PKR downregulation prevents neurodegeneration and β-amyloid production in a thiamine-deficient model

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Brain thiamine homeostasis has an important role in energy metabolism and displays reduced activity in Alzheimer’s disease (AD). Thiamine deficiency (TD) induces regionally specific neuronal death in the animal and human brains associated with a mild chronic impairment of oxidative metabolism. These features make the TD model amenable to investigate the cellular mechanisms of neurodegeneration. Once activated by various cellular stresses, including oxidative stress, PKR acts as a pro-apoptotic kinase and negatively controls the protein translation leading to an increase of BACE1 translation. In this study, we used a mouse TD model to assess the involvement of PKR in neuronal death and the molecular mechanisms of AD. Our results showed that the TD model activates the PKR-elf2α pathway, increases the BACE1 expression levels of Aβ in specific thalamus nuclei and induces motor deficits and neurodegeneration. These effects are reversed by PKR downregulation (using a specific inhibitor or in PKR knockout mice).

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**Abbreviations:** AD, Alzheimer’s disease; TD, thiamine deficiency; CSF, cerebrospinal fluid; SmTN, submedial thalamus nuclei; VLN, ventral lateral nuclei; BACE1, β-site APP-cleaving enzyme 1; PKR, double-stranded RNA-dependent protein kinase; eIF2α, eukaryotic translation initiation factor-2α; PERK, protein kinase RNA-like endoplasmic reticulum kinase; BACE1, β-site APP-cleaving enzyme 1; PACT, PKR activator; HRI, heme-regulated eIF2α kinase; GCN2, general control nonderepressible 2; MDA, malondialdehyde; NeuN, neuronal nuclei; F-J B, Fluoro-Jade B; IBA1, ionized calcium-binding adaptor molecule 1; APP, amyloid precursor protein

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Thiamine (vitamin B1) deficiency (TD) induces regionally selective neurodegeneration in the mammal’s brains, particularly in specific thalamus nuclei (submedial thalamus nuclei (SmTN) and ventral lateral nuclei (VLN)) and cerebellum.1,2 TD-induced neuronal death is associated with a chronic impairment of oxidative metabolism and neuroinflammation associated with glial activation.3–5 Diminished thiamine-dependent processes in humans is not only associated with Wernicke–Korsakoff syndrome but also accompany other neurodegenerative disorders, such as Alzheimer’s disease (AD).6 Experimental TD has been used to model the pathogenesis and investigate the cellular mechanisms of neurodegenerative disorders. In mice, TD-induced oxidative stress enhances Aβ accumulation by regulating β-site APP-cleaving enzyme 1 (BACE1) maturation. These effects are amplified in AD mouse model.7,8

The double-stranded RNA-dependent protein kinase (PKR) is one of the four mammalian serine–threonine kinases—the others being HRI (heme-regulated eukaryotic translation initiation factor-2α (elf2α) kinase), GCN2 (general control nonderepressible 2) and PERK (protein kinase RNA-like endoplasmic reticulum kinase)—that catalyzes the phosphorylation of the α subunit of eIF2 in response to stress signals, leading to an inhibition of protein synthesis.9,10 Activation of PKR is induced by various triggers such as viral infection and endoplasmic reticulum or oxidative stresses11,12 and could be controlled by an interaction with its specific activator PACT (PKR activator), also named RAX in rodents. PKR phosphorylation is known to have a significant role in AD,13 and cerebrospinal fluid (CSF) PKR levels could be used as a potential diagnostic biomarker in AD patients.14,15 PKR is a pro-apoptotic kinase16 and can exert a number of toxic effects on neurons that could contribute to the functional and pathological alterations in AD brains. PKR also contributes to neurodegeneration and to the pathological molecular mechanisms observed in AD. PKR can mediate Tau phosphorylation induced by Aβ exposure in cell cultures.17 Additionally, several investigators have demonstrated that elf2α phosphorylation, via PKR-induced cellular stress, leads to increased BACE1 mRNA translation when, paradoxically, global protein translation is inhibited.18–21 These alterations of BACE1 translational control could be explained by a stress-dependent phenomenon of translation initiation.22–24 Moreover, PKR and elf2α are highly phosphorylated in SmTN and the cerebellum of TD mouse model. Analyses performed on cerebellar granule neurons exposed to a thiamine metabolic antagonist suggest that TD-induced neuronal death could be partially mediated by PKR activation.25

To date, all studies that have explored the deleterious role of PKR activation in neurodegeneration indicate that inhibition of PKR is a promising approach to inhibit pathological mechanisms. Moreover, recent studies have shown that the genetic lack of PKR enhances learning and memory in several behavioral tasks while increasing network excitability.26 The goal of this study was to assess the role of PKR
downregulation on neurodegeneration and Aβ production in a mouse model of neurodegeneration.

Results

**TD induces regional oxidative stress and inflammation.**
To assess the consequences of the TD-induced stress cascade in our model, we evaluated the expression of markers of oxidative stress and microglial activation by immunofluorescence in the brain of mice exposed to 10 days of thiamine deprivation (TD mice) compared with wild-type mice (WT). Malondialdehyde (MDA) is the end product of lipid peroxidation and is commonly used as an oxidative stress marker. Using an MDA antibody, immunolabelings in TD mice brain showed a strong increase in staining in the cortical (not shown) and the thalamic neurons (37% of MDA-positive neurons). In the thalamus, this staining is mainly found in two well-defined thalamic regions (Figure 1a): the SmTN, which is located between bregma levels –0.94 and –1.94, and the VLN, between bregma levels –0.94 and –1.94.27 SmTN and VLN exhibit an increase of MDA-positive neurons, +57% and +42%, respectively (Figure 1c). Neurons in WT mice showed weak MDA staining (5.8% of MDA-positive neurons; Figures 1b and c). Microglia cells in their inactive state are characterized by a highly branched (ramified) morphology. After activation, they gradually undergo a transition from ramified to round feature (amoeboid). The ratio of these two phenotypical states both stained by an IBA1 antibody (ionized calcium-binding adaptor molecule 1) can be used as an indicator of microglial activation. Quantified immunofluorescence analyses of IBA1-positive microglia (IBA1+) in the thalamus revealed a 247% increase in the number of amoeboid cells in TD mice compared with controls (Figures 1d and e). The percentage of activated microglia was even greater in the SmTN (+343%) and in the VLN (+289%) (Figure 1d).

**TD enhances PKR activation in the thalamus and cerebellum.** To determine whether the inflammatory responses and oxidative stress affect the phosphorylation and expression of PKR in TD mice, immunoblotting with antibodies directed against the threonine (446) phosphorylation site or the total PKR protein were performed in the thalamus, cerebellum, cortex and hippocampus (Figure 2a). Quantification of the

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**Figure 1** Markers of oxidative stress (MDA) and microglial cell activation (IBA1) in the TD thalamus. (a) Schematic sagittal representation of thalamic nuclei distribution: SmTN, and VLN. (b) MDA immunohistochemistry (green) on the sagittal sections of WT and TD thalamus. (c) Percentage of MDA-positive cells in total thalamus, SmTN and VLN in WT (n = 6) and TD mice (n = 6). (d) IBA1 labeling (red) on the sagittal sections of WT and TD thalamus. (e) Quantification of relative proportion of IBA1-positive microglial cell phenotypes: ramified shape (inactivated state) and amoeboid shape (activated state) in WT (n = 7) and TD mice (n = 7). Unclassified shape corresponds to intermediate state or cells that were unable to be quantified. Scale bars: 10 μm. **P < 0.01, ***P < 0.001
pPKR_{Thr446}/PKR ratio showed a strong increase after TD in the thalamus, cerebellum and cortex of 139, 95 and 43%, respectively (Figure 2b). By contrast, there was no significant change in pPKR in the hippocampus (Figure 2b).

To explore the signaling pathway at the origin of PKR activation in TD, we performed immunoblots of the PKR main stress-dependent inducer, RAX, and the data revealed a significant increase in RAX in the thalamus and cerebellum (Supplementary Figure S1).

PKR knockout (PKR$^{-/-}$) mice are viable, phenotypically indistinguishable from their wild type$^{28}$ and have been used in several studies to test the protective role of PKR.$^{26,29}$ As expected, we did not detect a 65-kDa PKR band in brain samples of PKR$^{-/-}$ mice (Supplementary Figure S2). We extended the above TD findings by counting pPKR_{Thr446}-positive immunoreactive cells in the whole thalamus and in specific nuclei in the SmTN and VLN known to be more strongly affected by TD (Figures 2c and d) in animals.

**Figure 2** Activation of PKR in TD brain studied by immunoblotting and immunohistochemistry in the thalamus and cerebellum. Immunoblot analyses (a) and corresponding quantification of pPKR_{Thr446} and full PKR (b) in WT and TD mice protein samples, in the thalamus, cerebellum, hippocampus and cortex normalized on Tubulin levels. WT ($n=14$) and TD ($n=14$). *$P<0.05$, **$P<0.01$, ***$P<0.001$. (c) Immunohistochemistry (green) with a pPKR_{Thr446} antibody on WT, TD and TD pretreated with PKRinh mice in sagittal sections of (c) SmTN and (e) cerebellum. (d) The positive cell counting confirmed that PKR is activated after TD treatment and its increase is attenuated by PKRinh administration in total thalamus, specifically in SmTN and VLN. (e) Cerebellar pPKR_{Thr446} patterns in WT, TD and TD+PKRinh. pPKR_{Thr446} staining is almost absent in granular layer (GL) and molecular layer (ML) but exhibits strong signals and increased after TD in Purkinje cells (PC), both in cell bodies and processes (arrow). (f) The percentages of PC with pPKR_{Thr446} nuclear localization have been quantified, and as expected, pPKR_{Thr446} staining is decreased in PC after PKRinh administration WT ($n=8$) and TD ($n=8$). Scale bars: 10 $\mu$m.

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pretreated (30 mg/kg per os) or not with a PKR inhibitor (PKRinh) provided by Sanofi. There was a significant enhancement of pPKRThr446 expression after TD exposure. This effect was decreased after PKRinh pretreatment (−51% in total thalamus, −52% in SmTN, −54% in VLN) (Figure 2d) were observed in both structures. A comparable TD-induced increase of pPKRThr446 staining was detected in the cerebellum where pPKRThr446 is mostly present in Purkinje cell bodies and their arborizations (arrows) (+192%) (Figures 2e and f). In TD mice pretreated with PKRinh, the number of PKR-positive neurons was significantly reduced (58.9%; Figure 2f).

PKR inhibition reduces eIF2α activation in TD. To test whether TD-induced PKR activation could control eIF2α phosphorylation, we performed immunoblot analyses in mice pretreated or not with PKRinh or in PKR−/− mice. PKRinh inhibits eIF2α phosphorylation in the thalamus (−31%) and cerebellum (−39%) (Figures 3a and b). After genetic knockdown of PKR, TD-eIF2α activation was also reduced (Figure 3b). Quantified immunofluorescence of peIF2αSer51-positive neurons has confirmed these effects in the studied structures (Figures 4a and b). We also found that immunostaining for peIF2αSer51 has a neuronal localization in the thalamus in the ventral posterolateral nucleus (VPL), with the strongest labeling in the SmTN and to a lesser extent in the VLN (Figure 4c).

In the cerebellum, TD-dependant activation of peIF2αSer51 was predominantly observed in cell bodies and processes of Purkinje cells. As expected, peIF2αSer51 staining is reduced by downregulation or inhibition of PKR (Figures 4b and d). The expression of peIF2αSer51 signals showed a high degree of colocalization with pPKRThr446 staining in the thalamus (not shown) and in cerebellar Purkinje cells (Figure 4e). An increased eIF2α phosphorylation was also detected by immunoblot or immunofluorescence in cortical structures (Supplementary Figures S3a and b) but not in the hippocampus (Supplementary Figure S3a). Of the four known eIF2α kinases, the most likely to be induced by vitamin B1 deficiency is PKR. Therefore we tried to determine whether PERK, activated by the ER stress/unfolded protein response, GCN2 and HRI could contribute to TD-eIF2α activation. Immunoblots of the thalamus, cerebellum and hippocampus samples from TD and control mice did not reveal any statistically significant variations of the pPERKThr980/PERK ratio and HRI levels (Supplementary Figures S3c and d). However, we observed a 38.3% increase \((P = 0.044)\) of pGCN2Thr399/GCN2 ratio in the compared with control (Supplementary Figure S3e).

PKR inhibition reduces neuronal loss and neurodegeneration owing to TD. The neuronal nuclei (NeuN) antigen was used as a specific neuronal marker to quantify neuronal cell loss under various pathological conditions. Cell counting of NeuN-positive neurons in TD and control mice did not reveal any significant decrease in the various brain areas evaluated: cortex, cerebellum, hippocampus, and thalamus (data not shown). However, the evaluation in two specific thalamus nuclei (Figure 5a), SmTN and VLN, displayed a significant decrease of neuron density (54 and 37%, respectively) (Figure 5b). The pretreatment of mice with PKRinh before TD induction or TD context in knockdown PKR (PKR−/−) significantly reduced neuronal loss observed in the TD-SmTN (−73%). Similar protective effect have been found by pretreating TD mice with PKRinh in SmTN (−42%) and VLN (−39%). We then investigated the presence of damaged neurons in different thalamic areas using FluoroJade B (F-J B), an anionic fluorescein derivative that has been reported to specifically stain degenerating neurons. As expected, we did not detect any staining in Wt mice (Figure 5c). In the TD thalamus, 97% of F-J B+cells were present in the SmTN or VLN. PKR inhibition did not affect the total number of F-J B-stained cells (Figure 5d). To take into account the decreased neuronal density owing to TD, we have adjusted the number of F-J B-positive neurons to the total number of neurons counted in Figure 5b. Downregulation of PKR significantly reduced the number of damaged neurons in TD SmTN (−62%) and VLN (−46%). Similar results were obtained in TD mice pretreated with PKRinh (57% and 30%, respectively) (Figure 5e).

On the contrary, we did not observe any effect of PKR inhibition on oxidative stress (Supplementary Figure S4) or microglial activation (Supplementary Figure S5). In summary, PKR inhibition prevents neuronal loss induced by TD but has no effect on TD-induced oxidative stress or neuroinflammation.

Finally, to determine whether TD could lead to caspase 3-mediated apoptosis mainly in the SmTN, we quantified the number of neurons positive for a cleaved-caspase 3 antibody (Figure 6a). There was a large increase in apoptosis (+98% of positive cells) in the SmTN (Figure 6b). In addition, cleaved-caspase 3 stainings was also increased in TD Purkinje cells (+46% of positive cells) (Figures 7d and e). Interestingly, cleaved-caspase 3 and pPKRThr446 staining showed a high degree of colocalization in the SmTN (Figure 6c) and cerebellum (not shown). Inhibition of PKR, either by PKRinh or in PKR−/− mice, significantly reduced TD-induced caspase 3 levels in SmTN neurons (35 and 44%, respectively) (Figure 6b) but not in Purkinje cells (Figures 6d and e).

PKR downregulation inhibited amyloidogenic amyloid precursor protein (APP) processing in vivo and Aβ production. Previous studies have shown that PKR-induced eIF2α phosphorylation could increase BACE1 translation.\(^\text{18,19}\) To evaluate the functional importance of the PKR-eIF2α pathway in amyloidogenic signaling in TD mice, APP processing was assessed by immunoblotting of APP, BACE1 mature form and amyloid oligomers (Figure 7a). Quantified immunoblot demonstrated that BACE1 protein and amyloid β levels are significantly increased in the TD thalamus (by 110 and 52%, respectively) (Figures 7b and d). The TD-induced Aβ increase was confirmed by using ELISA assay for soluble Aβ (Figure 7e). By contrast, there were no changes in mRNA levels of BACE1 measured by real-time quantitative PCR (Figure 7c) or in APP protein expression in TD mice compared with control mice (Figure 7f). After 10 days of TD, PKR−/− and PKRinh pretreated mice both exhibit a significantly reduced levels of Aβ levels in the thalamus (−39 and −33%). In summary, PKR blockade attenuates Aβ levels in TD mice.

We also investigated a potential role of PKR downregulation on Tau hyperphosphorylation by quantifying the levels of pTau on three hyperphosphorylated sites (AT8, AT100 and AT180)
and of two kinases, JNK and GSK-3β, known to phosphorylate Tau in stress conditions. Comparisons of TD with control mice did not reveal any difference in AT100, AT180, GSK-3β and JNK activation ratio (Supplementary Figure S6).

Decreasing PKR activation mitigates motor deficits. Marked motor deficits have already been described in several TD models. We therefore assessed motor coordination and balance in our TD mice using a rotarod apparatus (Figure 8a). Analyses of latency to fall in TD and control mice on an accelerating rotarod revealed a significant progressive decline of motor performance from day 8 in TD mice (Figure 8b). PKR downregulation, either by PKRinh administration or by using PKR−/− mice, significantly reduced motor decline at day 8 in TD mice (Figure 8b). After 10 days, motor performance scores of deficient mice were strongly impaired and were not modified by PKR inhibition. Rotarod results of TD mice have to be related to the general status of the mice, as evaluated in this study by body weight measurement (Supplementary Figure S7). In comparison to non-deficient mice, TD mice exhibit a dramatic weight loss starting at day 8 that is not reduced by PKR inhibition. Finally, we also measured pPKRThr446 and peIF2αSer51 in the gastrocnemius muscle in order to evaluate a role of PKR-eIF2α pathway inhibition in the muscle. Immunoblot analyses did not reveal any variation in the phosphorylation ratios of these two proteins (Supplementary Figure S8).

Discussion
In the present study, we provide experimental data supporting the concept that ‘in vivo’ inhibition of the double-stranded RNA activated protein kinase PKR, originally identified as a sensor of virus infection, has a neuroprotective action and can decrease Aβ production in a mouse model of neurodegeneration. TD induces a strong increase of brain PKR activation, particularly in the thalamus and cerebellum. These observations are consistent with the previous study of Wang et al., which showed a progressive increase of PKR phosphorylation...
on threonine 446 and 451, in these two structures, from the fifth day of TD. In our work, we have described the histological localization of pPKR Thr446 overexpression in TD brain. In the thalamus, TD-induced, PKR activation is mainly detected in the submedial and ventrolateral nuclei, which are also the brain areas most strongly affected by oxidative stress and inflammation. The expression of activated PKR in the cerebellum is mainly in the Purkinje layer that is partly comparable to the findings described by our group in the human brain affected by prion diseases that can trigger PKR activation.32 This expression pattern is also in line with the PKR mRNA profile visible on the Allen Brain Atlas, which shows a strong expression in Purkinje cells. Additionally, it is known that PACT, or its murine counterpart RAX, is responsible for cellular PKR activation in response to a variety of extracellular stresses.33,34 After ethanol exposure, which shares common features with TD, RAX protein located in Purkinje layer interacts with PKR to reduce neuronal survival.35 After TD, we have also observed a significant increase of RAX levels in the cerebellum and at a lower magnitude in the thalamus, suggesting that RAX could mediate PKR activation in this TD cerebellum.

In vivo inhibition of PKR has been recently shown to exhibit significant positive effects such as enhancing memory and learning in behavioral tasks while increasing network excitability26,36–38 and protecting animals from high-fat diet-induced insulin resistance and glucose intolerance.29 Previous studies in neuronal cultures have shown that PKR inhibition can attenuate neuronal death induced by various stresses, including Aβ neurotoxicity.39,40 In the present study, we used a well-described PKR knockout model28 and a selective inhibitor of PKR activation (provided by Sanofi) to further evaluate the role of PKR in neurodegeneration in vivo. One of the major findings of this work is the validation that PKR inhibition can protect against neuronal loss in a model characterized by oxidative stress and neuroinflammation. Both stresses have been previously linked to PKR activation.31,42 We demonstrated here that they act upstream

Figure 4  eIF2αSer51 cellular localization in the thalamus and cerebellum in TD mice. (a) and (b) Double-labeling of DAPI (4,6-diamidino-2-phenylindole; blue) and peIF2αSer51 (green) in the (a) SmTN and (b) cerebellum of WT, TD, TD+PKRinh and TD+PKR−/− mice. peIF2αSer51 in cerebellum is present in Purkinje cells’ cytoplasm and mainly increase after TD at the level of the cells bodies. Scale bars: 10 μm. (c and d) Quantification of positive neurons in SmTN and (e) VLN and (d) in Purkinje cells suggests that peIF2αSer51 phosphorylation is dependent of pPKRThr446 activation. *P < 0.05, **P < 0.01, ***P < 0.001. (e) Confocal analysis showed a colocalization of pPKRThr446 (green) and peIF2α (red) in TD cerebellum of the cerebellum. Boxes represent higher magnification at the level of Purkinje cells, where colocalization (yellow) is mainly present in cell bodies
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Figure 5  Regional TD-induced neuronal loss and neurodegeneration in the thalamus reduced by modulation of PKR activation. (a) Immunohistochemistry of neuronal nuclear marker NeuN (green) shows a decreased number of neuron in the TD thalamus compared with control mice, in particular in SmTN and VLN (surrounded areas). (b) The number of marked NeuN cells’ evaluation indicates that TD-induced neuronal loss is reversed by PKR inhibition in the SmTN and in VLN. WT (n = 8), TD (n = 8) TD+PKRinh (n = 8) and TD+PKR−/− (n = 7). (c) F-J B histofluorescence in thalamic regions (surrounded nuclei: SmTN and VLN) reveals, as visible in higher magnification boxes, neurodegenerative neurons in TD mice. Note that WT thalamus did not present a positive signal. (d and e) Relative analysis as number of F-J B+ cells in SmTN and VLN (d) or the percent normalized on the mean number of NeuN+ (e). The percentage of TD-damaged neurons is decreased by PKR inhibition (TD+PKRinh and TD+PKR−/−). WT (n = 8), TD (n = 8) TD+PKRinh (n = 8) and TD+PKR−/− (n = 7); ∗P < 0.05, **P < 0.01, ***P < 0.001.

of PKR-eIF2α pathway activation. However, these results did not allow us, at this stage, to determine the respective role in the TD model of neuroinflammation and oxidative stress in PKR activation.

Thalamic regional neuronal loss in TD rodents was described several times by the group of Gibson. Nonetheless, the cellular mechanisms underlying this neuronal loss are not fully understood. One of the major hypotheses to consider is the...
The proapoptotic effect of PKR activation, through triggering of a downstream caspase cascade. By suppressing PKR in our TD model, we probably reduce neuronal loss due to apoptosis and, consequently, the proportion of damaged neurons. This possibility is consistent with the immunohistochemical results revealing that enhanced cleaved caspase 3 levels are strongly colocalized with pPKR\textsubscript{Thr446} and significantly reduced after PKR downregulation in the thalamus. In the cerebellum, the increase of cleaved caspase 3 detected in Purkinje cells was not associated with a reduced number of these neurons. Ten days of TD in our model may not be sufficient to observe this phenomenon, which is detected in the human brains affected by Wernicke's encephalopathy. Moreover, inhibition of PKR also leads to a decrease in the number of damaged neurons marked by F-J B staining in TD SmTN and VLN.

PKR can control the inflammasome and is involved in the general process of innate immunity and inflammation. However, we did not detect any modifications of the levels of microglial activation or oxidative stress in TD mice pretreated with PKRinh or in PKR\textsuperscript{−/−} mice. This finding suggests that, in the TD model, PKR inhibition acts downstream of the initial abnormal pathways associated with the lack of vitamin B1 and that the neuroprotective and functional effects obtained with PKR inhibition could be more related to the lack of activation of intrinsic or extrinsic apoptotic pathways or by a late regulation of autophagy.
During the past years, *in vitro* studies have demonstrated a role for eIF2α kinases and, in particular PKR, in the translational control of BACE1 levels and in the amyloidogenic pathway.\(^{18-20}\) Here we demonstrate a similar role for PKR on amyloid processes *in vivo*. The activation of eIF2α by TD is strongly colocalized with pPKR\(\text{Thr}^{446}\) and is reduced by PKR downregulation. In parallel, the levels of the other major stress-induced eIF2α kinases, activated PERK and HRI remain
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unchanged in the TD thalamus and TD cerebellum. Activated GCN2 is slightly increased only in the TD thalamus. Together the finding from PKR−/−− mice and the use of PKR inhibitor suggest that eIF2α phosphorylation is mainly under the control of PKR in this TD model. A previous report has revealed that TD increases β-secretase activity without effect on γ-secretase activity, which leads to an increase of Aβ production. Our results support this action on BACE1 levels and that PKR inhibition can attenuate this increase in amyloid production and BACE1 expression. A previous work has shown that OS inhibitors in the TD model will be worth trying. Taken together, these results suggest that the increase of BACE1 levels in TD is regulated, at least in part, by a translational mechanism initiated by the PKR/eIF2α pathway. Zhao et al., have demonstrated that pyrithiamine treatment and diet-induced TD increases β-secretase activity without effect on γ-secretase activity, which leads to an increase of Aβ production. Our previous work has shown that OS inhibition can attenuate this increase in amyloid production results support this action on BACE1 levels and that PKR inhibition. This effect could be related to the neuroprotective

Figure 7 Inhibition of PKR-eIF2α pathway by PKRinh or in PKR−/− mice decreases BACE1 levels and Aβ (Amyloid β) production. (a) Immunoblot on thalamus extracts and quantification of (b) BACE1, (d) Aβ oligomers and (f) APP. The blots show that TD induces BACE1 maturation and Aβ production without altering APP levels. PKR inhibition partially reduced the effects of TD. WT (n = 8), PKRinh (n = 8), PKR−/− (n = 7) TD (n = 8), TD+PKRinh (n = 8) and TD+PKR−/− (n = 7). (c) Transcriptional activity on BACE1 is not modified in the TD thalamus. Effect of TD on BACE1 mRNA levels is assessed with quantitative reverse transcriptase–PCR. BACE1 mRNA levels were normalized to GAPDH (glyceraldehyde 3-phosphate dehydrogenase) mRNA levels. WT (n = 6), PKRinh (n = 5), PKR−/− (n = 5) TD (n = 5), TD+PKRinh (n = 5) and TD+PKR−/− (n = 5). (e) The TD-dependent increase of Aβ1–42 peptide was confirmed by a sandwich enzyme-linked immunosorbent assay (ELISA) method and is expressed as means ± S.E.M. Aβ1–42. The values from TD mice are expressed relative to the values from WT mice that were set to 1. Immunoblots and ELISA were both performed on six different groups of mice: WT (n = 8), PKRinh (n = 8), PKR−/− (n = 7) TD (n = 8), TD+PKRinh (n = 8), and TD+PKR−/− (n = 7). *P < 0.05, **P < 0.01, ***P < 0.001

Figure 8 The rotarod motor performance impaired in TD mice is rescued after PKRinh administration or in PKR−/− mice. (a) Diagram illustrating latency to fall measured on separate days reveals a strong decrease of motor performance in the three groups of TD mice. (b) Statistical analysis of measures for each animal of the delay between the first day of the experiment and on days 7, 8, 9 or 10. At the beginning of the decline (day 8), downregulation of PKR maintains motor performance of animals with TD. WT (n = 14), PKRinh (n = 14), PKR−/− (n = 12) TD (n = 14), TD+PKRinh (n = 14) and TD+PKR−/− (n = 12). *P < 0.05
effect of PKR inhibition in the thalamus and Purkinje cells and could be explained in several ways. (i) Neuronal loss occurs in the SmTN, which is a part of VPL and VLN of the thalamus. VLN serves as a central integrative functional division in the thalamic nuclei for motor control, receiving inputs from the cerebellum, striatum and cortex.53 (ii) Biochemical abnormalities of the PKR, elf2α and caspase 3 in Purkinje cells, which are known to regulate and coordinate motor movements. Further electrophysiological studies analyzing Purkinje cells in TD model will be needed in the future. The strong and global imbalance of the physiological state of TD animals, as illustrated by the dramatic weight loss observed in all TD mice (with or without PKR suppression) could explain this lack of motor effect of PKR downregulation at days 9 and 10 of TD. Other motor performance tests carried out at the beginning of the decline could strengthen this hypothesis. Moreover, we cannot exclude that a part of the progressive motor impairment observed in our model could be due to metabolic changes induced by TD, in particular heart damage.

In conclusion, our results demonstrate that in vivo inhibition of PKR can allow neuroprotection and functional improvement in a TD model of neurodegeneration. In addition, PKR can modulate the in vivo production of Aβ. It is known that PKR is activated in the AD brains and other human neurodegenerative disorders32,34 and that activated PKR is increased in the CSF of AD patients.15 In the future, pharmacological inhibition of PKR might therefore represent a valid target to slow neurodegeneration in several human diseases.

Materials and Methods

Mice. All animal experiments in this study were performed in accordance with the guidelines of the French Agriculture and Forestry Ministry for handling animals (decree 87/849, license A75-05-22). PKR knockout (PKR−/−) mice were obtained by disruption of the Pkr gene introduced in embryonic stem cells, which were injected into C57BL/6 blastocysts. The chimeric mice were bred to C57BL/6 mice and the progeny homozygous for the disrupted allele, as previously published.28

For brain preparation, mice were anaesthetized with pentobarbital (40 mg/kg, i.p.) and received an intracardial perfusion with cold PBS. Brains were removed and dissected on ice and then placed in 4% (w/v) paraformaldehyde in PBS for 4–6 h at 4 °C to allow fixation. The brain was then cryoprotected in 30% (w/v) sucrose in PBS, before being sectioned at 30 μm with a freezing microtome. Sections were processed for immunohistofluorescence; blocking solution was PBS containing 0.1% (v/v) Triton X-100 and 2% (w/v) normal goat serum. Bound proteins were visualized using the Odyssey Imaging System (Li-Cor Biosciences, Lincoln, NE, USA) and quantified with the Multi Gauge software (Fuji Film, Tokyo, Japan). ELISA was used to assess the Aβ1-42 levels in thalamus samples. Frozen extracts were centrifuged, and the soluble supernatant fractions were collected. The insoluble materials were then dissolved with lysis buffer containing 8% SDS, urea and 5 mM EDTA. Aβ levels were quantified using the mouse Aβ1-42 ELISA Kit (Innogenetics) following the manufacturer’s instructions. The absorbance was read at 450 nm using a 96-well plate reader, and Aβ levels were calculated from a standard curve.

F-J B histofluorescence. F-J B (a high-affluence fluorescent marker for the localization of neuronal degeneration) (Millipore) staining was carried out according to the method described in a previous study.29 In brief, the sections were first immersed in a solution containing 1% sodium hydrosulphite in 80% alcohol. They were then incubated in a solution of 0.06% potassium permanganate and transferred to a 0.0004% F-J B staining solution. With this method, neurons that undergo degeneration brightly fluoresce in comparison to the background.

Inmunohistochemistry and confocal imaging. The other hemibrain from each mouse was sagittally sectioned on a cryostat apparatus at 10 μm. Sections were processed for immunohistochemistry; blocking solution was PBS (phosphate buffered saline)0.25% gelatin5% normal donkey serum0.3% Triton. Primary antibodies and secondary donkey antibodies were incubated at 4 °C for 24 h and at RT for 2 h, respectively, in PBS/0.125% gelatin0.3% Triton. The primary antibodies used for immunohistochemistry were as follows: rabbit anti-pPKR-Thr446 (Sigma), rabbit anti-pelf2α Ser51 (Cell Signaling), rabbit anti-caspase 3 active fragment (BD Bioscience, San Jose, CA, USA), mouse anti-NeuN (Millipore), rabbit anti-MDA (Abcam), and rabbit anti-ionized binding molecule adaptor 1 (iba1) (Wako, Osaka, Japan). Secondary antibodies were donkey anti-mouse Alexa Fluor 488 (Invitrogen) or donkey anti-rabbit Cy3 (Jackson Laboratory, Bar Harbor, ME, USA). For double confocal immunohistochemistry with antibodies from the same species, we used labelled Fab techniques. This approach need to label individually with the Zenon Kit (Sigma) one of the primary antibodies with fluorescent labeled-Fab, according to manufacturer’s protocol.

Standard epifluorescence images were acquired on a Leica DM5000 microscope using a high-resolution camera (CoolSnap HQ, Roper Scientific, Sarasota, FL, USA). Confocal imaging was performed using a Leica SP2 confocal microscope (Leica Microsystems, Solms, Germany). The Metamorph software (Roper Scientific) was used for image acquisition.
Counting of all the positive cells for pPKR, pellFox, caspase 3, MDA, F-L B and NeuN were performed with the NIH Image software (NIH Image, Bethesda, MD, USA), as previously described. DAPI and the cyanine 3 or alexa 488 pictures were both background corrected using the rolling ball method. A threshold was then chosen using the 'Auto threshold' function of ImageJ. The distinct phenotypes of IBA1-positive microglial cells were blindly evaluated by manual counting and classified into three groups: 'amoeboid', 'ramified', and 'intermediate' (also referred to as 'unclassified').

Quantitative RT-PCR. Total RNA was isolated using the TRIzol reagent (Invitrogen) from SH-SY5Y cells. SuperScript II Reverse Transcriptase Kit (Invitrogen) with random decamer priming was used to synthesize the first-strand cDNA from samples with an equal amount of total RNA (1 μg), according to the manufacturer’s instructions. cDNA samples were forwarded to amplification with specific primers for BACE1 and GAPDH using SYBR Green Technology (Invitrogen) and the Stratagene MX3000P PCR cycyter (Agilent Technologies, Masssay, France). Human BACE1 primers (forward 5'-GACGGCTACTACCTGAGA-GA-3' and reverse 5'-GATCCACGAGGATGTGAC-G3'), human GAPDH primers, (forward 5'-AATCC CA/TCATCTGTCCTCC-3' and reverse 5'-GGACTCCACGACGTA/CTACA-3') were used in the present studies. cDNA amplification was carried out as follows: denaturation at 94 °C for 2 min, followed by 35 cycles of denaturation at 94 °C for 15 s, primer annealing at 62 °C for 30 s, and extension at 72 °C for 40 s. A final extension was carried out at 72 °C for 10 min, ending with a 4 °C hold cycle. All assays were performed in triplicate, and BACE1 and PKR mRNA levels were denaturation at 94 °C for 2 min, followed by 35 cycles of denaturation at 94 °C for 15 s, primer annealing at 62 °C for 30 s, and extension at 72 °C for 40 s. A final extension was carried out at 72 °C for 10 min, ending with a 4 °C hold cycle. All assays were performed in triplicate, and BACE1 and PKR mRNA levels were normalized with the expression levels of GAPDH. Relative levels and gene copy numbers were calculated using the deltaCp method, as described by Pfaffl.

Statistical analysis. Statistical analysis was performed using GraphPad Prism 5 (GraphPad Softwares, Inc., La Jolla, CA, USA). Results were considered significant when P < 0.05 using a non-parametric Mann and Whitney test.

Conflict of Interest
Professor Jacques Hugon is a consultant for Roche, Sanofi, Novartis, Xigen, Elsai and Lundbeck. Dr. Claire Paquet is a consultant for Lilly and Novartis. Dr. Julien Dumurgier is a consultant for Novartis. Dr. Patrick Bernardelli, Dr. Véronique Taupin, Dr. Thomas Rooney and Dr. Laurent Pradier are fully employed by Sanofi. The remaining authors declare no conflict of interest.

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References
1. Baker KG, Harding AJ, Halliday GM, Kril JJ, Harper CG. Neuronal loss in functional zones of the cerebellum of chronic alcoholics with and without Wernicke’s encephalopathy. Neurosci 1998; 91: 429–438.
2. Chornyy S, Parkhomenko J, Chorna N. Thiamine deficiency caused by thiamine antagonists triggers upregulation of apoptosis inducing factor gene expression and leads to caspase 3-mediated apoptosis in neurally differentiated rat PC-12 cells. Acta Biochim Pol 2007; 54: 379–386.
3. Calingasan NY, Huang PL, Chun HS, Fabian A, Gibson GE. Vascular factors are critical in selective neuronal loss in an animal model of impaired oxidative metabolism. J Neuropathol Exp Neurol 2000; 59: 207–217.
4. Karapapagounder SS, Shi Q, Xu H, Gibson GE. Changes in inflammatory processes associated with selective vulnerability following mild impairment of oxidative metabolism. Neurobiol Dis 2007; 26: 353–362.
5. Ke ZJ, Gibson GE. Selective response of various brain cell types during neurodegeneration induced by mild impairment of oxidative metabolism. Neurochem Int 2004; 45: 361–369.
6. Heroux M, Raghavendra Rao VL, Lavioe J, Richardson JS, Butterworth RF. Alterations of thiamine phosphorylation and of thiamine-dependent enzymes in Alzheimer’s disease. Metab Brain Dis 1996; 11: 81–88.
7. Karapapagounder SS, Xu H, Shi Q, Chen LH, Pedrini S, Pechin D et al. Thiamine deficiency induces oxidative stress and exacerbates the plaque pathology in Alzheimer’s mouse model. Neurobiol Aging 2009; 30: 1567–1609.
8. Zhang Q, Yang Q, Li W, Fan Z, Sun A, Luo J et al. Thiamine deficiency increases beta-secretase activity and accumulation of beta-amyloid peptides. Neurobiol Aging 2011; 32: 42–53.
9. Dever TE. Gene-specific regulation by general translation factors. Cell 2002; 108: 545–556.
37. Medinas DB, Hetz C. Proteostasis impairment: at the intersection between Alzheimer’s disease and diabetes. Cell Metab 2013; 18: 771–772.
38. Jiang Z, Belforte JE, Lu Y, Yabe Y, Pickel J, Smith CB et al. elf2alpha Phosphorylation-dependent translation in CA1 pyramidal cells impairs hippocampal memory consolidation without affecting general translation. J Neurosci 2010; 30: 2582–2594.
39. Chang RC, Wong AK, Ng HK, Hugon J. Phosphorylation of eukaryotic initiation factor-2alpha (elf2alpha) is associated with neuronal degeneration in Alzheimer’s disease. Neuroreport 2002; 13: 2429–2432.
40. Page G, Rioux Bilan A, Ingrand S, Lafay-Chebassier C, Pain S, Perault Pochat MC et al. Activated double-stranded RNA-dependent protein kinase and neuronal death in models of Alzheimer’s disease. Neurobiology 2006; 139: 1343–1354.
41. De Felice FG, Ferreira ST. Inflammation, defective insulin signaling, and mitochondrial dysfunction as common molecular denominators connecting type 2 diabetes to Alzheimer disease. Diabetes 2014; 63: 2262–2272.
42. Goh KC, deVeer MJ, Williams BR. The protein kinase PKR is required for p38 MAPK activation and the innate immune response to bacterial endotoxin. EMBO J 2000; 19: 4292–4297.
43. Calingasan NY, Chun WJ, Park LC, Uchida K, Gibson GE. Oxidative stress is associated with region-specific neuronal death during thiamine deficiency. J Neuropathol Exp Neurol 1999; 58: 946–958.
44. Ke ZJ, Calingasan NY, DeGiorgio LA, Volpe BT, Gibson GE. CD40-CD40L interactions promote neuronal death in a model of neurodegeneration due to mild impairment of oxidative metabolism. Neurochem Int 2005; 47: 204–215.
45. Ke ZJ, DeGiorgio LA, Volpe BT, Gibson GE. Reversal of thiamine deficiency-induced neurodegeneration. J Neuropathol Exp Neurol 2003; 62: 195–207.
46. Garcia MA, Gil J, Ventoso I, Guerra S, Domingo E, Rivas C et al. Impact of protein kinase PKR in cell biology: from antiviral to antiproliferative action. Microbiol Mol Biol Rev 2006; 70: 1032–1060.
47. Heneka MT, Kummer MP, Stutz A, Delekate A, Schwartz S, Vieira-Saecker A et al. NLRP3 is activated in Alzheimer’s disease and contributes to pathology in APP/PS1 mice. Nature 2013; 493: 674–678.
48. Meng Y, Yong Y, Yang G, Ding H, Fan Z, Tang Y et al. Autophagy alleviates neurodegeneration caused by mild impairment of oxidative metabolism. J Neurochem 2013; 126: 855–861.
49. Shen S, Niso-Santano M, Ardjeman S, Takehara T, Malik SA, Minoux H et al. Cytoplasmic STAT3 represses autophagy by inhibiting PKR activity. Mol Cell 2012; 48: 667–680.
50. Tamagno E, Bardini P, Oibili A, Vitali A, Borghi R, Zaccaro D et al. Oxidative stress increases expression and activity of BACE in NT2 neurons. Neurobiol Dis 2002; 10: 279–288.
51. Zhao J, Sun X, Yu Z, Pan X, Gu F, Chen J et al. Exposure to pyrithiamine increases β-amyloid accumulation, Tau hyperphosphorylation, and glycogen synthase kinase-3 activity in the brain. Neurotox Res 2011; 19: 575–583.
52. Jones BJ, Roberts DJ. The quantitative measurement of motor inco-ordination in naive mice using an accelerating rotarod. J Pharm Pharmacol 1968; 20: 302–304.
53. Tlamsa AP, Brumberg JC. Organization and morphology of thalamocortical neurons of mouse ventral lateral thalamus. Somatosens Mot Res 2010; 27: 34–43.
54. Band Y, Omuki R, Katayama T, Manabe T, Kudo T, Taira K et al. Double-strand RNA dependent protein kinase (PKR) is involved in the extrastriatal degeneration in Parkinson’s disease and Huntington’s disease. Neurochem Int 2006; 48: 11–18.
55. Liu JY, Timm DE, Hurley TD. Pyrithiamine as a substrate for thiamine pyrophosphokinase. J Biochem 2006; 139: 6601–6607.
56. Schmued LC, Hopkins KJ. Fluoro-Jade B: a high affinity fluorescent marker for the localization of neuronal degeneration. Brain Res 2000; 874: 123–130.
57. Schmuck M, Temme T, Heinz S, Baksmeier C, Mosig A, Colomina MT et al. Automatic counting and positioning of 5-bromo-2-deoxyuridine (BrdU) positive cells in cortical layers of rat brain slices. Neurotox Res 2014; 43: 127–133.
58. Pfaffl MW. A new mathematical model for relative quantification in real-time RT-PCR. Nucleic Acids Res 2001; 29: e45.