for the eIF2Bβ(V316D) mutant agree closely with those of the authors of reference 29, who found that the corresponding yeast mutation (V341D) also impaired complex formation and activity.

To try to explain the basis of the reduced activity of these VWM eIF2B complexes, we examined their ability to bind to eIF2, the substrate for eIF2B. Three mutations altered the binding of eIF2B to eIF2. The eIF2Bε(W628R) mutation markedly decreased the ability of the eIF2B complex to bind eIF2. The fact that this residue lies within the catalytic domain of eIF2Bε (9) may explain why it impairs substrate binding. It is important that this tryptophan residue is not one of the conserved tryptophans within the cluster of aromatic and acidic residues that constitute the eIF2β binding site at the extreme C terminus of eIF2Bε (1). This decrease in substrate binding may explain the reduced nucleotide exchange activity of eIF2B complexes containing the eIF2Bε(W628R) mutant. Both the eIF2Bβ and eIF2Bε missense mutants tested here showed increased binding to eIF2. This appears surprising since their activities are also lower than for the WT complex. However, their expression increases the level of phosphorylation of eIF2α, and the increased binding may thus reflect the higher affinity of eIF2α(P) for eIF2B than that of the nonphosphorylated factor (30). This would lead to inhibition of eIF2B activity. Thus, the observed effects of these eIF2Bβ mutants on the ATF4-based reporter may reflect both the reduced activity of complexes containing them and the increased phosphorylation of eIF2α (Fig. 6). It is unclear why expression of mutant eIF2β increases basal eIF2α phosphorylation. The accumulation of (unfolded) mutant eIF2Bβ may elicit a stress response leading to increased eIF2α phosphorylation, or the lower activity of the VWM mutant eIF2B complexes may lead to increased translation of stress genes (e.g., ATF4). It will be important to study whether eIF2α phosphorylation and stress protein expression are increased in cells from VWM. Our data provide no information to suggest that the VWM mutations affect the susceptibility of eIF2B to regulation by eIF2α phosphorylation.

Episodes of infection accompanied by fever cause deterioration of VWM patients, and it has been suggested (37) that VWM mutations might confer on the eIF2B complex a thermosensitive phenotype. However, in extensive studies that used a range of complementary approaches, we did not detect any general enhancement of the temperature sensitivity of VWM eIF2B. Like the WT complex, its activity was not affected by temperatures up to 44°C. Our data agree with those of Richardson et al. (29), who also found no evidence for thermosensitivity of yeast eIF2B complexes containing VWM mutant subunits. The link between VWM and pyrexia thus
appears more complex than a simple model where eIF2B complexes containing VWM mutant subunits are thermosensitive; perhaps the already decreased eIF2B activity in brain tissue of VWM patients is further reduced due to the increased phosphorylation of eIF2α that occurs during pyrexia (reviewed in reference 32) or after traumatic brain injury (26). This may have marked effects either on total protein synthesis or, more likely, on the translation of specific mRNAs.

It is known from other studies that inhibition of eIF2B due to increased phosphorylation of its substrate eIF2 leads to upregulation of the translation of certain mRNAs, e.g., those for GCN4 in yeast (16) and ATF4 in mammals (13). eIF2α phosphorylation—and thus regulation of eIF2B activity—also appears to be important for the control of other transcription factors (15, 17, 33). As both proteins are transcriptional regulators, this effect could in principle affect the expression of multiple genes. Given that VWM mutations impair the activity of eIF2B, we asked whether expression of VWM mutant eIF2B had a similar effect. Using a reporter construct in which the ATF4 5′ UTR was cloned upstream of the coding region for an HA-tagged version of EGFP, we showed that both the eIF2Bβ point mutants do indeed enhance the expression of the reporter protein. Thus, one anticipates that the translation of mRNAs that are regulated in this way would be upregulated in VWM patients. Examples of such mRNAs are already known in mammals (e.g., ATF4), and others likely await discovery. If such mRNAs were expressed in a tissue-specific manner (e.g., in brain), this could explain why the phenotype of VWM is primarily neurological. Alternatively, it could be that overall protein synthesis in neuronal tissue is especially sensitive to perturbations in eIF2B activity.

Further progress requires cells (in particular, brain cells) that homozygously express VWM eIF2B, e.g., the generation of transgenic knock-in mice bearing VWM mutations. Work towards this goal is already under way.

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