Alternative sulfonylurea receptor expression defines metabolic sensitivity of K-ATP channels in dopaminergic midbrain neurons

Birgit Liss, Ralf Bruns and Jochen Roeper

Institute for Neural Signal Transduction, Centre for Molecular Neurobiology, Martinistraße 52, 20246 Hamburg, Germany

1 Present address: MRC Anatomical Neuropharmacology Unit, Oxford University, Mansfield Road, Oxford OX1 3TH, UK

2 Corresponding author
e-mail: roeper@plexus.uke.uni-hamburg.de

R.Bruns contributed experiments shown in Figure 6C and D

ATP-sensitive potassium (K-ATP) channels couple the metabolic state to cellular excitability in various tissues. Several isoforms of the K-ATP channel subunits, the sulfonylurea receptor (SUR) and inwardly rectifying K channel (Kir6.X), have been cloned, but the molecular composition and functional diversity of native neuronal K-ATP channels remain unresolved. We combined functional analysis of K-ATP channels with expression profiling of K-ATP subunits at the level of single substantia nigra (SN) neurons in mouse brain slices using an RT–multiplex PCR protocol. In contrast to GABAergic neurons, single dopaminergic SN neurons displayed alternative co-expression of either SUR1, SUR2B or both SUR isoforms with Kir6.2. Dopaminergic SN neurons expressed alternative K-ATP channel species distinguished by significant differences in sulfonylurea affinity and metabolic sensitivity. In single dopaminergic SN neurons, co-expression of SUR1 + Kir6.2, but not of SUR2B + Kir6.2, correlated with functional K-ATP channels highly sensitive to metabolic inhibition. In contrast to wild-type, surviving dopaminergic SN neurons of homozygous weaver mouse exclusively expressed SUR1 + Kir6.2 during the active period of dopaminergic neurodegeneration. Therefore, alternative expression of K-ATP channel subunits defines the differential response to metabolic stress and constitutes a novel candidate mechanism for the differential vulnerability of dopaminergic neurons in response to respiratory chain dysfunction in Parkinson’s disease.

Keywords: ATP-sensitive potassium channel/Parkinson’s disease/single-cell RT–PCR/substantia nigra/weaver

Introduction

Chronic metabolic stress is believed to play an important role in neurodegenerative disorders such as Parkinson’s or Huntington’s disease (Beal, 1996; Hanna and Bhatia, 1997). Different neuronal populations are not affected equally by metabolic challenges (Schreiber and Baudry, 1995), and resistant subpopulations within degenerating nuclei have been identified (Hirsch et al., 1997). Therefore, it is of considerable interest to define the molecular basis of differential neuronal vulnerability. Neuronal ATP-sensitive potassium (K-ATP) channels might be one of the promising targets acting as direct functional response elements to metabolic stress (Roeper and Ashcroft, 1995; Trapp and Ballanyi, 1995; Fujimura et al., 1997). K-ATP channels couple membrane excitability to cellular metabolism by directly sensing and integrating intracellular concentration changes of nucleotides such as ATP and ADP (Ashcroft, 1988; Quale et al., 1997; Aguilar-Bryan et al., 1998; Babenko et al., 1998; Yokoshiki et al., 1998).

Sulfonylurea binding and electrophysiological studies have characterized neuronal K-ATP channels with different properties in a variety of cell types, including hippocampal CA1 neurons, dorsal vagal neurons and hypothalamic neurons (Trapp and Ballanyi, 1995; Dunn-Meynell et al., 1997; Fujimura et al., 1997; Spanswick et al., 1997). We and others have identified functional somatodendritic K-ATP channels on dopaminergic substantia nigra (SN) pars compacta neurons. These neuronal K-ATP channels were activated by different paradigms of metabolic stress such as hypoxia, sodium loading, glucose removal or dialysis with ATP-free pipette solutions (Mercuri et al., 1994; Roeper and Ashcroft, 1995; Watts et al., 1995; Seutin et al., 1996; Stanford and Lacey, 1995). We have shown previously that inhibition of mitochondrial complex I by rotenone leads to a tonic activation of K-ATP channels, complete cessation of spontaneous electrical activity and membrane hyperpolarization in dopaminergic SN neurons (Roeper and Ashcroft, 1995). This is of particular interest, as complex I inhibition is one of the proposed trigger mechanisms of degeneration of dopaminergic midbrain neurons in Parkinson’s disease (Beal, 1996; Hanna and Bhatia, 1997; Hirsch et al., 1997).

The cloning of K-ATP channel subunits (Aguilar-Bryan et al., 1995; Sakura et al., 1995; Chutkow et al., 1996; Inagaki et al., 1995a, 1996) allowed us to probe for the molecular identity of neuronal K-ATP channels in SN neurons. K-ATP channels are heterooligomeric proteins composed of two types of subunits, a sulfonylurea receptor (SUR) and an inwardly rectifying K channel subunit (Kir6.X), which assemble in a 4:4 stoichiometry (Clement et al., 1997). Two related isoforms of the SUR have been identified, SUR1 and SUR2; the latter exists in at least two alternatively spliced forms (SUR2A and SUR2B) (Chutkow et al., 1996; Inagaki et al., 1996; Isomoto et al., 1996; Yamada et al., 1997). In addition, two related Kir6.X channels have been identified, Kir6.1 and Kir6.2 (Inagaki et al., 1995b; Sakura et al., 1995). Co-expression experiments of different SUR and Kir6 isoforms in heterologous systems suggested that different functional properties and pharmacological profiles of recombinant K-ATP channels are mediated by alternative subunit combinations (Inagaki et al., 1996; Isomoto et al., 1996; Gribble et al., 1997; Yamada et al., 1997; Okuyama et al., 1998).
The aim of this study was to identify the molecular composition and functional diversity of native K-ATP channels in midbrain neurons at the level of single cells. In addition to wild-type neurons, we also analysed K-ATP channel expression profiles of dopaminergic SN neurons in the weaver mouse, a genetic model of selective degeneration of a subpopulation of dopaminergic SN neurons induced by a pore mutation in a G-protein-activated inwardly rectifying K channel (Girk2; Graybiel et al., 1990; Bayer et al., 1995; Patil et al., 1995). We combined electrophysiological analysis with single-cell RT–multiplex PCR (RT–mPCR) protocols (Jonas et al., 1994; Monyer and Lambolez, 1995; Audinat et al., 1996; O’ Dowd and Smith, 1996; Surmeier et al., 1996; Cauli et al., 1997). In contrast to γ-aminobutyric acid (GABA)ergic SN neurons, single wild-type dopaminergic SN neurons displayed alternative expression profiles of K-ATP channel subunits (SUR1 + Kir6.2 and SUR2B + Kir6.2) and functional K-ATP channels with different pharmacological and metabolic properties. The co-expression of SUR1 + Kir6.2, but not of SUR2B + Kir6.2, was correlated with functional K-ATP channels highly sensitive to complex I inhibition. In addition, surviving dopaminergic SN neurons in homozygous weaver mice exclusively expressed SUR1 + Kir6.2.

**Results**

**RT–multiplex PCR**

An RT–mPCR protocol (Cauli et al., 1997) was designed simultaneously to detect, at the single-cell level, the mRNAs of the K-ATP channel subunits SUR1, SUR2, Kir6.2 and Kir6.1, as well as the mRNAs for the Girk2 subunit of the G-protein-activated inwardly rectifying K channel (Lesage et al., 1994), and the marker genes tyrosine hydroxylase (TH) (Iwata et al., 1992), glutamate decarboxylase (GAD67) (Szabó et al., 1996) and glial fibrillary acidic protein (GFAP) (Brenner, 1994). The expression profiles of the marker genes were used to classify analysed SN cells as dopaminergic (TH+) and GABAergic (GAD67+) neurons, and to exclude astroglial cells (GFAP+). Girk2, which is known to be expressed in dopaminergic SN neurons, was included to control the ability to detect low abundance ion channel transcripts in single neurons. For this technique, two rounds of PCR amplification, an initial mPCR and a subsequent nested PCR, were used after reverse transcription. The sensitivity of the protocol was optimized using highly diluted midbrain cDNA to allow reliable simultaneous amplification of all eight transcripts from estimated subfemptomolar transcript concentrations, which is likely to be in the range found in single cells. Figure 1A shows that the eight specific mRNAs were detected, each generating a PCR fragment of the size predicted by its mRNA sequence. The identities of all PCR fragments were verified by subcloning and sequencing or, alternatively, by direct sequencing. Alternative primer pairs were used to differentiate between the two SUR2 isoforms, SUR2A and SUR2B. In contrast to heart, where both splice variants SUR2A and SUR2B were expressed, only SUR2B is present in ventral midbrain (Figure 1B). Similar results were obtained from single-cell RT–mPCR experiments (see below), indicating that the SUR2A subunit which is believed to be one of the components of the cardiac K-ATP channel (Inagaki et al., 1996) is not expressed in SN neurons.

The established single-cell RT–mPCR protocol was then used to analyse the K-ATP channel subunit expression profiles in identified SN neurons. Three different neuronal populations were analysed. Our main interest was focused on the dopaminergic neurons located predominantly in the SN pars compacta (SNpc) (Condé, 1992). For comparison, we also studied a population of small interneurons from the SNpc (Lacey et al., 1989) and GABAergic projection neurons in the SNpars reticulata (SNpr) (Condé, 1992). Recent electrophysiological studies have suggested that two of these neuronal populations, the dopaminergic SNpc (Condé, 1992; Roep and Ashcroft, 1995; Watts et al., 1995; Seutin et al., 1996; Stanford and Lacey, 1995) and the GABAergic SNpr neurons (Schwanstecher and Panten, 1993; Stanford and Lacey, 1996), possess functional K-ATP channels, whereas the presence of K-ATP channels in SNpc interneurons has not yet been investigated. The different neuronal populations could be identified easily in acute coronal midbrain slices by their morphology and localization within the SN subnuclei. After electrophysiological characterization in the whole-cell patch-clamp mode, the cytoplasm of the cells was harvested carefully under visual control and stable seal conditions for single-cell RT–mPCR. In most cases, the nucleus was also harvested, and the patch pipette repositioned a few times to maximize the cellular yield. To rule out the possibility of genomic contamination by DNA of the single nucleus acting as a potential template for PCR amplification, we performed control single-cell mPCR amplifications without prior reverse transcription, obtaining no detectable PCR products (n = 10; Figure 1C). Also, we harvested the cellular contents carefully excluding the nucleus.
(n = 5). After reverse transcription and PCR, we could detect all the cDNAs routinely detected in cells which were harvested, including the nucleus. These control experiments clearly indicate that under our conditions, the genomic DNA of the single nucleus was not acting as a possible template for PCR amplification and that, consequently, we were studying the genuine mRNA expression profiles of single SN neurons.

**Electrophysiological properties of substantia nigra neurons with a defined marker expression profile**

As expected, the large (30–80 pF) neurons located in the SNpc demonstrated TH, but not GAD67 nor GFAP expression (n = 77), indicating their dopaminergic phenotype. At the single-cell level, the TH\(^+\) expression profile was correlated consistently with the well-characterized electrophysiological phenotype of dopaminergic SN neurons (Grace and Onn, 1989; Lacey et al., 1989; Richards et al., 1997) (Figure 2A), i.e. low-frequency pacemaker activity (1.8 ± 0.1 Hz, n = 70), broad action potentials (8.0 ± 0.5 ms, n = 40), with depolarized thresholds (–36.3 ± 0.8 mV, n = 40) and the presence of a strong \(h\)\(^-\)-dependent sag-current component. In addition, our voltage-clamp analysis of SN neurons showed that a dominant A-type current was found in all cases in TH\(^+\) SN neurons (n = 77), but was not observed in TH\(^-\), GAD67\(^+\), i.e. GABAergic SN neurons (n = 22). The small interneurons of the SNpc (10–25 pF) showed a marker expression profile (TH\(^+\), GAD67\(^+\), GFAP\(^-\); n = 11) indicating their previously only presumed GABAergic phenotype (Lacey et al., 1989; Johnson and North, 1992). These neurons were either electrically silent or displayed spontaneous discharge with higher frequencies compared with TH\(^+\) SNpc neurons (8.8 ± 0.6 Hz, n = 10; Figure 2B). In voltage-clamp experiments, they possessed no fast-inactivating outward component in the subthreshold range and showed slowly activating outward currents at higher membrane potentials (Figure 2B). The medium sized (20–40 pF) neurons of the SNpr possessed a marker expression profile (TH\(^-\), GAD67\(^+\), GFAP\(^-\); n = 11) identifying them as GABAergic. In accordance with previous reports (Richards et al., 1997), these SNpr neurons showed either high frequency discharge (11.4 ± 0.7 Hz, n = 10) or were electrically silent (Figure 2C). In voltage-clamp, these neurons also lacked a subthreshold A-type current but showed a small, fast-inactivating outward current component at more depolarized potentials (Figure 2C). Girk2 transcripts, which code for one subunit of the G-protein-activated Kir channels, were detected in all dopaminergic (n = 77) and GABAergic SNpc neurons (n = 11), but only in seven of 11 GABAergic SNpr neurons.

**Expression of K-ATP channel subunits in single GABAergic SN neurons**

Electrophysiological studies reported the presence of functional K-ATP channels on SNpr neurons of unknown
We found no functional evidence that these GABAergic (TH\(^{--}\), GAD67\(^{--}\)) neurons demonstrated a homogeneous pattern of K-ATP channel subunits (Figure 3B). These cells seemed, therefore, devoid of classical K-ATP channels acting as metabolic sensors. These results show that the K-ATP channel subunit expression profiles are different for two populations of GABAergic SN neurons but homogeneous within each population.

### Alternative expression profiles of K-ATP channel subunits in dopaminergic SN neurons

The single-cell RT–mPCR analysis of K-ATP channel subunit expression in phenotypically defined dopaminergic SN neurons (TH\(^{++}\), GAD67\(^{++}\), GFAP\(^{--}\)) revealed a heterogeneous expression pattern for the transcripts mediating the sulfonylurea receptors, SUR1 and SUR2. Three different expression pattern could be identified in the overall population of analysed dopaminergic SN neurons (Figure 4; \(n = 54\)). One subpopulation showed co-expression of SUR1 with Kir6.2, which represented 39\% (\(n = 21\) of 54) of the total population (Figure 4A). In a second major subpopulation of dopaminergic SN neurons (37\%, \(n = 20\) of 54), we detected the co-expression of SUR2 and Kir6.2 (Figure 4B). Finally, a third population of dopaminergic SN neurons (24\%, \(n = 13\) of 54) demonstrated co-expression of both SUR1 and SUR2 with Kir6.2 (Figure 4C). In accordance with the midbrain RT–PCR results, the use of SUR2 isofrom-specific primer pairs showed that SUR2B but not SUR2A was expressed also at the single-cell level (\(n = 5\); Figure 4D). Kir6.1 expression was not detected in dopaminergic SN neurons. These results showed that at the level of mRNA transcripts, dopaminergic in contrast to GABAergic SN neurons displayed alternative expression profiles of K-ATP channel subunits. This raised two questions. First, do dopaminergic SN neurons in contrast to GABAergic SN neurons also display a heterogeneity at the level of functional somatodendritic K-ATP channels? Secondly, which combination(s) of K-ATP channel transcripts codes for the highly sensitive metabolic sensor in dopaminergic SN neurons and is therefore responsible for coupling metabolic stress to altered excitability?

### Dopaminergic SN neurons express K-ATP channel subtypes with different sulfonylurea sensitivities

Our single-cell RT–mPCR data of dopaminergic SN neurons demonstrated a heterogeneous expression profile of K-ATP channel subunits. The alternative mRNA expression pattern of SURs in dopaminergic SN neurons predicted differences in sulfonylurea sensitivity for the level of functional K-ATP channels. Heterologous studies showed that SUR1-mediated K-ATP channels possess a higher affinity for sulfonylureas like tolbutamide compared with those containing SUR2A or SUR2B (Ishimoto and Kurachi, 1997; Babenko et al., 1998; Gribble et al., 1998). To study their sulfonylurea sensitivities, neuronal K-ATP channels were activated by combining pre-incubation of the slice with 100 nM of the respiratory chain complex I blocker rotenone (>30 min) with dialysis of an ATP-free pipette solution. After 5 min of dialysis with ATP-free pipette solutions, stable current levels were obtained in all cells, although we observed differences in the washout kinetics of the K-ATP currents. Subsequently, several molecular composition (Schwanstecher and Panten, 1993; Stanford and Lacey, 1996). The phenotype of these cells was not characterized further, but they were believed to be GABAergic. These K-ATP channels were reported to be activated by metabolic inhibition and the K channel opener diazoxide, and blocked by sulfonylureas. Our single-cell RT–mPCR analysis showed that phenotypically identified GABAergic (TH\(^{++}\), GAD67\(^{++}\)) SNpc neurons presented a homogeneous pattern of K-ATP channel subunit expression (\(n = 11\)). In all analysed GABAergic SNpc neurons, we detected the co-expression of Kir6.2 and SUR1. SUR2 and Kir6.1 were not detected in this cell type (Figure 3C). SUR1 + Kir6.2 co-expression at the transcript level might, therefore, suggest that these subunits form the molecular basis of functional K-ATP channels in GABAergic SNpc neurons. No previous studies have addressed the question of the presence of K-ATP channels on small interneurons in the SNpc. We found no functional evidence that these GABAergic (TH\(^{++}\), GAD67\(^{++}\), GFAP\(^{--}\)) interneurons in the SNpc possessed K-ATP channels.
Expression of K-ATP channel subunits in midbrain neurons

Fig. 4. Alternative SUR expression profiles in single dopaminergic SNpc neurons. mPCR expression profiling of single dopaminergic (TH⁺, GAD67⁻) SNpc neurons defined three dopaminergic subpopulations (A–C) with heterogeneous K-ATP subunit expression profiles. TH and Girk2 were expressed in a homogeneous fashion in all dopaminergic neurons. (A) Agarose gel analysis of a dopaminergic SNpc neuron of the first subpopulation where four PCR products were detected corresponding to the K-ATP channel subunits SUR1 (401 bp) and Kir6.2 (448 bp), and to the marker transcripts TH (377 bp) and Girk2 (595 bp); 21 of 54 tested neurons showed this expression profile. (B) Agarose gel analysis of a dopaminergic SNpc neuron of the second subpopulation where four PCR products were detected corresponding to the K-ATP channel subunits SUR2 (215 bp) and Kir6.2 (448 bp), and to the marker transcripts TH (377 bp) and Girk2 (595 bp); 20 of 54 tested neurons showed this expression profile. (C) Agarose gel analysis of a dopaminergic SNpc neuron of the third subpopulation where five PCR products were detected corresponding to the K-ATP channel subunits SUR1 (401 bp), SUR2 (215 bp) and Kir6.2 (448 bp), and to the marker transcripts TH (377 bp) and Girk2 (595 bp); 13 of 54 tested neurons showed this expression profile. (D) SUR2B is expressed in dopaminergic SNpc neurons as revealed by the use of SUR2 splice variant-specific primers in the single-cell RT–mPCR. Note that in this and four other dopaminergic neurons, only SUR2B (337 bp) but not SUR2A were detected (compare with Figure 1B).

Concentrations (0.1 µM–1 mM) of the sulfonylurea tolbutamide were tested by local application via a buffer pipette. Consistent with their homogeneous SUR1 + Kir6.2 mRNA expression pattern, all GABAergic SNpr neurons displayed K-ATP channels with high sulfonylurea sensitivity. As shown in Figure 5A, the concentration dependence of tolbutamide inhibition of the whole-cell K-ATP current in GABAergic SNpr neurons was described by a Hill equation with an IC₅₀ value of 8.7 ± 1.7 µM and an associated Hill coefficient of 1.1 ± 0.1 (n = 8). In contrast,
dopaminergic SN neurons \((n = 23)\) displayed different sensitivities to tolbutamide inhibition (Figure 5B). A total of 40% (nine of 23) of the cells possessed a high affinity for tolbutamide which was described by a Hill equation with an \(IC_{50}\) value of 8.8 ± 1.0 \(\mu\)M and a Hill coefficient of 1.2 ± 0.1. K-ATP channels in a second group \((n = 6)\) of dopaminergic SN neurons showed a 7-fold lower sulfonylurea sensitivity \((IC_{50} = 60.6 ± 5.1 \mu\)M; Hill coefficient = 1.0 ± 0.1, \(n = 6)\). In a third population of dopaminergic SN neurons \((n = 8)\), functional K-ATP channels were activated which possessed an intermediate sulfonylurea sensitivity, with an \(IC_{50}\) of 27.9 ± 1.9 \(\mu\)M and a Hill coefficient of 1.1 ± 0.1. For these cells, the concentration dependence of tolbutamide inhibition was not well described by the sum of two Hill equations with a high and low \(IC_{50}\) value but rather with a single Hill equation with an intermediate sulfonylurea sensitivity. The tolbutamide sensitivities of the three populations of dopaminergic SN neurons were significantly different \((p > 0.0005)\) and those with the highest sensitivity were similar to that of the GABAergic SNpr neurons (Figure 5C). In accordance with our single-cell RT–mPCR data, these pharmacological results demonstrated that GABAergic SNpr neurons possess a homogeneous population of K-ATP channels with a high SUR1-like sulfonylurea sensitivity. In contrast, dopaminergic SN neurons functionally express multiple K-ATP channel species distinguished by different affinities for sulfonylureas.

**K-ATP channel subtypes in dopaminergic SN neurons display large differences in metabolic sensitivities**

The results presented so far indicate that dopaminergic SN neurons express molecularly and pharmacologically distinct K-ATP channels. Our main interest was to ask whether these K-ATP channel types also possess functionally relevant differences. One of the most important differences between alternative K-ATP channel types would concern their metabolic sensitivity. Major differences in the metabolic sensitivity of K-ATP channels in dopaminergic SN neurons were apparent from two sets of experiments: either when the slices were pre-incubated with nanomolar concentrations of the complex I inhibitor rotenone or when 10 \(\mu\)M rotenone was applied locally to a patch-clamped neuron. In acute experiments, application of 10 \(\mu\)M rotenone led to activation of a K conductance which was voltage and time independent and showed little to no rectification in the subthreshold voltage range between −120 and −40 mV in a subpopulation of dopaminergic SN neurons. Current-clamp recordings showed that rotenone induced a mean hyperpolarization to −60.9 ± 1.9 mV from resting potentials in the range of −40 to −45 mV \((n = 14)\). The hyperpolarization was accompanied by a complete cessation of spontaneous electrical activity and was inhibited by 100 \(\mu\)M tolbutamide (Figure 6A). Co-application of glibenclamide (100 \(\mu\)M, \(n = 38\); Figure 6C) or tolbutamide (100 \(\mu\)M, \(n = 5\); Figure 6B) completely blocked the induced K conductance, and the current amplitudes returned to control levels. Whereas the block by tolbutamide was readily reversible, inhibition by glibenclamide was irreversible even after prolonged periods (>40 min) of washout. These experiments demonstrated that complex I inhibition selectively activated K-ATP channels with a sulfonylurea sensitivity which corresponded to the most sensitive subtype defined by the pharmacological experiments (Figure 5B). Voltage ramps showed that the K-ATP whole-cell conductances induced by complex I inhibition \((2.2 ± 0.2 \, nS, n = 12)\) reversed to \(E_K\) (104.0 ± 1.1 mV, \(n = 12\); Figure 6B). We also identified a subpopulation \((37\%, n = 25\) of 68) of dopaminergic SN neurons which showed no detectable change in membrane conductance during acute application of 10 \(\mu\)M rotenone (Figure 6D). Application of 10 \(\mu\)M rotenone to neurons which were dialysed with a pipette solution containing 2 mM ATP in the standard whole-cell configuration was not sufficient to activate the other K-ATP channel species with lower sulfonylurea sensitivities. It is likely that the standard whole-cell configuration per se will alter the metabolic threshold for K-ATP channel activation by washout of activating metabolites and the coupling of the cell to a virtually unlimited pool of ATP via the patch pipette.

To overcome these limitations, we investigated the threshold for K-ATP channel activation by complex I inhibition in metabolically intact cells by pre-incubation of the slice for >30 min in rotenone concentrations ranging from 1 \(nM\) to 10 \(\mu\)M. A 10 \(\mu\)M concentration of rotenone was the upper limit because pre-incubation with higher concentrations induced a fast disintegration of the midbrain slice. After the pre-incubation period, K-ATP currents and membrane potential of dopaminergic SN neurons were analysed immediately after obtaining the standard whole-cell configuration to minimize secondary washout effects. This second type of metabolic experiment confirmed that dopaminergic SN neurons indeed possess different thresholds for K-ATP channel activation. In one group of neurons, K-ATP channel activation and membrane hyperpolarization accompanied by loss of spontaneous activity were already observed in the presence of low nanomolar concentrations of rotenone. In 100 \(nM\) rotenone, these neurons showed large K-ATP currents and membrane hyperpolarizations greater than −65 mV (Figure 7A, \(n = 8\)). In addition, K-ATP channels were fully activated because dialysis with ATP-free pipette solutions did not enhance further the outward currents that were observed immediately after establishing the whole-cell configuration (Figure 7A). The mean rotenone concentration dependence of membrane hyperpolarization for this dopaminergic subpopulation was described by a Hill equation with an \(EC_{50}\) of 14.5 \(nM\), a Hill coefficient of 0.78 and a maximal membrane hyperpolarization to −68.1 mV (Figure 7B, \(n = 6–10\)). The mean rotenone concentration dependence of activation of this K-ATP channel subpopulation was described by a Hill equation with an \(EC_{50}\) of 16.2 nM, a Hill coefficient of 0.94 and a maximal K-ATP conductance of 7.1 nS (Figure 7C, \(n = 6–10\)). Co-pre-incubation of the slice with 100 \(nM\) rotenone and 100 \(\mu\)M tolbutamide completely prevented the activation of K-ATP channels in all dopaminergic SN neurons (subthreshold slope conductance: control: 1.5 ± 0.1 \(nS\), \(n = 12\); 100 \(nM\) rotenone: 7.7 ± 0.9 \(nS\), \(n = 8\); 100 \(nM\) rotenone + 100 \(\mu\)M tolbutamide: 1.6 ± 0.1 \(nS\), \(n = 10\)).

A second population of dopaminergic SN neurons was not affected by pre-incubation of nanomolar concentrations of rotenone, and K-ATP channels were only partially
Expression of K-ATP channel subunits in midbrain neurons

Fig. 6. Activation of K-ATP channels with high sulfonylurea affinity in a subpopulation of dopaminergic SN neurons by local application of 10 µM rotenone. (A) Current-clamp recording of a dopaminergic SN neuron. Application of 10 µM rotenone (indicated by the bar) led to hyperpolarization and complete cessation of spontaneous pacemaker activity which was completely reversed by the co-application of 100 µM tolbutamide. (B–D) Current responses evoked by 10 mV depolarizing voltage steps from a holding potential of –60 mV of voltage-clamped dopaminergic SNpc neurons (drug applications indicated by bars). (B) Application of 10 µM rotenone activated outward K currents which were completely and reversibly blocked by co-application of 100 µM tolbutamide. (C) Rotenone-activated K currents were completely blocked by 100 nM glibenclamide (insert: current responses to voltage ramps from –120 to –40 mV before and during application of rotenone showed that the rotenone-evoked current reversed close to EK). Forty-three of 68 dopaminergic SN neurons showed K-ATP currents activated by acute application of rotenone. (D) A subpopulation of dopaminergic SN neurons did not respond to local rotenone application. This and another 24 of 68 tested dopaminergic SNpc neurons showed no current activation by acute application of 10 µM rotenone (insert: current responses to voltage ramps from –120 to –40 mV before and during application of rotenone).

activated by rotenone concentrations of 1–10 µM. In 100 nM rotenone, these neurons presented subthreshold slope conductances and pacemaker activity similar to those observed under metabolic control conditions (Figure 7D). K-ATP channels were closed in the presence of 100 nM rotenone and only activated by dialysis with ATP-free pipette solutions (Figure 7D). The mean rotenone concentration dependence of membrane hyperpolarization for this dopaminergic subpopulation was described by a Hill equation with an EC50 of 1.2 µM, a Hill coefficient of 1 and a maximal membrane hyperpolarization to –58.3 mV (Figure 7E, n = 6–10). The mean rotenone concentration dependence of K-ATP channel activation in this population was described by a Hill equation with an EC50 of 3.6 µM (Hill coefficient = 1.0, maximum K-ATP conductance = 3.9 nS), corresponding to >200-fold lower metabolic sensitivity compared with the K-ATP channels activated in the nanomolar range (Figure 7F). Co-pre-incubation of 10 µM rotenone with 100 µM tolbutamide only partially prevented the activation of K-ATP channels in this subpopulation (subthreshold slope conductance: 10 µM rotenone: 4.5 ± 0.2 nS, n =
8; 10 µM rotenone + 100 µM tolbutamide: 2.8 ± 0.2 nS, n = 8). The partial block of K-ATP channels with low metabolic sensitivity by 100 µM tolbutamide indicated that they correspond to those with low sulfonylurea affinity (IC₅₀ tolbutamide = 60 µM; Figure 5B).

Like complex I inhibition, pre-incubation of the slice for >30 min in iodoacetate, an inhibitor of glycolysis in concentrations ranging from 1 to 300 µM, also activated two types of K-ATP channels with high (EC₅₀ = 17 µM, Hill coefficient = 2.5, maximum K-ATP conductance = 5.6 nS, n = 4–6) and low (EC₅₀ = 1.2 mM, Hill coefficient = 1, maximum K-ATP conductance = 5.5 nS, n = 4–6) metabolic sensitivity. Again, only K-ATP channels with high affinity for iodoacetate were blocked completely by co-pre-incubation with 100 µM tolbutamide (subthreshold slope conductance: control; 1.4 ± 0.1 nS, n = 8; 100 µM iodoacetate: 7.2 ± 0.6 nS, n = 6; 100 µM iodoacetate + 100 µM tolbutamide: 1.5 ± 0.2 nS, n = 7). In summary, these experiments clearly indicated that dopaminergic SN neurons possess at least two functionally different types of K-ATP channels with large differences in their sensitivities to metabolic inhibition (>200-fold for complex I inhibition; >70-fold for inhibition of glycolysis).

**Differential sensitivity to metabolic inhibition correlates with alternative SUR mRNA expression at the level of single dopaminergic SN neurons**

To test directly the hypothesis that alternative SUR expression is correlated with differential metabolic sensitivity, we combined rotenone pre-incubation with our single-cell RT–mPCR protocol. After pre-incubation with 100 nM rotenone, membrane potentials and subthreshold conductances were analysed immediately after obtaining the whole-cell configuration. Subsequently, the cytoplasm of the cells was harvested and used for single-cell RT–
mPCR. Dopaminergic SN neurons which showed the highest sensitivity for metabolic inhibition, evident by a large K-ATP whole-cell conductance and strong membrane hyperpolarization, expressed the K-ATP channel subunit combination SUR1 + Kir6.2 (Figure 8A). This is consistent with our finding that K-ATP channels with the highest metabolic sensitivity were also characterized by a high affinity for sulfonylureas. In contrast, for cells that were unresponsive to pre-incubation with 100 nM rotenone, co-expression of SUR2B + Kir6.2 was determined (Figure 8B). An intermediate response with a smaller hyperpolarization to about −55 mV and increases of subthreshold conductances to ~5 nS was found for cells co-expressing both sulfonylurea receptors SUR1 + SUR2B with Kir6.2 (Figure 8C). Under metabolic control conditions, no electrophysiological differences were apparent between neurons with alternative SUR expression pattern (subthreshold slope conductance: SUR1: 1.5 ± 0.1 nS, n = 12; SUR2B: 1.4 ± 0.2 nS, n = 7; SUR1 + SUR2B: 1.5 ± 0.3 nS, n = 4). The data for the 19 cells studied in this combined metabolic–molecular experiment are summarized in Figure 8D and E. They demonstrate that the alternative co-expression of sulfonylurea receptors in dopaminergic SN neurons is correlated with highly significant differences (SUR1 versus SUR2: p < 0.0005) in the functional response to inhibition of mitochondrial metabolism.

Surviving dopaminergic SN neurons in the weaver mouse exclusively express SUR1-mediated K-ATP channels

To test whether a particular SUR expression profile is correlated with the surviving population of dopaminergic SN neurons, we analysed their K-ATP channel expression profiles in 14-day-old homozygous weaver mice. The weaver mouse is a genetic model of selective degeneration of a subpopulation of highly vulnerable dopaminergic midbrain neurons caused by a pore mutation in a G-protein-activated inwardly rectifying K channel (Girk2) (Patil et al., 1995). All analysed dopaminergic SN neurons in homozygous weaver (w/vv) mouse exclusively expressed SUR1 + Kir 6.2 (Figure 9A; n = 22). In contrast, dopaminergic SN neurons of unaffected littermates (+/+) displayed a similar distribution of the
Expression profiling of K-ATP subunits in phenotypically identified single midbrain neurons by RT–mPCR

In the present study, we have determined the mRNA expression profiles of K-ATP channel subunits in three populations of wild-type and weaver mouse midbrain neurons at the level of single neurons using an RT–mPCR approach (n = 128). The sensitivity of the single-cell RT–mPCR protocol which combined two rounds of PCR (multiplex + nested) was optimized using highly diluted mouse midbrain cDNA to obtain an estimated subfemtomolar detection threshold suitable for single-cell transcripts. The neurons were characterized phenotypically by co- amplification of a panel of marker transcripts (TH, GAD67 and GFAP) in order to differentiate between dopaminergic and GABAergic neurons, and to exclude astroglial cells. Identified dopaminergic (TH⁺) SN neurons uniformly displayed their previously described typical electrophysiological properties, i.e. slow spontaneous pacemaker activity, broad action potentials with depolarized thresholds and a strong Iₚ component (Grace and Onn, 1989; Lacey et al., 1989; Conde, 1992; Richards et al., 1997). The small GAD67⁺ neurons within the SNpc most likely correspond to the previously reported SNpc type II neurons which displayed faster spontaneous activity and possessed no Iₚ or subthreshold A current (Lacey et al., 1989; Johnson and North, 1992). The larger GAD67⁺ neurons recorded in the SNpr showed similar properties to GABAergic projection neurons studied in rat brain slices (Stanford and Lacey, 1996; Richards et al., 1997).

Our single-cell RT–mPCR protocol probed for the mRNA expression of the cloned subunits currently thought to be involved in the formation of different K-ATP channels (Bryan and Aguilar-Bryan, 1997; Quale et al., 1997; Yokoshiki et al., 1998). GABAergic neurons of the SNpr showed a homogeneous K-ATP channel subunit profile, with SUR1 and Kir6.2 co-expression detected in all cells. This subunit combination leads to β-cell-like K-ATP channels in heterologous expression systems (Inagaki et al., 1995a; Gribble et al., 1997). SUR1 + Kir6.2-mediated K-ATP channels are activated by diazoxide and metabolic inhibition, and are blocked with high affinity by sulfonylureas such as glibenclamide and tolbutamide. In accordance with our single-cell RT–mPCR data, all GABAergic SNpr neurons expressed K-ATP channels on SNpr neurons (Schwanstecher and Panten, 1993; Stanford and Lacey, 1996). In contrast, GABAergic interneurons in the SNpc did not express any of the K-ATP channel subunits. In these cells, we could, in contrast, detect the presence of Girk2 transcripts, indicating that the amount of harvested material was sufficient to detect low abundance ion channel mRNA. As we also did not observe functional K-ATP channels in these neurons, our results suggest that GABAergic SNpc interneurons do not express SUR1- or SUR2B-mediated K-ATP channels.

Molecular and pharmacological diversity of K-ATP channels in dopaminergic SN neurons

In contrast to the homogeneous K-ATP subunit expression profiles detected in GABAergic SN neurons, the pattern alternative SUR isoform expression as those from C57Bl/6J mice (Figure 9B and C). These SUR1-containing K-ATP channels were already partially activated under metabolic control conditions in wv/wv dopaminergic SN neurons and were completely blocked by 100 µM tolbutamide (wv/wv K-ATP conductance: 1.2 ± 0.3 nS, n = 9). These results suggest that alternative SUR expression is relevant in the context of neurodegeneration and that expression of SUR1-containing K-ATP channels might play a protective role for dopaminergic SN neurons during the active period of neurodegeneration in the weaver mouse.

Fig. 9. Homogeneous SUR1 expression in surviving dopaminergic SN neurons of a homozygous weaver mouse. Distribution of SUR isoform expression in dopaminergic (TH⁺) SN neurons from (A) homozygous weaver (wv/wv) mice (n = 22), (B) homozygous normal (+/+ ) controls (n = 7) and (C) C57Bl/6J mice (n = 54). Note the absence of SUR2 expression in wv/wv dopaminergic SN neurons and the similar SUR isoform distribution in dopaminergic SN neurons from C57Bl/6J and (+/+) B6/CBA-A⁻/⁻AC mice.

Discussion
Expression of K-ATP channel subunits in midbrain neurons

Document:

of K-ATP subunit expression was more complex for dopaminergic SN neurons. We defined three populations with alternative K-ATP subunit expression profiles. In the two major populations, the transcripts for the pore-forming part of the K-ATP channel, Kir6.2, was co-expressed with the mRNA for either SUR1 (39%) or SUR2B (37%). The remaining population displayed co-expression of both SUR1 and SUR2B with Kir6.2 (24%). In correspondence with the single-cell RT–mPCR results, we observed functional K-ATP channels with significant differences in sulfonylurea sensitivities in dopaminergic SN neurons. The K-ATP channel population with the highest tolbutamide sensitivity (IC\textsubscript{50} = 8.8 µM) was indistinguishable from that found in GABAergic SNpr neurons. This K-ATP channel subtype is therefore likely to be mediated by SUR1 + Kir6.2 co-expression. In heterologous expression systems, SUR2-mediated K-ATP channels show a lower sulfonylurea sensitivity compared with SUR1-mediated channels (Isomoto and Kurachi, 1997; Babenko et al., 1998; Gribble et al., 1998). Thus, it is reasonable to assume that the K-ATP channels with the lowest tolbutamide sensitivity (IC\textsubscript{50} = 60.6 µM) are formed by SUR2B + Kir6.2 co-expression in dopaminergic SN neurons. It is unknown whether SUR1 and SUR2B form functional heterooligomeric K-ATP channels in the presence of Kir6.2. However, both our single-cell RT–mPCR results demonstrating SUR1 + SUR2B co-expression in single neurons and the identification of K-ATP channels with intermediate sulfonylurea sensitivity (IC\textsubscript{50} = 27.9 µM) suggest this possibility.

Our analysis demonstrated the diversity of K-ATP subunit expression profiles in midbrain neurons. Two recent reports analysing the K-ATP subunit expression in three rat dorsal vagal neurons (SUR1 + Kir6.2) and five rat striatal interneurons (SUR1 + Kir6.1) further illustrate the molecular complexity of neural K-ATP channels (Karschin et al., 1998; Lee et al., 1998).

Alternative sulfonylurea receptor expression correlates with metabolic sensitivity of K-ATP channels in single dopaminergic SN neurons

Our results show that K-ATP channels with very large differences in metabolic sensitivity (70- to 200-fold) were activated in dopaminergic SN neurons by inhibition of both mitochondrial complex I and glycolysis. K-ATP channels which were highly responsive to metabolic inhibition were completely blocked by 100 µM tolbutamide, indicating that they possess a high affinity for sulfonylureas, while low sensitivity K-ATP channels were only partially blocked by this concentration of tolbutamide. Although sulfonylurea sensitivity of K-ATP channels might be modulated by other factors in addition to subunit composition (Brady et al., 1998), the correlation between expression profiles and pharmacological properties demonstrated for GABAergic and dopaminergic SN neurons in this study suggested that SUR1 is involved in the formation of K-ATP channels acting as highly sensitive metabolic response elements. We tested this hypothesis directly by combining rotenone pre-incubation with the single-cell RT–mPCR protocol. Our results clearly demonstrate, at the level of individual neurons, that cells with fully activated K-ATP channels and large membrane hyperpolarizations in response to 100 nM rotenone co-express SUR1 + Kir6.2, while unresponsive cells co-expressed SUR2B + Kir6.2. The neurons which possessed both SUR1 and SUR2B at the transcript level showed an intermediate metabolic sensitivity. Although the molecular mechanisms remain undefined, a higher metabolic sensitivity has also been reported for heterologously expressed SUR1-containing channels compared with SUR2-containing channels (Ashcroft and Gribble, 1998), indicating that differences in metabolic sensing might indeed be linked directly to alternative expression of SUR isoforms. Much has been speculated about the functional response of dopaminergic SN neurons to partial inhibition of mitochondrial complex I activity which is present in Parkinson’s disease (Hanna and Bhatia, 1997). The selective activation of SUR1-containing K-ATP channels by low nanomolar concentrations of rotenone, which corresponds to a partial complex I inhibition, identifies this K-ATP channel subtype as a sensitive target for the metabolic challenge of Parkinson’s disease.

K-ATP channel expression in surviving dopaminergic SN neurons of weaver mouse: alternative SUR expression as a candidate mechanism for differential vulnerability in neurodegeneration?

In striking contrast to the alternative SUR expression profiles detected in dopaminergic SN neurons of wild-type mice [C57Bl/6J and (+/+)] B6/CBA-A\(^{wv}/\)AC\(^{wv}\)], all analysed neurons in homozygous weaver (wv/wv) mice exclusively expressed SUR1 + Kir6.2. In addition, these SUR1-containing K-ATP channels were already partially activated under metabolic control conditions in wv/wv dopaminergic SN neurons. The weaver mouse is a genetic model for the selective degeneration of a highly vulnerable subpopulation of dopaminergic SN neurons (Bayer et al., 1995; Patil et al., 1995). At postnatal day 14, which is the age we investigated, 35% of dopaminergic SN neurons are already lost (Verney et al., 1995). Our findings that SUR2 expression was never detected in surviving dopaminergic wv/wv neurons and that 37% of wild-type dopaminergic SN neurons express only SUR2 might indicate that the SUR2-expressing subpopulation of dopaminergic SN neurons corresponds to that possessing high vulnerability. Furthermore, all surviving dopaminergic wv/wv neurons exclusively expressed SUR1. This suggests that the activity of SUR1-containing K-ATP channels might have a neuroprotective role in the weaver mouse. There are great similarities in the pattern of differential vulnerability of dopaminergic subpopulations between the weaver mouse and Parkinson’s disease (Gaspar et al., 1994). Thus, it is conceivable that alternative SUR expression also constitutes a molecular mechanism involved in the differential vulnerability of dopaminergic neurons in Parkinson’s disease.

Materials and methods

Slice preparation

Young C57Bl/6J (12–16 days old), unaffected (+/+) and homozygous (wv/wv) weaver (B6/CBA-A\(^{wv}/\)AC; 14 days old) mice were anaesthetized deeply with halothane and then decapitated. Weaver mice were genotyped by genomic DNA sequencing. Brains were removed quickly, immersed in ice-cold solution, and then blocked for slicing. Thin coronal midbrain slices (200–250 µm) were cut with a Vibroslice (Campden Instruments,
London, UK) while bathed in an ice-cold artificial cerebrospinal fluid (ACSF) containing: 125 mM NaCl, 25 mM NaHCO$_3$, 2.5 mM KCl and 25 mM glucose, bubbled with a mixture of 95% O$_2$/5% CO$_2$. After sectioning, midbrain slices were maintained submerged in a holding chamber filled with gassed ACSF, and allowed to recover for >30 min at room temperature (22–24°C) before the experiment. The midbrain slices were first subjected to Student’s t-test in SIGMAPLOT (Jandel).

**Whole-cell recordings and data analysis**
For patch-clamp recordings, midbrain slices were transferred to a chamber continuously perfused at 2–4 ml/min with ACSF bubbled with O$_2$/CO$_2$. All experiments were performed electronically in an external solution containing: 145 mM NaCl, 2.5 mM KCl, 10 mM HEPES, 2 mM CaCl$_2$, 2 mM MgCl$_2$, pH 7.4. At the end of the procedure, Pipettes were then quickly removed from the cell, washed in an ice-cold artificial cerebrospinal fluid (ACSF) containing: 125 mM NaCl, 25 mM NaHCO$_3$, 2.5 mM KCl and 25 mM glucose, bubbled with a mixture of 95% O$_2$/5% CO$_2$. After sectioning, midbrain slices were maintained submerged in a holding chamber filled with gassed ACSF, and allowed to recover for >30 min at room temperature (22–24°C) before the experiment. The midbrain slices were first subjected to Student’s t-test in SIGMAPLOT (Jandel).

**Multiplex and nested PCR**
Multiplex and nested PCR was used for the experiments.

**Cytoplasm harvest and reverse transcription**
For single-cell RT-PCR experiments, the patch pipettes were filled with 6 µl of autoclaved internal RT-PCR solution containing: 140 mM KCl, 5 mM HEPES, 5 mM EGTA, 3 mM MgCl$_2$, pH 7.3. At the end of the recording (<15 min), the cell contents (including the nucleus, in most cases) were aspirated as completely as possible into the patch pipette under visual control (40x objective × 2–4x zoom) by application of gentle negative pressure. Cells were only analysed further when the whole-cell configuration remained stable throughout the harvesting procedure. Pipettes were then quickly removed from the cell, washed twice through the solution interface, and the pipette contents were expelled immediately into a 0.5 ml test tube containing the contents for reverse transcription. First strand cDNA was synthesized for 1 h at 37°C in a total reaction volume of 10 µl containing random hexamer primers (Boehringer Mannheim, final concentration 5 µM), diithiothreitol (final concentration 10 mM), the four deoxyribonucleotide triphosphates (Pharmacia, final concentration 0.5 mM each), 20 U of ribonuclease inhibitor (Promega) and 100 U of reverse transcriptase (Superscript™II; Gibco-BRL). The single-cell cDNA was kept at −70°C until PCR amplification.

**Multiplex and nested PCR**

**Whole-cell recordings and data analysis**
For patch-clamp recordings, midbrain slices were transferred to a chamber continuously perfused at 2–4 ml/min with ACSF bubbled with a mixture of 95% O$_2$/5% CO$_2$. After sectioning, midbrain slices were maintained submerged in a holding chamber filled with gassed ACSF, and allowed to recover for >30 min at room temperature (22–24°C) before the experiment. The midbrain slices were first subjected to Student’s t-test in SIGMAPLOT (Jandel).

**Cytoplasm harvest and reverse transcription**
For single-cell RT-PCR experiments, the patch pipettes were filled with 6 µl of autoclaved internal RT-PCR solution containing: 140 mM KCl, 5 mM HEPES, 5 mM EGTA, 3 mM MgCl$_2$, pH 7.3. At the end of the recording (<15 min), the cell contents (including the nucleus, in most cases) were aspirated as completely as possible into the patch pipette under visual control (40x objective × 2–4x zoom) by application of gentle negative pressure. Cells were only analysed further when the whole-cell configuration remained stable throughout the harvesting procedure. Pipettes were then quickly removed from the cell, washed twice through the solution interface, and the pipette contents were expelled immediately into a 0.5 ml test tube containing the contents for reverse transcription. First strand cDNA was synthesized for 1 h at 37°C in a total reaction volume of 10 µl containing random hexamer primers (Boehringer Mannheim, final concentration 5 µM), diithiothreitol (final concentration 10 mM), the four deoxyribonucleotide triphosphates (Pharmacia, final concentration 0.5 mM each), 20 U of ribonuclease inhibitor (Promega) and 100 U of reverse transcriptase (Superscript™II; Gibco-BRL). The single-cell cDNA was kept at −70°C until PCR amplification.

**RNA isolation and cDNA preparation for control reactions**
Poly(A)$^+$ RNA was prepared from fresh ventral midbrain and heart of 13–day-old C57Bl/6J mice using the Micro-FastTrack™ Kit (Invitrogen). The reverse transcription was performed with 500 ng of poly(A)$^+$ RNA as described above. For the positive controls carried out in parallel with each single-cell amplification, the resulting midbrain cDNA stock was diluted 1000-fold and 1 µl was used as template for the PCR. All eight PCR fragments were detected routinely in the positive control reactions. Negative controls were carried out in parallel to single-cell experiments excluding only the harvesting procedure, and resulted in no detectable bands. To probe for possible amplification of genomic DNA from the harvested single nuclei, in particular from the intronless Kir6.2 gene, single-cell mPCR amplifications were carried out without prior reverse transcription. For all analysed neurons ($n$ = 10), no PCR products were detectable. All other primer pairs were designed to be intron spanning.

**Set of primers**

**Categorization of primers**
Set of primers (from 5’ to 3’): Kir6.2 (accession No. D505581) sense, AAAGAAAGCACTGCAACGT (position 243), antisense, CCCCATGAATCTGTCAGC (position 1066); Kir6.1 (accession No. D88159) sense, GTCCCTAGGACCCATGTG (position 126), antisense, TATCACAGGGGCTACGC (position 786); SUR1 (accession No. rat 2091), antisense, GTGATGTTCTCCTCCACGCGT (position rat 2022); SUR2 (accession No. D86038) sense, GAGACGAGAAGATTGCGCAT (position 2148), antisense, CTATGATCCAGTACGCGT (position 2802); SUR2A/B sense, TGTCGACATATTGACGCA (position 4235), antisense, CTCCAATCCACGTCAAGG (position 4900 for SUR2A and SUR2B, respectively); TH (accession No. M692090) sense, CACCTGGAGATCTTGTGCG (position 387), antisense, CTTGTTGGTGATCCCATGT (position 1525); Girk2 (accession No. U11859) sense, CCTACGGATCTGTCGGACAC (position 711), antisense, GGTGCTGTTCTCATACTGGTC (position 1580); GFP (accession No. K10347) sense, AGAAACACATGCTGTGCTAT (position 407), antisense, CTCAATACACAGCTTTCG (position 1213); GAD67 (accession No. Z49976) sense, TGACGTCATGACCCATACC (position 731), antisense, GGTTTAGAGATGCCATACCG (position 1835). First multiplex-PCR was performed as hot start in a final volume of 100 µl containing the 10 µl reverse transcription reaction, 100 pmol of each primer, 0.2 µl each dNTP (Pharmacia), 1.8 mM MgCl$_2$, 50 mM KCl, 20 mM Tris–HCl (pH 8.4) and 3.5 µl of Tyg polymerase (Gibco-BRL) in a Perkin Elmer Thermal Cycler 480C with the following cycling protocol: after 3 min (94°C), 35 cycles (94°C, 30 s; 58°C, 60 s; 72°C, 3 min) of PCR were performed followed by a final elongation period of 7 min at 72°C. The nested-PCR amplifications were carried out in eight individual reactions, in each case with 2.5 µl of the first PCR product under similar conditions with the following modifications: 50 µl of each primer, 2.5 µl of Tyg polymerase, 1.5 mM MgCl$_2$ and a shorter extension time (60 s) using the following primer pairs: Kir6.2 sense, GCTGATCCTTCATGAAAGC (position 625), antisense, TTGGGACCGATGAGGTTGTA (position 1262); Kir6.1 sense, GCACAAGAAAAGGCAACTGCAACGT (position 936), antisense, GCCTGAAATACGTGTCCT (position 786); SUR1 sense, TCTAGAGTGTTGAGCCTGGA (position 2186), antisense, GTTCTCTGATGCTGAGGCC (position 2586); SUR2 sense, GGACGACAACTGAAATCGTC (position 2550), antisense, TCTGCTCAGACAAACAGG (position 2764); SUR2A/B sense, CTCTGACACGCTTCACCC (position 4463), antisense, ACAATGTCGACGGAACAG (position 4972), antisense, GAGACGAAACAACCATCGC (position 936), antisense, TCTGACACGAGTACCCCG (position 1312); Kir2 sense, AGACAGAAACACCATCGC (position 936), antisense, GAACGCCGTCTCATGCA (position 1531); GFP sense, AGGAAGGTTGAATGCGCTGGA (position 472), antisense, CCAGGCTCATTTAAGG (position 988); GAD67 sense, CATATGAAATGACCCCTTG (position 761), antisense, CGTGGTACGAGAACCTGCAG (position 1462). The nested-PCR products were amplified in 15 µl aliquots of PCR products were separated and visualized in an ethidium bromide-stained agarose gel (2%) by electrophoresis. The predicted sizes (bp) of the eight PCR-generated fragments were: 215 (SUR2), 337 or 513 (SUR2B and SUR2A, respectively), 298 (Kir6.2), 377 (TH), 401 (SUR1), 448 (Kir6.1), 517 (GFP), 595 (Kir2) and 702 (GAD67). All individual PCR products were verified several times (n > 3) by direct sequencing or by subcloning and sequencing.
Expression of K-ATP channel subunits in midbrain neurons


Received October 20, 1998; revised and accepted December 18, 1998