Transgenic Activation of Ras in Neurons Promotes Hypertrophy and Protects from Lesion-induced Degeneration

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Abstract. Ras is a universal eukaryotic intracellular protein integrating extracellular signals from multiple receptor types. To investigate its role in the adult central nervous system, constitutively activated V12-Ha-Ras was expressed selectively in neurons of transgenic mice via a synapsin promoter. Ras-transgene protein expression increased postnatally, reaching a four- to fivefold elevation at day 40 and persisting at this level, thereafter. Neuronal Ras was constitutively active and a corresponding activating phosphorylation of mitogen-activated kinase was observed, but there were no changes in the activity of phosphoinositide 3-kinase, the phosphorylation of its target kinase Akt/PKB, or expression of the anti-apoptotic proteins Bcl-2 or Bcl-XL. Neuronal Ras activation did not alter the total number of neurons, but induced cell soma hypertrophy, which resulted in a 14.5% increase of total brain volume. Choline acetyltransferase and tyrosine hydroxylase activities were increased, as well as neuropeptide Y expression. Degeneration of motorneurons was completely prevented after facial nerve lesion in Ras-transgenic mice. Furthermore, neurotoxin-induced degeneration of dopaminergic substantia nigra neurons and their striatal projections was greatly attenuated. Thus, the Ras signaling pathway mimics neurotrophic effects and triggers neuroprotective mechanisms in adult mice. Neuronal Ras activation might become a tool to stabilize donor neurons for neural transplantation and to protect neuronal populations in neurodegenerative diseases.

Key words: Ras • neuron • transgenic • protection • mouse

Introduction

The Ras/mitogen-activated protein kinase (MAPK)1 (extracellular signal–regulated kinase [ERK]) pathway is an evolutionary conserved signaling pathway found in all eukaryotic cells, including yeast (Marshall, 1996). In mammals, the Ras family comprises Ha-Ras, Ki-Ras, and N-Ras that are all lipid-anchored proteins in the inner face of membranes, cycling between the inactive GDP-bound and the signaling competent GTP-bound conformation. A network of upstream mechanisms converge onto the activation of Ras by using distinct types of Ras GTPase activating proteins (GAPs) or Ras-guanine nucleotide exchange factors (Wittinghofer, 1998) that respond to extracellular ligand–receptor triggered signaling cascades and intracellular second messengers (Reuther and Der, 2000). The mammalian RAS-guanine nucleotide exchange factor son of sevenless (mSOS1,2) is activated by receptor tyrosine kinases, whereas the guanine nucleotide release factors (Ras-GRF1,2) and the guanine nucleotide release protein are involved in calcium channel–mediated or G-protein–coupled receptor mechanisms. Conversely, a family of distinct types of Ras-GAP proteins regulate GTP hydrolysis: besides 120Ras-GAP and GAP1, neurofibromin (NF)-1, a protein that is deficient in patients with von Reckling-
housein disease type 1, is regulated by lipids (Bollag and McCormick, 1991; Lockyer et al., 1999). The brain-specific synGAP is negatively regulated by intracellular calcium concentrations (Kennedy, 1998).

In cultured neural cells, the intracellular functions of Ras strictly depend on the cell type investigated: in transformed astrocytes proliferation depends on Ras (Gua et al., 1997), yet in nerve growth factor responsive–PC12 pheochromocytoma tumor cells, Ras has been shown to stop proliferation and induce neurite outgrowth (Bar-Sagi and Feramisco, 1985; Noda et al., 1985). Moreover, neuronal survival has been demonstrated to depend on Ras, as shown by the intracellular application of function blocking anti-Ras Fabs fragments into cultured embryonic neurons in vitro (Borsaio et al., 1993) or inactive Ras (N17 Ras) (Xue et al., 2000). Consistently, cultured neurotrophin-deprived embryonic neurons derived from dorsal root ganglia, superior cervical ganglion, nodose ganglia, ciliary ganglia, or spinal cord are rescued from apoptosis by the intracellular application of activated Ras protein or cDNA (Borsaio et al., 1989; Nobes et al., 1996; Weng et al., 1996; Xue et al., 2000).

How can Ras activation direct specific downstream functions in neurons? The major pathways induced by Ras are Raf-1/B-raf/MAPK, Ral-GDS/phospholipase D/Rho, and phosphoinositol 3-kinase (PI 3-kinase)/Akt-1/phosphoinositide-dependent protein kinase (PKB) (Katz and McCormick, 1997). The latter signaling branch has been shown to promote survival in many neuronal systems (Williams and Doherty, 1999). In addition to PI 3-kinase, MAPK (ERK) activation is necessary for neuronal survival, but this pathway may regulate distinct apoptotic signaling cascades (Xue et al., 2000). In cultured cerebellar neurons, MAPK (ERK) activates the ribosomal Rsk-kinase, which, in turn, phosphorylates and thereby inactivates the proapoptotic activity of Bad (Bonni et al., 1999). The latter belongs to the Bcl-2 family of apoptosis-regulating proteins, forming homotypic or heterotypic dimers.

An in vivo function of Ras in the intact nervous system has been implicated from genetically induced deficiencies: expression of a dominant-negative mutant of c-Raf increased apoptosis during early development of the retina (Pimentel et al., 2000). Furthermore, inactivation of the neuronal-specific Ras-guanylate nucleotide release factor 1 in mice resulted in an impairment of long-term potentiation in the amygdala (Brambilla et al., 1997). The latter signaling branch has been shown to regulate survival in many neuronal systems (Williams and Doherty, 1999). In addition to PI 3-kinase, MAPK (ERK) activation is necessary for neuronal survival, but this pathway may regulate distinct apoptotic signaling cascades (Xue et al., 2000). In cultured cerebellar neurons, MAPK (ERK) activates the ribosomal Rsk-kinase, which, in turn, phosphorylates and thereby inactivates the proapoptotic activity of Bad (Bonni et al., 1999). The latter belongs to the Bcl-2 family of apoptosis-regulating proteins, forming homotypic or heterotypic dimers.

NGF-mediated survival of sympathetic chain neurons is completely uncoupled from Ras activity (Borsaio et al., 1993; Markus et al., 1997). Here, we expressed constitutively activated Ha-Ras selectively in neurons by using the neuronal promoter for the synapsin I gene, allowing us to directly investigate the effects of enhanced Ras activity on neuronal survival postnatally. Moreover, we asked if neuronal Ras activation could mimic typical neurotrophic effects, such as neuronal hypertrophy, induction of neurotropic Y (NPY), or increase in neurotransmitter synthesizing enzyme activities.

Materials and Methods

Establishment of the Transgene and Transgenic Mouse Lines

The 5’ nontranslated regions of the human Ha-Ras (Capon et al., 1983) and the rat Synapsin I (Sauerwald et al., 1990) genes were fused. The 3’ flanking region of the Ha-Ras gene, including its polyadenylation signal, was removed and substituted with a fragment containing internal ribosomal entry site (IRES)/LucZ (Kalinski et al., 1983; Ghnatt et al., 1991). A linear 1.1-kb DNA fragment, without vector sequences, was recovered and was suitable for mouse embryo manipulation.

Pronucleus DNA injections and embryo transfers were carried out according to standard procedures. 15-μg DNA from tail biopsies of each progeny aged 3 wk were restricted with KpnI, separated on 1.1% agarose gels, blotted to nylon membranes, and hybridized to a radioactively labeled Ras probe. Detected 1.9-kb DNA fragments represented 1.6 kb of the Synapsin I promoter and 0.3 kb of the Ha-Ras gene at their connection point, indicating transgene specificity. Outbred lines B6CBF1, B6C3F1, XMF1, and HimOF1XM1 were used as embryo donors and recipients. Founders were further bred with lines HimOF1, C57BL/6 (Institute for Animal Breeding and Genetics, University of Veterinary Medicine, Vienna, Austria) and NMRI (Central Institute for Laboratory Animal Breeding, Hannover, Germany).

Lines 46 and 50 were crossed back to NMRI background. They contained one integration site each with multiple transgene integrants. Whereas line 46 was bred to homozygocity, putative homozygous animals of line 50 could not be obtained. Ras-transgenic mice (syn Ras-TG) of lines 46 and 50 developed normally and did not show obvious behavioral differences to wild-type (w) littermates.

Western Blots

Tissue was lysed in ice-cold lysis buffer (50 mM Tris-HCl, pH 7.4, 150 mM NaCl, 40 mM NaF, 5 mM EDTA, 5 mM EGTA, 1% [%vol/vol] NP-40, 0.1% sodium deoxycholate, 0.1% SDS, 10 μg/ml aprotinin, 1 mM PMSF) using a 1-ml Dounce homogenizer. After centrifugation (13,000 g, 15 min, 4°C), the supernatants were collected. The protein concentration was determined using the DC protein assay (Bio-Rad Laboratories). Equal amounts of protein were separated on 12% polyacrylamide gels, as described by Laemmli. After blotting to 0.2-μm nitrocellulose (Schleicher & Schuell), the blots were blocked with 5% dry milk in TBS (10 mM Tris-HCl, pH 7.6, 150 mM NaCl, 0.05% Tween 20). The immunoreactive bands were detected using the indicated antibodies, horseradish peroxidase–conjugated secondary antibodies (Sigma–Aldrich) and the enhanced chemiluminescence substrate detection kit (Amersham Pharmacia Biotech). The following primary antibodies were used: anti–Ha-Ras (OP23; Calbiochem), anti–pan-Ras (OP40; Calbiochem), anti–phospho-MAPK (9106S; New England Biolabs), anti–synapsin I, anti–phospho-Akt (9271S; New England Biolabs), and HimOF1XM1, anti–Bcl-XL (clone 4; Signal Transduction Laboratories).

Glutathione-S-transferase–Ras-binding Domain Fusion Protein Pulldown Assay and PI 3-kinase Assay

Precipitation of GTP–Ras was performed essentially as described previously (de Rooij and Bos, 1997). For the determination of PI 3-kinase activity, tissues were Dounce homogenized on ice in lysis buffer (1% NP-40, 140 mM NaCl, 20 mM Tris-HCl, 1 mM MgCl2, 1 mM CaCl2, 1 mM NaVO4, 10% glycerol, 1 mM PMSF, 10 μg/ml aprotinin, pH 7.4). After clearing the lysate by centrifugation, the protein concentration was deter-
Primary Neuronal Cultures

Embryonic brains (embryonic day [E]14) were dissected and the midbrain was transferred to a tube containing MPBS** (10 mM Heps-NaOH, pH 7.4, 137 mM NaCl, 2.7 mM KCl, 8 mM Na2HPO4, 1.5 mM KH2PO4, 10 mM glucose, 1 mM sodium pyruvate, 1 mg/ml BSA, 2 mM glutamax [Sigma–Aldrich]), 6 μg/ml DNAse I, 5 μg/ml phenol red, 100 U/ml penicillin, 100 μg/ml streptomycin, 5.8 mM MgCl2). After incubation with trypsin (0.25%, 15 min, 37°C) [GIBCO BRL], the tissue was washed twice with DME/10% FCS and then followed by a 15 min wash in DME/10% FCS. Cells were treated either with 150 mM 6-hydroxydopamine (6-OHDA) for 90 min after 6 d in vitro or with 14-phenylpyridinium (MPP+) twice with MPBS.

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In Situ Hybridization and Anatomical and Morphometric Analysis

For in vivo magnetic resonance imaging measurements, five synRas-TG and five wt adult littermates were analyzed on a 4.7 tesla 30-cm bore Bruker DBX (Bruker), according to a published method (Allegrini and Sauer, 1992) adapted for mice.

For histochemical analysis, mice were killed (overdose of Nembutal, 60 mg/kg body weight i.p.) and perfusion-fixed, as described previously (Obst and Wahle, 1995). A surface view of each brain was drawn at 16× final magnification with a camera lucida. To process wt and synRas-TG brains at the same time, the hemispheres were paired, embedded in Tissue Tek, frozen, cut in 30-μm-thick serial sections, and processed for free-floating in situ hybridization, as described previously (Obst and Wahle, 1995; Wahle et al., 2000). For detection of the Ras-TG transcript, a 827-bp region of the lacZ cDNA cloned into pBS SK was used to generate digoxigenin-UTP-labeled antisense riboprobes (Roche). Furthermore, riboprobes against glutamic acid decarboxylase (GAD) mRNA and NPY mRNA were employed (Wahle et al., 2000). β-Galactosidase (lacZ) staining of fixed-brain sections (in 2% paraformaldehyde/0.2% glutaraldehyde, 30 min at 4°C) was performed using a 5-bromo-4-chloro-3-indolyl-β-D-galactopyranoside (X-gal)–staining solution. The morphometric analysis was carried out on thionin-stained brain sections of pairs of mice (siblings of one wt and one synRas-Tg) aged postnatal day (P) 13 and 40, as well as 2 and 6 mo. Somata of large pyramidal neurons of neocortical layer V of motor and somatosensory cortex and Purkinje cells of the cerebellum were outlined with a camera lucida at 1,000× magnification. Drawings were digitalized, and a somatic area in square micrometer was determined with an AutoCAD system for at least 150 somata of every experiment, which were then statistically compared using a nonparametric test. Some sizes were fed into size-frequency histograms.

Results

Neuronal Expression of Constitutively Active Ras in Neurons Leads to an Increase in Postnatal Cortical Brain Volume Without Affecting Neuronal Cell Number

The synapsin I promoter was used to drive the selective expression of V12-Ha-Ras in postmitotic neurons (Sauermann et al., 1990; Hoesche et al., 1993; Lietz et al., 1998) and minimize the risk of a deregulated cell number or tumor formation (Sweetser et al., 1997; Holland et al., 2000). The transgene encoded for a dicistronic mRNA, which directs the synthesis of two separate proteins, V12-Ha-Ras and LacZ (Fig. 1 A), and was tested for its Ras activity by transfection and induction of fiber outgrowth in PC12 cells (data not shown).

In the resulting heterozygous syn-Ras-TG mice (line 50), the levels of activated Ha-Ras (Ras-TG) protein were very low in the embryo, as revealed by Western blots (not shown) using antibodies selectively recognizing Ha-Ras, but not N-Ras or Ki-Ras protein. The major increase in V12-Ha-Ras expression occurred postnatally in the cortex of synRas-TG mice, which start to rise at P4, elevate three-fold by P13, and reach four- to fivefold basal levels by P40 (Fig. 1 B). In wt littermates, endogenous Ha-Ras increased only moderately during postnatal development. There was
Ras-TG expression was detected in the claustrum and cells by smooth, rounded nuclei with a large clear nucleolus. Restricted to neurons. The latter were distinguished from glia by Nissl-staining. Note the intense expression of the transgene in the region of neuronal somata of the hippocampus and also in the cortex (black arrows).

**Figure 1.** Generation of Ras-transgenic mice: developmental expression of constitutively activated Ras and increased brain volume in adult mice. (A) Schematic diagram of the bicistronic transgene construct. The Ras-TG expression was driven by the neuron-specific promoter of the synapsin I gene. The structure gen of the construct consisted of the gene for constitutively activated V12-Ha-Ras and the lacZ gene. An “internal ribosomal entry site” (IRES) is located downstream of the Ras stop codon. The bicistronic mRNA generated by this construct is terminated at the polyadenylation site of LacZ. Synapsin I promoter was from Rattus norvegicus; V12-Ha-Ras, human Val12–Ha-Ras oncogene (R. Jaggi, Inselspital Bern, Bern, Switzerland); IRES of encephalomyocarditis virus (J.E. Majors, Washington University School of Medicine, St. Louis, MO); LacZ was from *Escherichia coli.* (B) Developmental Ha-Ras expression in various brain regions and at different ages of Ras-transgenic (synRas-TG) mice and wt littermates. Homogenates of brain cortices or cerebella derived from synRas-TG mice and wt littermates were analyzed by Western blotting using an antibody recognizing endogeneous and activated transgenic Ha-Ras. Ras-TG expression was hardly detectable prenatally. In the cortex, there was a rapid postnatal increase reaching a 4.2-fold expression of activated Ras-TG, relative to endogeneous Ha-Ras at 40 d. Only moderate postnatal Ha-Ras increases were seen in wt cortex or in cerebella of wt or synRas-TG mice. Data are the means of two independent experiments, as obtained from densitometric measurements. (C) Horizontal sections of paired brain hemispheres derived from wt (left) or synRas-TG (right) mice after fixation and X-gal staining. Note the intense expression of the transgene in the region of neuronal cell somata of the hippocampus and also in the cortex (black arrows). The cortical brain volume is enlarged, leading to a caudal displacement of the cerebellum (compare white arrows). Bar, 1.8 mm. (D) Total brain volume and cortical brain volume are enlarged as determined from 13 contiguous coronal slices. Morphometric evaluation of wt (*n* = 5) or synRas-TG (*n* = 5) mice was performed using NMR imaging and a semiautomatic imaging software on a Silicon Graphics Indy computer.
Brain Region–Specific Expansion of Neuronal Cell Soma Diameter

The size-frequency histogram of cross-sectional areas of the layer V pyramidal neurons in the motor and somatosensory cortex showed a significant shift to larger soma areas, including size classes above the normal range at 6 mo old animals ($P \leq 0.0001$) (Fig. 2 A). This neuronal hypertrophy was developmentally regulated, corresponding to the neuronal Ras-TG expression levels. At P4, changes were barely detectable, i.e., differences were not significant, whereas, at P13, the increases in soma areas were highly significant ($P \leq 0.0001$) (data not shown). Neuronal hypertrophy and cortical expansion were most prominent in the frontal or parietal cortex (Fig. 2 C). In the cerebellum, the cross-sectional areas of the individual Purkinje neurons were not changed in their size distribution (Fig. 2 B).

Significant ($P \leq 0.0001$) increases in cortical soma area in layer V pyramidal neurons was also found in line 46 at 6 mo old.

Constitutive Activation of Ras and MAPKs (ERK1 and ERK2), but Not PI 3-kinase and Akt/PKB

To test whether the Ras-TG protein was active in neurons, we captured active GTP-Ras selectively with a peptide derived from Raf kinase (de Rooij and Bos, 1997). A major Ras activity was found in extracts of the hippocampus and cortex, but only low levels were detected in the cerebellum of synRas-TG mice. In wt littermates, the basal activity of endogenous Ras was hardly detectable under these experimental conditions (Fig. 3, second row). Because chronic treatment of responsive PC12 cells with NGF increases synapsin I synthesis (Romano et al., 1987), we tested if neuronal Ras-TG activity regulates endogenous synapsin levels in the brain. Fig. 3 A shows that the synapsin levels were not different between synRas-TG mice and wt littermates in all regions investigated. Moreover, the Ras-TG expression pattern reflected the endogeneous synapsin promoter activity, with low expression in the cerebellum and high levels in hippocampus or cortex, indicating that Ras-TG activity does not regulate the endogeneous synapsin I promoter.

As expected from the increased Ras activities in synRas-TG mice, phosphorylation of MAPKs (ERK-1 and/or ERK-2) was constitutively elevated in all three brain regions without any changes in MAPK protein expression (Fig. 3 A). The differential increase in phosphorylation of ERK-1 or ERK-2 was especially pronounced in the hippocampus and cortex where mainly ERK 1 was activated in all cases tested ($n = 3$).

Besides the MAPK pathway, Ras activates the catalytic subunit of PI 3-kinase subtypes as a downstream target

Figure 2. (A) Size-frequency distributions of cortical, hippocampal, and cerebellar neurons. Size-frequency distribution of large pyramidal neurons of layer V in the motor and somatosensory cortex to 66.74 ± 5.76 ($\times 10^3$ mm$^{-3}$, $P < 0.05$) in synRas-TG.
were determined in coimmunoprecipitates using either anti-pTyr antibodies or anti-Ras monoclonal antibody 13-259. Neither the phosphotyrosine-associated, nor the Ras-associated, PI 3-kinase activity was altered. Akt (PKB) phosphorylation: crude brain lysates were separated by SDS-PAGE, blotted on nitrocellulose. The amount of signaling active GTP-bound conformation of Ras proteins was detected using a pan-Ras antibody. Transgenic V12-Ha-Ras expression results in a major increase of GTP-Ras in the cortex and hippocampus, but only an attenuated increase is observed in the cerebellum. Endogenous synapsin I expression: in adult cerebellum, the relative levels of Synapsin expression are low compared with cortex and hippocampus. Note that the Ras activity in synRas-TG mice does not regulate the endogenous Synapsin levels. MAPK phosphorylation: crude brain lysates were separated by SDS-PAGE, blotted on nitrocellulose, and phosphorylation of the PI 3-kinase target Akt was detected using an antibody specific for MAPK (ERK1 and ERK2) phosphorylated on the activation-competent threonine and tyrosine. In synRas-TG mice, MAPK tyrosine phosphorylation in neurons is permanently enhanced, though protein levels (ERK1 and ERK2) are not affected (see below). MAPK phosphorylation is enhanced most prominently in the hippocampus, but only moderately in the cerebellum, thus, corresponding to the different synRas-TG expression levels. In the cortex, we observe a more selective ERK1 over ERK2 phosphorylation. MAPK—total protein content: equal amounts of protein were loaded in each slot, resulting in similar signals for total amounts of ERK1 or ERK2. PI 3-kinase activity: the levels of phosphatidylinositol 3-phosphate (PI[3]P) reaction products were determined in coimmunoprecipitates using either anti-pTyr antibodies or anti-Ras monoclonal antibody 13-259. Neither the phosphotyrosine-associated, nor the Ras-associated, PI 3-kinase activity was altered. Akt (PKB) phosphorylation: crude brain lysates were separated by SDS-PAGE, blotted on nitrocellulose, and phosphorylation of the PI 3-kinase target Akt was detected using an antibody specific for Akt phosphorylated on serine 473. The phosphorylation of Akt is not altered by the Ras-TG activity in neurons.

Enhanced Chat or TH Activities and Increased Neuropeptide Y Expression

Besides induction of neuronal hypertrophy, neurotrophic actions include the regulation of neurotransmitter synthesizing enzymes, such as Chat or TH. Specific Chat activity in homogenates containing the septal region from the basal forebrain of synRas-TG mice was 89% higher than in homogenates of wt littermates (942 ± 181 pmol/mg/min, n = 6 versus 498 ± 149 pmol/mg/min, n = 9, P < 0.001). TH activity was elevated by 30.6 ± 31% (n = 5, P < 0.05) in the mesencephalic forebrain region containing the dopaminergic neurons of the substantia nigra.

Next, we asked if the activated Ras-TG also affects the neurochemical properties of cortical inhibitory interneurons known to express TrkB-receptors (Gorba and Wahle, 1999). Fig. 4 shows that the staining intensity and the density of cortical NPY-expressing neurons clearly increased throughout the cortex, despite cortical expansion (from 54 to 80 cells/mm² in sagittal sections, P ≤ 0.01). In contrast to neocortex, NPY mRNA expression appeared to remain identical in hilus interneurons in the hippocampus and in neostriatal interneurons (not shown). The staining intensity and density of GAD (Fig. 4) or parvalbumin mRNAs (not shown) were not affected.

The motorneurons of the facial nucleus were investigated in more detail. In situ lacZ–mRNA hybridization revealed that in the facial nucleus region the typical large diameter motorneurons were clearly lacZ positive (Fig. 5 A). No positive glial cells were found throughout the brain (not shown). As a functional consequence of the Ras-TG expression, we found a 25.6% (P < 0.001, n = 100) increase in the cross-sectional area of the motorneuron somata (Fig. 5 B). Similarly, in the weak Ras-TG–expressing mouse line 46, an attenuated, yet significant, increase in cross-sectional area of the motorneuron somata was found (P < 0.005, n = 143, data not shown). Finally, we investigated if the Ras-TG activity might regulate the basal activity of Chat in motorneurons, as it was found in the septum. The Chat activity per facial nucleus was indeed elevated by 39% (n = 3) (Fig. 5 B, insert), suggesting an increased Chat activity per motorneuron, because their number remained unchanged in the synRas-TG mice.
Ras-TG Expression in Motorneurons of the Facial Nucleus Completely Prevents Their Degeneration after Nerve Lesion

To test whether the Ras activation not only prevents embryonic apoptosis (see Introduction), but also promotes protection against lesion-induced neuronal degeneration, we performed facial nerve lesions in young-adult mice (10 wk old). There was on average a 34% reduction in cell number of wt mice, whereas there was no reduction (3%, not significant) in the synRas-TG littermates, as shown by counting the number of nucleoli-containing neurons in every third section of the facial nuclei from five mice each (Fig. 6, B and C). Morphologically, neurons appeared healthy on the lesioned side, with no signs of degeneration or shrinkage in synRas-TG mice (Fig. 6 A).

Ras-TG Expression Attenuates Neurotoxic Insults by 6-OHDA, MPP⁺, and MPTP in the Substantia Nigra

We next investigated whether the Ras-TG was expressed in the dopaminergic neurons of the substantia nigra of the synRas-TG mice. Fig. 7 A shows an example of a TH-expressing neuron of the substantia nigra that was Ras-TG–positive. We then asked if the Ras-TG expression could induce protective mechanisms against the neuronal-type–specific toxins 6-OHDA or MPP⁺ (Lotharius et al., 1999). Mesencephalic embryonic cultures containing TH-positive neurons were established and neurons were allowed to mature for a period of 5 d. Treatment with 6-OHDA for 90 min or MPP⁺ exposure for 36 h reduced the number of TH-positive neurons to 36 or 25% of untreated controls, respectively. In cultures of synRas-TG mice, the loss was partially prevented resulting in a 48 or 34% enhancement of survival after 6-OHDA or MPP⁺ treatment (Fig. 7 B).

We also tested the striatal projection area of dopaminergic substantia nigra neurons. In unlesioned animals, there was a 22% increased level of dopamine in synRas-TG mice compared with wt littermates (Fig. 7 C). After two consecutive applications of MPTP, dopamine levels were reduced from 10336 ± 1489 ng/g (n = 10) to 4029 ± 335 ng/g (n = 8) in wt and from 12599 ± 1461 ng/g (n = 11) to 7459 ± 1572 ng/g (n = 8) in synRas-TG mice, thus indicating a substantial protection.

Discussion

To investigate the physiological role of the proto-oncogene Ras in the adult mammalian nervous system, constitutively activated V12-Ha-Ras was expressed postnatally in the brain of transgenic mice. By using the synapsin I promoter, expression of the Ras-TG was restricted to neurons, giving rise to healthy transgenic mice with a normal life span.

In the synRas-TG mice, we observed pronounced neuronal hypertrophy, one of the earliest described neurotrophic effects in embryonic chick dorsal root ganglion neurons and in sympathetic neurons (Levi-Montalcini, 1987). This hypertrophy was present in several populations of Ras-TG–expressing neurons, i.e., in pyramidal cells of the cortex and in N7 motorneurons of the nucleus facialis. Because the Ras oncogene promotes tumors in cell cycle-
interneurons, reaching maximal levels at development, NPY mRNA is transiently increased in cortical trophic factor stimulates NPY synthesis, and, during de-
mimic neurotrophic actions. Similarly, brain-derived neu-
igra (Hagg, 1998), are increased in synRas-TG mice (Gnahn et al., 1983) and the TH activity of the substantia
same developmental profile is displayed by organotypic thalamocortical cocultures. An early “postnatal” applica-
tion of NT4/5 prevents the developmental downregulation (Wahle et al., 2000). Moreover, blocking spontaneous electrical activity also downregulates NPY mRNA (Wirth et al., 1998). Thus, Ras could mimic neuronal activity, resulting in a maintenance of elevated NPY mRNA levels in the neocortex of adult synRas-TG mice. In PC12 cells, such an increase in TrkB-mediated NPY protein expression is sensitive to blockade by the MEK-inhibitor PD98059, suggesting a role of the MAPK signaling pathway (Williams et al., 1998).

MAPKs (ERKs) are phosphorylated after direct binding of Raf kinases to activated Ras. In addition, some subtypes of PI 3-kinase have been shown to be direct effectors of GTP-bound Ras (Rodriguez-Viciana et al., 1996; Rubio et al., 1997), and PI 3-kinase is a preferred effector of the Ha-Ras isotype used in the synRas-TG mice (Yan et al., 1998). Activation of the PI 3-kinase/Akt-1 (PKB) pathway has been shown to phosphorylate and, thereby, inactive cell death regulatory proteins (Biggs et al., 1999; Brunet et al., 1999). However, in contrast to our recent results on the rapid increase of PI 3-kinase in response to NGF activation in peripheral neurons (Virdee et al., 1999), constitutive activa-
tion of Ras did not reveal any changes in PI 3-kinase activity. The levels of the phosphorylation of the down-
stream kinase PKB/Akt remained equally unaffected in synRas-TG mice. Thus, the phenotype of synRas-TG mice appears to be independent of a major PI 3-kinase activation in cerebellum, hippocampus, and cortex. Interestingly, PI 3-kinase–independent neuronal survival pathways have been described recently, suggesting that PI 3-kinase activation may not be the sole mechanism of preventing neuronal cell death (Vogelbaum et al., 1998; Virdee et al., 1999; Williams and Doherty, 1999).

One of the major motivations to establish this synRas-
TG mouse model was to investigate the possible role of Ras activity in lesion-induced degeneration of the adult nervous system. It has been shown previously that in cul-

Figure 5. Characterization of Ras-transgenic facial motorneurons. (A) Ras-TG expression in motorneurons of the facial nucleus in adult mice (6 mo old). (a and b) Low-power photomicrographs of coronal sections through the brainstem of a synRas-TG animal (a) and a wt littermate (b). At the level of the genu of the facial nerve, hybridization with lacZ mRNA reveals intense signals in N7 motor-neurons and in subsets of neurons in other brainstem structures in the synRas-TG, but not the wt mouse. (c and d) Higher magnification of the facial nucleus at a more posterior level in the synRas-TG animal reveals intensely lacZ mRNA–expressing neurons. The Ras-TG expression varies between the individual neu-
rons, and the reaction product may extend into primary dendrites (arrow head). No signals were observed in glia cells. Bars: (a and b) 200 μm; (c) 50 μm; (d) 20 μm. (B) Size-frequency distribution of facial motorneurons. The average facial motorneuron size is enlarged in synRas-TG mice. Insert: Chat activity in facial nuclei of synRas-TG mice and wt littermates. Expression of Ras-TG protein in facial motorneurons results in a significant increase in Chat activity.

Our finding that activities of neurotransmitter-synthe-
sizing enzymes regulated by neurotrophic factors, such as Chat activity of the sepal or nucleus facialis regions (Gnahn et al., 1983) and the TH activity of the substantia nigra (Hagg, 1998), are increased in synRas-TG mice fits with the notion that Ras activation in neurons could mimic neurotrophic actions. Similarly, brain-derived neu-
rotrophic factor stimulates NPY synthesis, and, during de-
velopment, NPY mRNA is transiently increased in cortical interneurons, reaching maximal levels at ~30 d postnatally and is then downregulated (Obst and Wahle, 1995). The same developmental profile is displayed by organotypic thalamocortical cocultures. An early “postnatal” application of NT4/5 prevents the developmental downregulation (Wahle et al., 2000). Moreover, blocking spontaneous electrical activity also downregulates NPY mRNA (Wirth et al., 1998). Thus, Ras could mimic neuronal activity, resulting in a maintenance of elevated NPY mRNA levels in the neocortex of adult synRas-TG mice. In PC12 cells, such an increase in TrkB-mediated NPY protein expression is sensitive to blockade by the MEK-inhibitor PD98059, suggesting a role of the MAPK signaling pathway (Williams et al., 1998).

MAPKs (ERKs) are phosphorylated after direct binding of Raf kinases to activated Ras. In addition, some subtypes of PI 3-kinase have been shown to be direct effectors of GTP-bound Ras (Rodriguez-Viciana et al., 1996; Rubio et al., 1997), and PI 3-kinase is a preferred effector of the Ha-Ras isotype used in the synRas-TG mice (Yan et al., 1998). Activation of the PI 3-kinase/Akt-1 (PKB) pathway has been shown to phosphorylate and, thereby, inactive cell death regulatory proteins (Biggs et al., 1999; Brunet et al., 1999). However, in contrast to our recent results on the rapid increase of PI 3-kinase in response to NGF activation in peripheral neurons (Virdee et al., 1999), constitutive activa-
tion of Ras did not reveal any changes in PI 3-kinase activity. The levels of the phosphorylation of the down-
stream kinase PKB/Akt remained equally unaffected in synRas-TG mice. Thus, the phenotype of synRas-TG mice appears to be independent of a major PI 3-kinase activation in cerebellum, hippocampus, and cortex. Interestingly, PI 3-kinase–independent neuronal survival pathways have been described recently, suggesting that PI 3-kinase activation may not be the sole mechanism of preventing neuronal cell death (Vogelbaum et al., 1998; Virdee et al., 1999; Williams and Doherty, 1999).

One of the major motivations to establish this synRas-
TG mouse model was to investigate the possible role of Ras activity in lesion-induced degeneration of the adult nervous system. It has been shown previously that in cul-
tures of embryonic neurons ligand-independent survival is achieved by intracellular application of activated Ras in chick dorsal root ganglia neurons, in superior cervical ganglion neurons, or in spinal motorneurons (Borasio et al., 1989; Nobes et al., 1996; Weng et al., 1996; Mazzoni et al., 1999). Our finding that constitutively activated Ras in neurons completely prevented axotomy-induced degeneration of adult facial motorneurons shows that the activation of the Ras pathway promotes protective functions in vivo.

Survival of motorneurons is also enhanced by expression of Bcl-2 in transgenic mice and in Bax knock out mice because axotomy-induced motorneuron death was prevented (Martinou et al., 1994; Deckwerth et al., 1996). In our system, considering the lack of changes in levels of Bcl-2 or Bcl-XL, together with the lack of activation of PKB/Akt, we assume that an as yet unidentified downstream target of Ras, other than the Bcl-2 family of proteins, may be involved in the Ras-mediated neuronal rescue.

In addition to motorneurons, dopaminergic neurons of the substantia nigra were investigated, because they serve as a model of neurons affected in Parkinson’s disease. 6-OHDA and MPP+ were used as toxins that are thought to be specific for neurons carrying the dopamine transporter. Although the mechanism by which MPP+ causes neuronal degeneration appears to be distinctly different (Lotharius et al., 1999), the adverse effects of both types of toxins could be attenuated in dopaminergic neurons of synRas-TG mice. Interestingly, suppression of Ras activity in PC12 cells rendered them more sensitive to toxic radicals (Guyton et al., 1996), and, similarly, in rat sympathetic neurons, NGF-induced suppression of radical formation was dependent on the concommittent activation of MAPK (Dugan et al., 1997).
were derived from E14 embryos and cultured under serum-free conditions. After treatment with MPP+ number of cultured midbrain dopaminergic neurons derived from synRas-TG mice or their wt siblings. Midbrain dopaminergic neurons could be tested for a possible enhancement of en neuronal-specific NF splice variant (Geist and Gutmann, 1997; Piccini et al., 1999), and methods for neuronal stabili- sation before transplantation are desirable. In addition, pharmacological downregulation of a recently described Ras-TG transcript is found in the TH-positive neuron. (B) Effect of 6-OHDA or MPP+ treatment on the number of cultured midbrain dopaminergic neurons derived from synRas-TG mice or their wt siblings. Midbrain dopaminergic neurons were derived from E14 embryos and cultured under serum-free conditions. After treatment with MPP+ (1 µM) for 36 h or 6-OHDA (150 µM) for 90 min, cells were fixed with methanol. Dopaminergic neurons were identified using a monoclonal antibody against TH (Roche), a biotin-labeled secondary antibody (Sigma-Aldrich), and a streptavidine–FITC conjugate. After counting dopaminergic neu- rons, the genotype was determined by Southern blot analysis. Treatment with the toxin resulted in degeneration of the dopaminergic neurons. This toxin-induced degeneration was markedly reduced in cultures derived from synRas-TG mice. The numbers of non- dopaminergic neurons remained unchanged (data not shown). (C) Determination of dopamine after in vivo treatments with MPTP in synRas-TG mice and their wt siblings (see Materials and Methods). In the synRas-TG mice, the MPTP-induced decrease of dopamine contents was profoundly reduced.

Figure 7. Characterization of transgenic midbrain dopaminergic neurons and protection against neurotoxin-induced degeneration. (A) Expression of the Ras-TG in dopaminergic neurons of the substantia nigra. (I) A coronal section through the midbrain at the level of the substantia nigra was hybridized with a lacZ riboprobe and showed staining in a subset of neurons. (II) Immunohistochemical analysis of the same section using a monoclonal antibody against TH (Roche) and a Texas red–labeled secondary antibody shows TH expres- sion in the neuron expressing the transgene. Bar, 20 μm. (III) Overlay of the immunofluorescence picture and the brightfield micro- graph, indicating that the Ras-TG transcript is found in the TH-positive neuron. (B) Effect of 6-OHDA or MPP+ treatment on the number of cultured midbrain dopaminergic neurons derived from synRas-TG mice or their wt siblings. Midbrain dopaminergic neurons were derived from E14 embryos and cultured under serum-free conditions. After treatment with MPP+ (1 µM) for 36 h or 6-OHDA (150 µM) for 90 min, cells were fixed with methanol. Dopaminergic neurons were identified using a monoclonal antibody against TH (Roche), a biotin-labeled secondary antibody (Sigma-Aldrich), and a streptavidine–FITC conjugate. After counting dopaminergic neu- rons, the genotype was determined by Southern blot analysis. Treatment with the toxin resulted in degeneration of the dopaminergic neurons. This toxin-induced degeneration was markedly reduced in cultures derived from synRas-TG mice. The numbers of non- dopaminergic neurons remained unchanged (data not shown). (C) Determination of dopamine after in vivo treatments with MPTP in synRas-TG mice and their wt siblings (see Materials and Methods). In the synRas-TG mice, the MPTP-induced decrease of dopamine contents was profoundly reduced.

In summary, we present a model of a neuroprotective mechanism promoted by constitutive activation of Ras in adult central nervous system neurons. It is now possible to test if neural stem cells derived from synRas-TG mice can be isolated, cultured, and transplanted to result in stabil- ized differentiated neurons due to the Ras-TG expression in the adult mouse brain. The success of transplants of dopaminergic embryonic mesencephalic neurons into Parkinson’s patients has been shown to depend on the preser- vation of TH activity of the transplant (Wenning et al., 1993; Piccini et al., 1999), and methods for neuronal stabiliza- tion before transplantation are desirable. In addition, pharmacological downregulation of a recently described neuronal-specific NF splice variant (Geist and Gutmann, 1996) could be tested for a possible enhancement of endo- genous Ras activity. Thus, investigations on the Ras-medi- ated neuroprotection in vivo may help to develop new therapeutic strategies against neurodegenerative diseases.

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