Multiple mechanisms repress N-Bak mRNA translation in the healthy and apoptotic neurons

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N-Bak is a neuron-specific BH3-only splice variant of pro-apoptotic Bcl-2 family member Bak. We have shown that its mRNA is stable in the neurons, whereas the protein cannot be detected by antibodies, suggesting a strong translational arrest of the mRNA. Here we identify two regulatory elements in the N-Bak mRNA that significantly repress translation in the luciferase reporter assay: an upstream open reading frame in the 5′-untranslated region (UTR) and naturally spliced exon–exon junction downstream of the premature translation termination codon in the 3′ UTR. We also show that N-Bak mRNA is stored in granular structures in the sympathetic neurons and stays in these granules during intrinsic apoptosis. Finally, we confirm the absence of N-Bak protein by quantitative mass spectrometry analysis in the healthy, apoptotic or stressed sympathetic and cortical neurons. We conclude that N-Bak mRNA is translationally repressed by multiple mechanisms, and the protein does not participate in the classical apoptosis or cellular stress response.

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Abbreviations: NGF, nerve growth factor; SCG, superior cervical ganglion; UTR, untranslated region; uORF, upstream open reading frame; EJC, exon junction complex; LNA, locked nucleic acid; DIV, days in vitro; PTC, premature stop codon; NMD, nonsense-mediated decay; NMTR, nonsense-mediated translational repression; Fluc, Firefly luciferase; Rluc, Renilla luciferase; eGFP, enhanced green fluorescent protein

Post-transcriptional regulation of mRNA is a crucial aspect determining gene expression pattern in tissues and organs. It covers not only the splicing, modification, translation and degradation, but also stabilization, proper localization and storage of the mRNA. Many mRNAs are not translated constituatively but are stored in the repressed mode and translated only under certain conditions. Various regulatory elements in the untranslated regions (UTRs) of the mRNAs are frequently involved in the control of mRNA translation and stability, and also targeting to specific cellular compartments. In the 5′ UTRs, short upstream open reading frames (uORFs) and stable stem-loop secondary structures can interfere with mRNA translation. In the 3′ UTRs, the sequence elements can be recognized by distinct proteins or regulatory RNAs, like microRNAs (miRNAs) that can cause translational arrest of the mRNA. Also, the mRNA processing history, reflected in the exon junction complexes (EJCs) at splice junctions, is involved in mRNA localization, surveillance and translational control. Of interest, nonsense-mediated translational repression (NMTR) has recently been described, where EJC in the context of premature termination codon (PTC) represses mRNA translation without affecting its level. Thus, the interplay of various RNA-binding factors and mRNA-specific regulatory elements determine the post-transcriptional fate of the mRNA.

Bak is the BH1-3 pro-apoptotic member of the Bcl-2 protein family that, together with the related protein Bax, triggers mitochondrial changes in the intrinsic apoptosis. When activated, Bak and Bax generate pores on the mitochondrial outer membrane whereby the mitochondrial proteins (cytochrome c, DIABLO/Smac, EndoG, AIF and so on) are released to the cytosol to advance apoptosis. We have shown that unlike all other studied cells, neurons do not express Bak as its pre-mRNA is entirely spliced into mRNA encoding a putative BH3-only protein, named N-Bak. Thus, the neurons express only Bax, and their mitochondrial apoptosis depends solely on Bax. In the same studies, we demonstrated that, despite the presence of mRNA, N-Bak protein is not detectable in the brain or cultured primary neurons. We also showed that the absence of N-Bak protein is not caused by rapid degradation of its mRNA or proteasome-mediated degradation of the protein. Instead, the N-Bak mRNA appears to be stable suggesting its translational arrest and storage in the neurons.

Here, we set up to address the mechanisms of N-Bak mRNA repression more in detail. We identified an inhibitory uORF at the 5′ UTR of N-Bak mRNA that represses luciferase reporter translation. Also, a naturally spliced exon–exon junction downstream of PTC in the 3′ UTR of N-Bak mRNA strongly represses translation of the luciferase reporter. We show that endogenous N-Bak mRNA is localized to granular structures in the neurons. Importantly, endogenous N-Bak mRNA was not translated in the neurons during mitochondrial apoptosis or cellular stress, as shown by antibodies and quantitative mass spectrometry analysis, suggesting specific and robust translational arrest.
Results

Regulatory elements of the N-Bak mRNA UTRs that repress its translation. To ascertain regulatory elements in the UTRs of N-Bak mRNA that could participate in its translational repression, we prepared reporter constructs where the UTRs with or without mutations in the candidate control regions were fused upstream or downstream of Firefly luciferase (Fluc) coding sequence (Figure 1). Sympathetic neurons were microinjected with these constructs together with the Renilla luciferase (Rluc) reporter plasmid and relative luciferase activities were measured 48 h later. In addition, the mouse neuroblastoma Neuro-2a cells were transfected with reporters and analyzed for the relative luciferase activities and reporter mRNA levels. Because of limited amount of the material, reporter mRNA levels were not analyzed from microinjected primary neurons.

Mouse Bak mRNA (NM_007523.2) (Figure 1) has a 5′ UTR of 253 nucleotides, present in both Bak and N-Bak transcripts. The 5′ UTR of Bak mRNA is not well conserved, but different species contain one or two uORFs, some with weak Kozak consensus sequence. We cloned full-length 5′ UTR in front of the Fluc reporter gene (5′UTR-Fluc, Figure 1). We choose to address the 189-nucleotide uORF in the mouse Bak/N-Bak mRNAs as its initiator AUG is in the context of the Kozak consensus sequence, whereas the second AUG of the 5′ UTR lies within the first uORF and lacks the Kozak sequence. We deleted a 38-nucleotide region from the N-Bak 5′ UTR encompassing the AUG of the first uORF (5′UTRmut-Fluc, Figure 1). This deletion significantly increased luciferase activity compared with the wild-type 5′ UTR (5′UTR-Fluc) in both cell types (Figures 2a and b). This is in line with our previous results showing the importance of Bak 5′ UTR on the translation of luciferase reporter in the superior cervical ganglion (SCG) neurons. Transcript levels for both 5′ UTR reporters, analyzed from transfected Neuro-2a samples by quantitative RT-PCR, were similar (Figure 2h). Hence, the 5′ UORF significantly represses translation in the luciferase reporter assay and may inhibit translation of the main ORF on the N-Bak mRNA in the neurons.

Figure 1 Diagram of Firefly luciferase reporter constructs used in the study. Two upper schemes present N-Bak and Bak mRNAs with 5′ upstream ORFs (5′uORF) and coding regions (black boxes), stop codon (STOP) and predicted microRNA sites on the 3′ UTR (black squares) are shown on the top. For N-Bak mRNA the addressed exon–exon junction (eej) and premature termination codon (PTC) are also shown. Lower schemes illustrate the reporter constructs. Firefly luciferase coding region (Fluc) is shown as a gray box. 5′UTRmut, deletion of 38 nucleotides from the 5′ UTR, including the initiation codon for the uORF. 3′UTRmut, the construct where the 3′ UTR region containing the predicted microRNA target sites is replaced with the heterologous sequence (gray line) about the same length. Fluc-Bak-3′UTR, the construct where the 191 nucleotides between the two stop codons 20 are not included. Also construct with wild-type 5′UTR and 3′UTR, named 5′UTR-Fluc-EJC-3′UTR, and construct carrying mutated critical regulatory elements in both UTRs, named 5′UTRmut-Fluc-3′UTR, are shown. EJC, the putative exon junction complex deposited at the eej after splicing off the 131-nucleotide intron (for clarity, intron is not shown on the scheme).
Figure 2  Dual luciferase reporter assays. Relative luciferase activities measured from the SCG neurons (a, c, and e) or Neuro-2a cells (b, d, and f) overexpressing the indicated reporter constructs. The neurons (3–4 repeats on the independent cultures) were microinjected with the reporter plasmids together with the plasmid for eGFP. The ratio of Firefly and Renilla luciferase (Fluc/Rluc) activity was determined 48 h later and was normalized to the number of eGFP-positive neurons, counted before the lysis. The Neuro-2a cells (6 repeats on the independent cultures) were transiently transfected and the Fluc/Rluc activity determined 48 h later. Shown are the means ± S.E.M. Statistical significance of the differences was estimated by Student’s t-test (a and b) or one-way ANOVA and post hoc Tukey’s test (b and c). *P<0.05, **P<0.01, ***P<0.001, the null hypothesis was rejected at P<0.05. (g) RT-PCR analysis of the indicated constructs with and without the 131-nucleotide intron in transiently transfected Neuro-2a cells using primers flanking the exon–exon junction sequence. Equal sizes of the amplified fragments show that the intron was correctly removed by splicing. Shown is the Fermentas GeneRuler 1-kb DNA Ladder molecular weight marker. (h) Quantitative RT-PCR analysis of the indicated constructs in transiently transfected Neuro-2a cells. Shown are the Cp values of Fluc constructs normalized to Rluc values (means ± S.E.M. of three independent experiments). There were no significant differences between the groups (one-way ANOVA and post hoc Tukey’s test)
Differently from Bak, 3'UTR of the N-Bak mRNA harbors a PTC in the second last exon followed by downstream exon–exon junction (Figure 1). Proteins of the EJC downstream of PTC can direct mRNA to rapid degradation via nonsense-mediated decay (NMD) or, in contrary, lead to repression of mRNA translation via NMT without affecting its stability.9,10 We tested if premature termination context in the N-Bak 3'UTR affects mRNA translation in the luciferase reporter assay. Three 3'UTR reporter constructs were generated (Figure 1), N-Bak 3'UTRs, with and without a 131-nucleotide intron between exons 5 and 6, were cloned downstream of the Fluc stop codon (Fluc-EJC-3'UTR and Fluc-3'UTR respectively, Figure 1). Splicing of the intron in Fluc-EJC-3'UTR should lead to EJC deposition near the spliced site and the stop codon of Fluc would be recognized as PTC, thus mimicking a natural 3'UTR of N-Bak mRNA. Third construct with full-length Bak 3'UTR (Fluc-Bak-3'UTR, Figure 1), starting from its natural stop codon, was also cloned downstream of Fluc and served as a control. RT-PCR analysis with primers flanking exon–exon junction sequence in the N-Bak 3'UTR reporters Fluc-EJC-3'UTR and Fluc-3'UTR revealed equal lengths of PCR products, confirming correct splicing of Fluc-EJC-3'UTR transcripts in the transiently transfected Neuro-2a cells (Figure 2g). Quantitative PCR analysis revealed equal mRNA levels for all 3'UTR reporters, indicating that PTC context in the 3'UTR does not affect reporter mRNA level in the transiently transfected Neuro-2a cells (Figure 2h). Importantly, the spliced Fluc-EJC-3'UTR showed significantly reduced luciferase activity compared with the intronless reporter Fluc-3'UTR or Fluc-Bak-3'UTR control in both SCG neurons and Neuro-2a cells (Figures 2c and d). Thus, spliced exon–exon junction downstream of PTC, the special feature of the 3'UTR of N-Bak but not Bak mRNA, strongly represses translation of the luciferase reporter and may also contribute to translational repression of N-Bak mRNA in vivo. Fluc-3'UTR and Fluc-Bak-3'UTR did not show significant differences in the luciferase activity in either cell type (Figures 2c and d), indicating that the sequence between two stop codons in the N-Bak 3'UTR is not a determinant for translational repression. Reporter construct with both 5'UTR and 3'UTR of N-Bak mRNA, that is, having intact uORF and exon–exon junction (5'UTR-Fluc-3'-EJC-UTR), showed significant decrease in luciferase activity values in the SCG neurons and Neuro-2a cells, compared with similar reporter construct harboring mutated uORF in the 5'UTR and absence of exon–exon junction in the 3'UTR (Figures 2e and f). Both reporter constructs had equal mRNA levels (Figure 2h).

TargetScan 5.1 (http://www.targetscan.org/, release 6, 2 June 2012) predicts conserved binding sites for miR-29ab (UGGUGCUA), miR125ab-5p (CUCAGGGA) and miR26ab (UACUUGAA) on the 3'UTR of Bak mRNA. We next explored the role of miRNA sites in mediating repression of reporter mRNA translation. We used Fluc-EJC-3'UTR and Fluc-3'UTR reporters and replaced the region encompassing all three predicted microRNA sites with the heterologous sequence about the same length, resulting in two reporter constructs, Fluc-3'UTRmut and Fluc-EJC-3'UTRmut (Figure 1). Transcript levels of both constructs were similar in transiently transfected Neuro-2a cells (Figure 2h). RT-PCR with intron flanking primers showed equal lengths for both reporter mRNAs, confirming that Fluc-EJC-3'UTR was correctly spliced. In the context of spliced reporter transcript, Fluc-EJC-3'UTRmut, the luciferase activity was increased in a small but significant scale in both cell types, compared with Fluc-EJC-3'UTR (Figures 2c and d), whereas intron-free reporter Fluc-3'UTRmut had luciferase activity similar to that of Fluc-3'UTR (Figures 2c and d). Thus, miRNA binding sites mediated slight repressive effect on translation of the Fluc reporter, but only in the context of spliced 3'UTR.

We conclude that two sequence elements of N-Bak mRNA, namely uORF in the 5'UTR and exon–exon junction downstream of PTC in the 3'UTR, have strong repressive effects on its translation. Also, miRNAs may potentially contribute to N-Bak mRNA translational repression in the neurons.

N-Bak mRNA has granular localization in the SCG neurons. Translational repressed mRNAs are often compartmentalized in the cell to allow orchestrated control through interaction with specific regulatory factors and processing enzymes.21–24 We studied the intracellular localization of endogenous N-Bak mRNA in the sympathetic neurons from the SCG by in situ hybridization, utilizing locked nucleic acid (LNA) oligonucleotide probes designed specifically to detect a 20-nucleotide exon N within the N-Bak mRNA. The sense probe for exon N served as the negative control. As shown in Figure 3, N-Bak-specific LNA probe hybridized to punctate, granular structures. Such granules were observed in all neurons, most densely located in the soma, whereas the nuclei were essentially free of them. The sense probe gave mostly faint diffuse signal (Figure 3) with some scattered dots that were always sparse, whereas the antisense probe-hybridized neurons always exhibited dense granular pattern. Nonneuronal cells lacking N-Bak mRNA were always free of signal (not shown).

To analyze whether the N-Bak mRNA is stored in the P-bodies, the known storage/degradation sites of translationally repressed mRNAs, we attempted to colocalize it with Dcp1a, the marker of P-bodies. We first injected the SCG neurons with plasmid encoding for GFP-Dcp1a fusion protein. Subsequent immunostaining with anti-Dcp1a antibody revealed the specificity of the antibodies as they clearly recognized GFP-Dcp1a fusion proteins (Figures 4a–d). These antibodies were then used to immunostain the neurons hybridized with N-Bak LNA probes. The N-Bak mRNA-containing granular structures and endogenous Dcp1a-positive signal did not colocalize (Figures 4e–h). Thus, N-Bak mRNA is associated with the granules whose identity remains currently elusive.

N-Bak protein is not found in the healthy, apoptotic or stressed neurons. N-Bak protein is not detectable by immunoblot in the healthy or apoptotic neurons,15,16 suggesting that its mRNA is not translated. However, we cannot exclude technical insufficiency to detect N-Bak protein by antibodies. We therefore turned to quantitative mass spectrometry as an independent approach. The samples of the cultured SCG and cortical neurons were separated on two parallel gels, one used for immunoblot and another for mass spectrometry. In the immunoblot, anti-Bak
antibodies again did not recognize N-Bak from the healthy SCG neurons (Figure 5a, NGF+ lane), although a N-Bak-specific band of ~20 kD was detected from the sample of N-Bak-L76E-transfected HeLa cells (Figure 5a, N-Bak-L76E lane). In the cortical neuron samples, a band of ~21 kD was noticed (Figure 5a). We have observed this band occasionally in some cortical culture samples, but never in the samples of SCG neurons. The regions corresponding to the size of N-Bak protein but excluding the regions for Bak protein were cut out from the parallel gel of the same samples (Figure 5) and analyzed by quantitative mass spectrometry. Notably, N-Bak protein was not found in the neuronal samples but was identified only in the sample of the HeLa cells overexpressing it (not shown). Thus, two independent techniques confirmed the absence of N-Bak protein in the healthy SCG and cortical neurons. The protein of ~21 kD in the cortical cultures appears to be nonspecifically recognized by the anti-Bak antibodies.

We conclude that N-Bak mRNA is translationally repressed in the neurons. We also reasoned that this repression could be released during apoptosis or under cellular stress. BH3-only proteins are induced or activated in the intrinsic apoptotic pathway.25 Of interest, the intrinsic apoptosis induced by cellular stress is preceded by increased phosphorylation of α-subunit of the eukaryotic translation initiation factor 2 (eIF2α), leading to increased translation of certain mRNAs, often via short uORFs in their 5'UTRs.26,27 We tested whether N-Bak protein could be induced in the apoptotic or stressed neurons using immunoblot and mass spectrometry techniques. Cellular stress was induced in the cortical neurons by thapsigargin, an inhibitor of sarco/endoplasmic reticulum Ca2+ ATPases, that causes endoplasmic reticulum stress because of Ca2+ depletion and causes subsequent activation of intrinsic apoptotic pathway. Treatment of neurons with 10 nM of thapsigargin for 1 h induced maximal level of eIF2α phosphorylation (Figure 5b, upper panel) and these conditions were used in the following experiments. To induce mitochondrial apoptosis, we deprived SCG neurons of nerve growth factor (NGF) for 48 h or treated cortical neurons with 4 μg/ml of etoposide, the inhibitor of topoisomerase II, for 24 h. All treatments were done in the presence of caspase inhibitors. As shown by immunoblot, N-Bak protein was not induced by any of these treatments (Figure 5a and lower panel of Figure 5b). The absence of N-Bak protein in the samples of Figure 5a was confirmed by quantitative mass spectrometry analysis (not shown). Thus, translational repression of N-Bak mRNA is not released in the neurons during classical apoptotic or cellular stress conditions.

We also checked whether the N-Bak-positive granular structures were lost during mitochondrial apoptosis. To that end, we deprived the SCG neurons of NGF for 24 h and blocked the death by caspase inhibitor BAF. By that time, the apoptotic mitochondrial changes such as translocation of Bax to the mitochondria and release of cytochrome c to the cytosol had already occurred in the majority of the neurons28–31 and the BH3-only proteins should have been activated,32 as shown for DP5/Hrk,33,34 Bim35,36 and PUMA.37 The cultures were then hybridized with N-Bak-specific LNA probes in the antisense or sense orientation as above. The dense N-Bak-containing granules were still clearly visible in the neurons as exemplified on Figure 3e, whereas the sense-probe again revealed only sparse signal (Figure 3f). It must be noted that when the neurons become highly pyknotic, they bound strongly both antisense and sense LNA probes (Figures 3e and f), most probably nonspecifically. However, before this stage, all neurons showed granular pattern of N-Bak mRNA. In summary, we conclude that N-Bak mRNA is not released from repression during classical mitochondrial apoptosis.
apoptosis caused by NGF deprivation, in good agreement with our findings of the general absence of N-Bak protein.

Discussion

In this study, we show that the mRNA for N-Bak is under strong translational repression in neurons. The evidences for this conclusion are as follows: (1) 5′UTR and 3′UTR of N-Bak mRNA contain regulatory elements (inhibitory uORF and PTC with downstream splice junction, respectively) that strongly repress translation in the luciferase reporter assay; (2) endogenous N-Bak mRNA is stored in the granular structures; and (3) N-Bak protein cannot be detected in the healthy and apoptotic/stressed neurons by two independent techniques. Together with our earlier findings that N-Bak mRNA is stable and not rapidly degraded, these data strongly suggest a stable translational repression of N-Bak mRNA in the neurons, and that N-Bak does not contribute to classical mitochondrial apoptosis.

We show that presence of exon–exon junction downstream of PTC on the N-Bak 3′UTR is repressive to the reporter mRNA translation, whereas reporter mRNA levels were not affected. Translational repression and stabilization of mRNA via PTC and downstream EJC has been described as NMTR.9,10,38 Thus, our data suggest that N-Bak mRNA is stored in the granular structures; and (3) N-Bak protein cannot be detected in the healthy and apoptotic/stressed neurons by two independent techniques. Together with our earlier findings that N-Bak mRNA is stable and not rapidly degraded,16 these data strongly suggest a stable translational repression of N-Bak mRNA in the neurons, and that N-Bak does not contribute to classical mitochondrial apoptosis.

When analyzing the effect of mutated miRNA-binding region in N-Bak 3′UTR to reporter translation, we detected minor increase in luciferase activities, which was significant only in the reporter mimicking natural N-Bak 3′UTR. Although the effect was small, the role of miRNAs in the repression of endogenous N-Bak mRNA should not be underestimated. For example, the overexpressed reporter constructs may highly exceed the capacity of miRNAs available in the neurons such that their possible effects remain below detection. Recently others45 have shown that miR29b can repress the mRNAs for several BH3-only proteins, including N-Bak, in mature SCG neurons. In that study the repressive effect was studied using both the luciferase reporter and exogenous miR29b.

![Figure 4](image-url)

**Figure 4** N-Bak mRNA is not localized to P-bodies in the SCG neurons. (a–d) Anti-Dcp1α antibody recognizes its antigen. Shown are three SCG neurons overexpressing GFP-Dcp1α (b) and immunostained with anti-Dcp1α antibody (c). The corresponding phase-contrast image (a) and merged image (d) are also shown. Note that all GFP-Dcp1α granules are recognized by anti-Dcp1α antibody. The endogenous Dcp1α is not visible because of underexposure. (e–h) Deconvoluted merged stacks of confocal microscopic images of typical neurons colabeled with anti-Dcp1α antibodies, the marker for the P-bodies (f) and N-Bak-specific LNA probe (g). Corresponding phase-contrast image (e) and merged image (h) are also shown. Note that the granules do not colocalize. Scale bar, 25 µm.
constructs in transfected HEK293T cells, whereas we studied the exogenous reporter constructs in the natural neuronal milieu. Validation of the role of miRNAs in translational repression of the N-Bak mRNA requires further studies that are, however, challenging in the primary neurons.

In the actively growing, unstressed cells, P-bodies concentrate proteins not only involved in the translational repression, surveillance and degradation of the mRNAs, but also miRNA-mediated mRNA silencing. N-Bak mRNA does not colocalize with P-body marker Dcp1a, indicating that it is directed to some other type of granular structures, most probably for translational silencing and storage. Indeed, diverse granules have been described in different cells, including the neurons, but their identity and biology is not well understood. Nevertheless, as different RNA granules are known to contain translationally inactive mRNAs, storage of the N-Bak mRNA in the cytoplasmic granules could be an additional mechanism for its translational control.

In summary, N-Bak mRNA translation seems to be controlled via multiple mechanisms, including repression by uORF, premature termination context and also potentially by miRNAs. Possible synergy between multiple regulatory elements may lead to deposition of N-Bak mRNA into cytoplasmic granules where its fate can be further controlled by mechanisms that remain currently unknown and need to be addressed in further studies.

The complete absence of N-Bak band in our immunoblots was remarkable, raising doubts that it could result from technical complications. For example, the N-terminal epitope on endogenous N-Bak, the epitope for most anti-Bak antibodies, could be modified and therefore not recognized any more. The anti-Bak antibodies against other epitopes did not recognize the protein in the presence of caspase inhibitor QVD-OPh, or left untreated. Immunoblot was probed with anti-Bak antibodies from the nonneuronal cells whose numbers vary from culture to culture. Possibly the same nonspecific band was reported earlier by others as N-Bak protein, but as it was observed inconsistently, it may originate from the nonneuronal cells whose numbers vary from culture to culture. Possibly the same nonspecific band was reported earlier by others as N-Bak in the cortical cultures.

By common understanding, the mRNAs are translationally suppressed only transiently and released from the repression in the relevant conditions. We were surprised to find that N-Bak protein was not induced in the apoptotic and stressed neurons where such BH3-only protein could be required. Thus, the neurons totally repress N-Bak mRNA and N-Bak does not contribute to classical mitochondrial apoptosis in the neurons. By our current model, the neurons entirely avoid BH3-only protein, the mRNA for BH1-3 Bak is translationally completely repressed. Moreover, the translational block is complete, as even sensitive mass spectrometry did not detect any sign of N-Bak protein. A nonspecific band close to the size of N-Bak was occasionally recognized by anti-Bak antibodies from the cortical neuron samples, and its level was changed under some stress conditions. We do not know the identity of this protein, but as it was observed inconsistently, it may origin from the nonneuronal cells whose numbers vary from culture to culture. Possibly the same nonspecific band was reported earlier by others as N-Bak in the cortical cultures.

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Materials and Methods

Cloning and expression plasmids. N-Bak 5’UTR-Luc reporter plasmid, published earlier is named here as 5’UTR-Luc reporter construct by deleting a fragment between Avf and Nru restriction sites, thereby deleting first 39 nt in the 5’UTR, containing also initiation codon for longest uORF. N-Bak full length 3’UTR region, lacking an intron, was amplified by PCR from mouse embryo (E10.5) DNA with forward 5’-CATCTAGAGGTTTGGACCGGTCTCCG-3’ and reverse 5’-ATACGATCCCGCTGGGATACGAGGTCTG-5’ primers. The full-length 3’UTR region of N-Bak containing intron and full-length 3’UTR were amplified by PCR from mouse genomic DNA using forward N-Bak 5’- CATCTAGAGGTTTGGACCGGTCTCCG-3’ or forward Bak 5’-GGATATCTGAGGGATTTCTGAAGGGTCCG-3’ primers. Respective PCR fragments were digested with XbaI/BamHI restriction enzymes.
cloned into XbaI/BamHI sites of the pGL 4.13 vector (Promega, Madison, WI, USA), resulting in Fluc-3'UTR, Fluc-EJC-3'UTR and Fluc-Bak-3'UTR constructs. The 1 kb Xmal/RpmI restriction fragment covering highly conserved miRNA sites in Fluc-3'UTR and Fluc-EJC-3'UTR were replaced with 0.8 kb HpaI fragment from pSIREN-dsRED DNR vector (Clontech Laboratories, Inc., Saint-Germain-en-Laye, France), resulting in Fluc-3'UTRmut and Fluc-EJC-3'UTRmut constructs. N-Bak 3'UTRs with and without the addressed intron were also cloned into XbaI/BamHI sites of the 5'UTR-Fluc and 5'mutUTR-Fluc vector. This resulted in construct with natural 5'- and 3'-UTRs, named 5'UTR-Fluc-EJC-3'UTR and construct carrying mutated critical regulatory elements in both 5'UTR, 5'mutUTR-Fluc-3'UTR.

All constructs were verified by direct sequencing. N-Bak-L76E construct has been published previously.15 pEGFP-F1-Dcp1 plasmid is a kind gift from J Peränen (University of Helsinki, Helsinki, Finland; unpublished).

Cell culture and transfection. Neuro-2a and HeLa cells were grown in DMEM (Invitrogen, Carlsbad, CA, USA) containing 10% fetal bovine serum (HyClone, Thermofscien, Oxford, UK) and 100 μg/ml of Normocin (InvivoGen, Toulouse, France). Cells were transiently transfected with Lipofectamine 2000 (Invitrogen) according to the manufacturer's instructions. Neuro-2a cells were co-transfected in 24-well plate wells with pGL4.13 (Promega)-based Fluc and pGL4-73 (Promega) Rluc reporter plasmid mixtures in a 1:50 ratio and lysed 48 h later for dual luciferase assay or 36 h later for RNA isolation. HeLa cells were transfected in 6 cm dishes with the expression construct N-Bak-L76E and lysed after 36 h for immunoblot analysis.

Primary cell culture and microinjection. Neonatal sympathetic SCG neurons were cultured on polycarbonate-laminin (Sigma, St. Louis, MO, USA)-coated dishes or glass coverslips in the Neurobasal medium containing B27 supplement (Invitrogen) and 30 ng/ml mouse 25S (Promega). The cultures were maintained with NGF or deprived at 5 to 6 days in vitro (DIV) for 24 or 48 h of NGF by adding fresh medium with 2 μg/ml of function-blocking anti-NGF antibodies MAB25020 (Merck Millipore, Billerica, MA, USA) and 25 μg/ml pan-caspase inhibitor boap-acetyl-fluoromethyl ketone (BAF) (Caymanchem/EMD Biosciences, Darmstadt, Germany). Microinjection was performed as published previously.15,16 Dual Luciferase reporter assay. SCG neurons were microinjected with the following expression plasmid mixtures: 10 ng Fluc reporter in pg Rluc reporter in pGL4.13 and 5 ng/μl of Rluc reporter in pGL4.73. To identify successfully injected neurons, 5 ng/μl pEGFP-N1 (Invitrogen) plasmid was also included. Approximately 220–230 neurons were injected per each plasmid combination. pEGFP-F1-Dcp1 plasmid was injected at a concentration of 10 ng/μl. The neurons were grown with 5 μg/ml of pan-caspase inhibitor Q-VD-OPh (R&D Systems, Minneapolis, MN, USA) for 24 h and then fixed for immunostaining.

Cortical neurons were cultured as published previously.16,18 in cell stress induction experiments, etoposide (Sigma) at 4 μg/ml or thapsigargin (Sigma) at 5 μM were added to culture medium for 24 h or other indicated periods. For immunoblot or mass spectrometry analysis, cells from 40 sympathetic ganglia or ~3 × 10⁵ cells from cortical neuron cultures were used for sample preparation.

Immunoblot analysis. Cells were incubated on ice in the lysis buffer containing 150 mM NaCl, 50 mM Tris-HCl, pH 7.5, 1 mM PMF, 1 μg/ml phosphatase inhibitor cocktail solution (Roche Applied Science, Burgess Hill, UK; 0490684501), 100 μg/ml protease inhibitor cocktail (Sigma 8340), 1% Na-deoxycholate, 1% NP-40 and 0.1% SDS, collected to microtube and homogenized through the 25G needle with 6 strokes. Then, 5 × Laemmli sample buffer was added, the lysates were boiled for 5 min and cooled on ice for 2 min before centrifugation at maximum speed for 8 min. Next, 1/2 from neuronal cell cultures, 1/4 of Neuro-2a and 1/8 of HeLa cell culture samples were loaded to gel. Samples were run in 15% SDS-PAGE gels followed by blotting onto nitrocellulose membranes or direct fixation and gel staining. Nitrocellulose membranes were blocked for 1 h at room temperature (RT) with 5% BSA/TBST (TBS-0.1% Tween-20). Anti-Bak (Sigma, B-5857) or anti-P-eliF2a (Cell Signaling, Hitchen, UK; S 3398) primary antibodies and HRP-donkey anti-rabbit secondary antibodies (GE Healthcare, Little Chalfont, UK) in blocking buffer were used to detect N-Bak or P-eIF2a, respectively. SuperSignal West Pico Chemiluminescent Substrate (Pierce Biotechnology, Rockford, IL, USA) and LAS3000 imager (FujiFilm, Tokyo, Japan) were used to develop immunoblot signals. For reprobing, the filters were washed with 62.5 mM Tris-HCl, pH 6.8, containing 2% SDS and 100 mM 2-mercaptoethanol for 15 min at 55 °C and continued with several washings at RT with PBS and TBST.

RT-PCR and quantitative PCR analysis. Total RNA from transfected Neuro-2a cells was isolated with NucleoSpin RNA II RNA isolation kit (Macherey-Nagel, Düren, Germany) according to the manufacturer’s instructions. DNase treatment was performed separately, 50 μl DNase treatment reaction was kept at 37°C for 20 min containing 5 μl RQ1 DNase buffer, 2 μl Rnase-free DNase (all from Promega). RNA was cleaned with RNA Clean-up kit (Macherey-Nagel). Q-PCR products were analyzed in 2% agarose electrophoresis with GeneRuler 1 kb DNA Ladder (Thermo Fisher Scientific, Rockford, IL, USA). Quantitative PCR was performed essentially as published,16 except that primer annealing at 58 °C for 10 s was used. Fluc reporter transcript levels were detected with forward 5'-TCTATCTAAGGATTGTGCGCTTCGCT-3' and reverse 5'-ACCAGCAAGTCTTGGGAT-3' primers; to detect correct splicing of the reporter transcripts, as shown by the amplicon length of 450 nt. PCR products were microinjected with the following expression plasmid mixtures: 10 ng Fluc and 5 ng/μl of Rluc reporter in pGL4.73. To identify successfully injected neurons, 5 ng/μl pEGFP-N1 (Invitrogen) plasmid was also included. Approximately 220–230 neurons were injected per each plasmid combination. pEGFP-F1-Dcp1 plasmid was injected at a concentration of 10 ng/μl. The neurons were grown with 5 μg/ml of pan-caspase inhibitor Q-VD-OPh (R&D Systems, Minneapolis, MN, USA) for 24 h and then fixed for immunostaining.

Luciferase reporter assay. Luciferase activity assay was performed using Dual-Luciferase Reporter Assay System (Promega) as published earlier.16 Briefly, 48 h after microinjections, enhanced green fluorescent protein (eGFP)-positive living neurons were counted, resulting in average 180–210 positive cells per experimental point. The neurons were collected to the microtubes and lysed in 20 μl of 1 × passive lysis buffer (Promega). Neuro-2a cells were lysed 48 h after transfections in 300 μl of 1 × passive lysis buffer. The relative luciferase activity (ratio of Fluc/Rluc) was determined with GloMax 20/20 luminometer (Promega). In neurons, the ratio of Fluc/Rluc was normalized to the number of eGFP-positive neurons for each experimental point. For statistical analysis,收购 was performed by using GraphPad InStat 3 program (GraphPad Software, Inc., La Jolla, CA, USA).

In situ hybridization. For in situ hybridization, sympathetic neurons were seeded to polycarbonate-laminin-coated glass coverslips with NGF for 5 DIV, then deprived of NGF for 24 h or not deprived. The neurons were fixed with 4% PFA and 5 mM MgCl₂ in PBS, acetylated with 0.25% acetic anhydride in 0.1 M triethanolamine buffer (pH 8.0), washed once with 5 mM MgCl₂-PBS buffer, permeabilized with 0.1% Triton X-100 in PBS, washed with 5 mM MgCl₂-PBS, then two times with 5 × SSC, pH 5.0, and incubated in prehybridization buffer (50% formamide, 5 × SSC, pH 5.0) for 1 h at RT. N-Bak mRNA in situ hybridization was carried out for 16 h at 80°C, in hybridization buffer (10% dextran sulfate, 50% formamide, 2 × SSC, 2 mM vanadylribonucleoside complexes solution (Sigma 94742), 50 mM phosphate buffer, pH 7.4) in the presence of 5 pmol of 5'-3'digoxigenin-labeled antisense or sense LNA-based DNA oligonucleotides purchased from Exiqon A/S (Vedbaek, Denmark). Antisense oligonucleotide, specific to exon N sequence, was 5'-CTGTCGATATGGTGTGCTGCTGC-3' and sense oligonucleotide with opposite direction was 5'-CCTGATCAAGAGCGAAGG3'- reverse 5'-GATGACGCGAATCCTTGCCG-3' primers; and Cyclophilin G (NM_004792) levels were detected with forward 5'-CAAAACCAAGGAGGAGAAAGG3'- and reverse 5'-CAAGCTGCGAACCTCCTAGCG-3' primers. Acquired Cp values of Fluc and Rluc were normalized to the Cyclophilin G levels to ensure equal loading of the material. Thereafter, Fluc transcript levels were normalized to Rluc levels. Data were acquired from three independent biological replicas.

Immunostaining. In situ hybridized samples were incubated for 1 h at RT in the blocking buffer (1% BSA in 150 mM NaCl, 100 mM Tris-HCl, pH 7.4) and overnight in same buffer with primary antibodies: horseradish peroxidase-conjugated sheep anti-digoxigenin (Roche Applied Science 11207733901), with or without mouse monoclonal anti-Dcp1α (Abnova, Taipei, Taiwan; M06, clone 3G4). Cells were rinsed with 1 × PBS and signal was amplified for 5 min in the amplification buffer (1:200 diluted biotinyl-horseradish peroxidase-conjugated goat antirabbit secondary antibody; Abcam, Cambridge, UK). Neurons were visualized by 0.1% DAB substrate solution and counterstained with 1% aqueous solution of 0.1% Hoechst 33342.
diluted in 100 mM Tris-HCl, pH 8.0). After three washings with TBST, the slides were incubated with Alexa 488-conjugated streptavidin (Invitrogen) in the blocking buffer for 2 h at RT in dark, followed by additional washings with TBST. GFP-Dcp1Δ13-overexpressing cells were fixed with 4% PFA, 5 mM MgCl2 in PBS for 15 min at RT, washed twice with 1 × PBS/5 mM MgCl2 and treated as in situ hybridized samples except that mouse monoclonal anti-GFP antibodies in the blocking buffer were used and signal was not amplified. Alexa 568-labeled anti-mouse secondary antibodies (Molecular Probes/Invitrogen) were used to detect Dcp1Δ13 primary antibodies. Samples were mounted into Prolong Gold Antifade reagent (Molecular Probes) and stored at -4°C. Each experimental group was analyzed on three independent coverslips.

Immunofluorescence analysis. The fluorescent images were acquired using the confocal microscope TCS SP5 (Leica Microsystems, Wetzlar, Germany) equipped with LAS AF 1.82. The objective was Leica HCX PL APO 63 ×/1.3 GLYC CORR CS (21°C). The laser for the fluorescent images was OPLS 488 nm/270 mW with the beam splitter RSP 500. Doubly labeled cells were imaged in sequential mode with lasers OPLS 488 nm/270 mW and HeNe 633 nm/12 mW and beam splitter DD 488/561. For the transmitted light images, the Dodt gradient sequential mode with lasers OPSL 488 nm/270 mW with the beam splitter RSP 500. Doubly labeled cells were imaged in transmitted light with a Dodt gradient sequential mode with lasers OPSL 488 nm/270 mW and the beam splitter DD 488/561. Deconvoluted images were identically processed with ImageJ and Photoshop CS5 (Adobe Systems, Inc., San Jose, CA, USA) programs.

Mass spectrometry analysis. For mass spectrometry analysis, total cell lysates were separated in 15% polyacrylamid gel, the gels were fixed in 50% ethanol, 10% acetic acid in water for 1 h and with 50% methanol, 10% acetic acid in water overnight at room temperature. Gels were stained in 0.1% Coomassie blue R250, 20% methanol, 10% acetic acid solution for 3 h at RT and destained in 50% methanol, 10% acetic acid by changing buffer after every 30 min for 6 to 7 times. Gels were equilibrated in 5% acetic acid storage buffer. By using molecular weight markers as the landmarks, areas surrounding the position of 20 kDa, but excluding 50% methanol, 10% acetic acid by changing buffer after every 30 min for 6 to 7 times. Gels were equilibrated in 5% acetic acid storage buffer. By using molecular weight markers as the landmarks, areas surrounding the position of 20 kDa, but excluding 5×25 kDa (the position of Bak), 15 kDa and neighboring lanes were cut out from the gels by clean sterile blades and gel pieces were stored separately in the sterile micropetri dishes in storage buffer. Further in gel digestion54,55, NanoLC-MS/MS analysis, peptide identification and peptide quantification were performed as described earlier.53

Conflict of Interest

The authors declare no conflict of interest.

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