Adenosine kinase is a target for the prediction and prevention of epileptogenesis in mice

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Astrogliosis is a pathological hallmark of the epileptic brain. The identification of mechanisms that link astrogliosis to neuronal dysfunction in epilepsy may provide new avenues for therapeutic intervention. Here we show that astrocyte-expressed adenosine kinase (ADK), a key negative regulator of the brain inhibitory molecule adenosine, is a potential predictor and modulator of epileptogenesis. In a mouse model of focal epileptogenesis, in which astrogliosis is restricted to the CA3 region of the hippocampus, we demonstrate that upregulation of ADK and spontaneous focal electroencephalographic seizures were both restricted to the affected CA3. Furthermore, spontaneous seizures in CA3 were mimicked in transgenic mice by overexpression of ADK in this brain region, implying that overexpression of ADK without astrogliosis is sufficient to cause seizures. Conversely, after pharmacological induction of an otherwise epileptogenesis-precipitating acute brain injury, transgenic mice with reduced forebrain ADK were resistant to subsequent epileptogenesis. Likewise, ADK-deficient ES cell–derived brain implants suppressed astrogliosis, upregulation of ADK, and spontaneous seizures in WT mice when implanted after the epileptogenesis-precipitating brain injury. Our findings suggest that astrocyte-based ADK provides a critical link between astrogliosis and neuronal dysfunction in epilepsy.

Introduction

Current epilepsy therapies largely rely on the suppression of spontaneous seizures by pharmacotherapy or surgical intervention, but to date no effective prophylaxis or true pharmacotherapeutic cure is available. Epileptogenesis, i.e., the process that leads to epilepsy and spontaneous seizures, is thought to be triggered by an initial acute brain injury (e.g., status epilepticus [SE]), followed by progressive neuronal cell loss, mossy fiber sprouting, and formation of an astrogliotic scar. However, it is presently unclear why some brain injuries evolve into epilepsy while others do not. Therefore, the identification of diagnostic markers to predict epileptogenesis is of utmost importance. The identification of astrogliosis as a hallmark in brain of epileptogenesis and the identification of astrocytes as important modulators of neuronal activity imply that dysfunction of astrocytes might play a key role in the pathogenesis of epilepsy.

Animal models of epileptogenesis have been developed that closely mimic the pathogenesis of human mesial temporal lobe epilepsy, a form of human focal epilepsy that is frequently associated with progression to chronic intractable epilepsy. These models rely on the intrahippocampal (4), intraamygdaloid (5), or intracerebroventricular (6) administration of small doses of kainic acid (KA). As a primary consequence of KA injection, SE is elicited, which in turn — during a time span of several weeks — leads to a characteristic pattern of hippocampal cell death, mossy fiber sprouting, astrogliosis, and spontaneous recurrent seizures (4–6).

Adenosine is an inhibitory modulator of brain activity. It acts at 4 G-protein–coupled receptor subtypes (A1, A2A, A2B, and A3) and in hippocampus exerts predominantly inhibitory functions by activation of A1 receptors (A1Rs) (8). The inhibitory actions of adenosine can be used therapeutically to suppress seizures (9) and are considered important for maintaining postictal depression (10) and for restoring the metabolic equilibrium following seizures (11). However, despite more than 20 years of research on the role of adenosine in experimentally induced seizures and the identification of adenosine as endogenous anticonvulsant of the brain (12–15), the pathogenetic role of the adenosine system in epileptogenesis remains understudied.

Ambient concentrations of adenosine are largely regulated by the activity of its major metabolic enzyme adenosine kinase (ADK) (16), which in the adult brain is predominantly expressed in astrocytes (17). Consequently, astrogliosis in epilepsy is associated with upregulation of ADK (18). Transgenic upregulation of ADK in brain leads to a reduction in the tone of ambient adenosine and therefore is associated with increased susceptibility to seizures (19) and ischemic cell death (20). These findings provide a neurochemical rationale for therapeutic intervention (21). Consequently, cultured cells engineered to release adenosine by disrupting their ADK genes were tested as therapeutic brain implants in the rat kindling model (22). These studies have provided the proof-of-concept that focal augmentation of adenosine by cellular brain implants can reduce induced (kindled) seizures and the progression of kindled seizure severity (22, 23). However, it remains to be demonstrated whether focal augmentation of adenosine can prevent the development of spontaneous seizures, i.e., true epileptogenesis.

To characterize ADK as a target for the prediction and prevention of epileptogenesis we followed 3 independent lines of research: characterization of the pathogenetic role of ADK in a mouse model of focal epileptogenesis, study of epileptogenesis in a panel of transgenic mice with altered brain expression levels of ADK, and intracerebral transplantation of ADK-deficient stem cells to

Nonstandard abbreviations used: ADK, adenosine kinase; KA, kainic acid; NP, neural precursor cell; SE, status epilepticus.

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modify epileptogenesis. Our results support the ADK hypothesis of epileptogenesis and define ADK as a promising target for the prediction and prevention of epileptogenesis.

Results

Astrogliosis, upregulation of ADK, and focal spontaneous seizures characterize epileptogenesis in WT mice. Adult male WT C57BL/6 mice (n = 6) received intraamygdaloid KA injections. As a consequence of the KA injection, all animals developed nonconvulsive electrographic SE and CA3-selective neuronal cell loss, as previously described (24). All animals survived a time span of 3 weeks, when electrodes were implanted into the ipsilateral CA3 region. During 8 hours of continuous monitoring, spontaneous focal seizures were recorded from the CA3 region of all animals, with 4.3 ± 1.5 seizures per hour and an average duration of 17.5 ± 5.8 seconds (Figure 1, bottom left), indicating epileptogenesis. It is important to note that (a) spontaneous seizures were observed in every KA-injected animal, (b) that electrographic seizures were never associated with convulsions, and (c) that seizure activity was specific to the injured ipsilateral CA3 region. In additional animals (n = 6 each) with electrodes placed in CA1 or onto the cortex, we were not able to record any seizures (Figure 1), demonstrating that epileptogenesis in WT animals is highly specific for the ipsilateral CA3 region, the region of acute neuronal cell loss. In a separate set of animals (n = 3) that were recorded daily after SE using CA3 electrodes, the first onset of spontaneous seizures was determined. In all animals, spontaneous CA3 seizures first emerged 12 days after SE (3.1 ± 0.9 seizures per hour, with a duration of 15.7 ± 6.0 seconds).
A

Adk transgene

\[ \text{Adktm1}^+/-\text{Tg(UbiAdk)} = \text{"Adk-tg"} \]

\[ \text{Adktm1}^+/-\text{Tg(UbiAdk):Emx1-Cre-tg3} = \text{"fb-Adk-def"} \]

B

Breeding strategy

C

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D

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<tr>
<td>BSCB</td>
<td>be-ch</td>
<td>be-ch</td>
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</table>

E

ADK (%)

After completion of EEG recordings, all animals were sacrificed for histology. Analysis of cresyl violet–stained brain sections showed ipsilateral CA3 injury (Figure 1, B and D), indicating the presence of the epileptogenesis-preventing acute injury, while the contralateral CA3 region was spared from injury (Figure 1, A and C). Fluorescence staining for nuclei revealed the presence of an increased number of nuclei in the ipsilateral compared with the contralateral CA3 (Figure 1, E and F), indicating an increase in nonneuronal cells within the injured CA3. To investigate whether neuronal cell loss in the injured CA3 (Figure 1D) was accompanied by a loss of neuronal processes, we performed an immunofluorescence staining with a marker for neuronal filaments (NFM) (Figure 1, G and H). Despite the reduction of neuronal cell bodies within the injured CA3, the amount of NFM immunoreactivity was unchanged, indicating a dense network of neuronal processes within the injured CA3 likely originating from neuronal cell bodies elsewhere. Since astrogliosis and overexpression of endogenous ADK may be linked to epileptogenesis, we analyzed expression of the astrocyte markers GFAP and ADK in the hippocampal formation in more detail. All animals displayed a striking and highly localized astrogliotic response covering the rostrocaudal extent of the ipsilateral CA3 (Figure 1, J, L, and N), with colocalized overexpression of GFAP and ADK (Figure 1, Q and R), while the contralateral hippocampus was spared from astrogliosis and upregulation of ADK (Figure 1, I, K, M, O, and P). Confocal imaging (Figure 1, O, P, Q, and R) revealed not only increased ADK immunoreactivity and the predominantly nuclear localization of the enzyme but also a redistribution and spread of ADK immunoreactivity to extranuclear locations (Figure 1R).

Generation of transgenic mice with altered levels of brain ADK. To provide experimental evidence that upregulation of ADK during epileptogenesis is causative for the development of spontaneous seizures in epileptogenesis, we created a panel of transgenic mice with either enhanced or reduced expression of brain ADK. To generate a mouse model with a forebrain-selective reduction of ADK, we bred Emx1-Cre-Tg3 mice, which express Cre recombinase in neurons and astrocytes of the telencephalon (25), with ADK transgenic mice, which carry a loxP-flanked ADK transgene (Adktm1\(^{+/-}\)) (19) (Figure 2A) in an otherwise lethal ADK KO (Adktm1\(^{-/-}\)) background (26). Adktm1\(^{+/-}\)Tg(UbiAdk) (referred to here as “Adk-Tg”) and Adktm1\(^{+/-}\)Tg(UbiAdk):Emx1-Cre-Tg3 (referred to here as “fb-Adk-def”) offspring were produced as littermates (Figure 2B). Genotyping was performed at weaning and again retrospectively from all experimental animals (Figure 2C).

To demonstrate that Cre recombination of the ADK transgene in fb-Adk-def mice resulted in a forebrain-selective reduction of ADK, the expression pattern of ADK was analyzed by immunoperoxidase staining on coronal sections from adult WT, fb-Adk-def, and Adk-Tg mice (n = 3 per group). In WT sections, strong ADK immunoreactivity was largely restricted to nuclei and cell bodies of evenly distributed astrocytes (Figure 3, A and B), as previously described (17), whereas pyramidal cells of the CA3 region (Figure 3B) lacked specific ADK immunoreactivity. In contrast, Adk-Tg mice were characterized by lack of the endogenous astrocytic ADK (lack of...
discrete staining of astrocytic nuclei and cell bodies due to genetic disruption of the endogenous Adk gene), which was replaced by a ubiquitous overexpression of transgenic ADK (Figure 3C), resulting in strong diffuse immunoreactivity in neurons and astrocytes all over the brain but lack of nuclear localization. Most importantly, a novel neuronal expression of ADK was found, particularly in pyramidal neurons of the CA3 region (Figure 3D). In fb-Adk-def mice, ADK reduction — induced by Cre-mediated deletion of the loxP-flanked transgene — became apparent in the whole dorsal telencephalon (Figure 3E), encompassing cerebral cortex and hippocampus, while basal brain regions continued to display the diffuse overexpression of ADK, in line with unaltered expression of the Adk transgene (Figure 3, C and F). Only a few individual cells with neuronal morphology within the forebrain remained positive for ADK (Figure 3G), most likely GABAergic interneurons, a cell type in which Emx1-Cre is not expressed (25).

To quantify the reduction of ADK in the dorsal telencephalon of fb-Adk-def mice, a western blot analysis was performed comparing the intensity of ADK immunoreactivity in dorsal telencephalon with brain stem plus cerebellum from tissue samples taken from WT, fb-Adk-def, and Adk-Tg mice (Figure 2D). The forebrains of fb-Adk-def mice had significantly reduced levels of ADK, with an average of 62% ± 27% (P < 0.01) compared with WT forebrain (100%), while ADK in forebrain of Adk-Tg mice was elevated to 147% ± 49% of WT levels (P < 0.05). In both fb-Adk-def and Adk-Tg mice, brainstem plus cerebellum ADK levels were elevated to an average of 141% ± 50% and 141% ± 30% of WT levels, respectively (Figure 2E).

Adk-Tg mice are characterized by spontaneous seizures in CA3 and lethal KA-induced SE. If upregulation of ADK in the CA3 region during epileptogenesis (Figure 1) is the cause for seizures rather than astrogliosis, then transgenic overexpression of ADK within the CA3 should result in spontaneous seizures. Adk-Tg mice display a particularly prominent overexpression of ADK within the CA3 region (Figure 3D). To test for spontaneous electrophysiological seizures, we therefore inserted bipolar recording electrodes directly into the CA3 region of Adk-Tg mice (n = 4). Eight hours of continuous EEG recordings yielded frequent spontaneous seizures (Figure 3E), with an average of 4.8 ± 1.5 seizures per hour, each seizure lasting 26.7 ± 13.2 seconds. Remarkably, transgenic overexpression of ADK yielded a similar seizure frequency as astrogliotic upregulation of ADK in epileptic WT mice (4.3 ± 1.5 seizures per hour).

To determine the susceptibility to seizures, Adk-Tg (n = 5) and WT (n = 6) mice received intraamygdaloid injections of KA. Following the KA injections, seizure activity was monitored in EEG recordings with a set of 3 cortical electrodes. After 30 minutes, seizures were terminated with lorazepam. Seizures were graded into type I–IV activities, as previously described (24). While WT mice developed typical type IV seizures (Figure 4), Adk-Tg mice developed type IV seizures of increased intensity (Figure 4). In Adk-Tg mice, the duration of type IV activity (713 ± 186 s) was significantly longer than that of WT mice (62% ± 27% (P < 0.01) compared with WT forebrain (100%), while ADK in forebrain of Adk-Tg mice was elevated to 147% ± 49% of WT levels (P < 0.05). In both fb-Adk-def and Adk-Tg mice, brainstem plus cerebellum ADK levels were elevated to an average of 141% ± 50% and 141% ± 30% of WT levels, respectively (Figure 2E).

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prolonged \( (P < 0.05) \) compared with WT (Table 1) and finally led to the death of 2 of 5 animals within 24 hours (Table 1). In an additional group of KA-injected Adk-Tg mice \((n = 5)\) we observed to investigate long-term consequences of KA-induced SE, all animals died within 3 days of the KA injection (Table 1).

To investigate the influence of overexpressed ADK on the development of acute seizure-induced hippocampal injury, 24 hours after KA injection all remaining Adk-Tg mice \((n = 3)\) were analyzed histologically by cresyl violet and TUNEL staining (Figure 5 and Table 1). KA-injected WT mice developed a well-defined lesion in the ipsilateral CA3 of the hippocampus, which was characterized by prominent TUNEL-positive cells, being indicative of ongoing cell death (Figure 5, B and C). KA-induced seizures affected neither the contralateral hippocampus (Figure 5A) nor any other brain region. In contrast, ADK overexpression in Adk-Tg mice led to highly aggravated cell loss in the ipsilateral CA3 region (Figure 5K and L, and Table 1) as well as to an extension of cell loss in the ipsilateral CA1 region (Figure 5N) and dentate gyrus (Figure 5K), and most remarkably, to significant cell loss in the contralateral hippocampus (Figure 5, J, M, and O, and Table 1).

**Table 1**
Mortality, electrographic seizure activity, and cell death 24 hours after intraamygdaloid KA injection

<table>
<thead>
<tr>
<th>Parameter</th>
<th>WT</th>
<th>Adk-Tg</th>
<th>fb-Adk-def</th>
<th>fb-Adk-def + DPCPX</th>
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</thead>
<tbody>
<tr>
<td>Mortality (24 h)</td>
<td>0/6</td>
<td>2/5</td>
<td>0/5</td>
<td>0/5</td>
</tr>
<tr>
<td>Mortality (3 d)</td>
<td>0/5</td>
<td>5/5</td>
<td>0/5</td>
<td>0/5</td>
</tr>
<tr>
<td>Average type IV seizure duration (s)</td>
<td>443.3 ± 147.2 ((n = 6))</td>
<td>713.3 ± 185.8 ((n = 3))</td>
<td>0 ((n = 5))</td>
<td>318.0 ± 100.6 ((n = 5))</td>
</tr>
<tr>
<td>TUNEL count, ipsilateral CA3B</td>
<td>73.8 ± 17.1</td>
<td>117.6 ± 31.1 (A)</td>
<td>0</td>
<td>51.8 ± 17.7</td>
</tr>
<tr>
<td>TUNEL count, ipsilateral CA1B</td>
<td>0</td>
<td>38.6 ± 4.0 (B)</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>TUNEL count, contralateral CA3B</td>
<td>0</td>
<td>28.3 ± 10.4 (C)</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>TUNEL count, contralateral CA1B</td>
<td>0</td>
<td>25.0 ± 5.6 (D)</td>
<td>0</td>
<td>0</td>
</tr>
</tbody>
</table>

\( ^A P < 0.05 \) compared with WT by \( t \) test. \( ^B \) The average number of TUNEL-positive cells per brain section was determined by counting TUNEL-positive cells in 3 brain sections from \( n = 3 \) mice per genotype. \( ^C P < 0.001; \) \( ^D P < 0.01 \) compared with WT by \( t \) test.

\( ^C P < 0.001; \) \( ^D P < 0.01 \) compared with WT by \( t \) test.

**Figure 5**
Epilepsy-associated cell loss. Representative micrographs of the hippocampal formation from coronal brain sections taken from WT mice (A–C), fb-Adk-def mice (D–F), fb-Adk-def mice pre-treated with 1 mg/kg i.p. DPCPX (G–I), and Adk-Tg mice (J–O) 24 h after intraamygdaloid injection of KA. Sections were stained either with cresyl violet or with TUNEL (green). (A–C) Typical cell death in the CA3 region of the hippocampus of WT mice ipsilateral (ipsi) to the KA-injected amygdala. (D–F) fb-Adk-def mice did not display any signs of cell loss. (G–I) After pretreatment with DPCPX, fb-Adk-def mice developed cell loss, which was comparable with WT animals. (J–O) Adk-Tg mice showed aggravated cell loss in the ipsilateral CA3 region and novel cell loss in the contralateral CA1 region (M), and the contralateral CA1 (N) and CA3 (O) regions. Scale bars: 300 \( \mu m \) (A, B, D, E, G, H, J, and K); 75 \( \mu m \) (C, F, I, L, M, N, and O).
seizures were never seen in fb-Adk-def mice (Figure 4). These mice developed normal types I and II activity, and type III activity of reduced intensity. This strong protection from seizures was due to elevated adenosine in the forebrain of the mutants acting via the $\text{A}_1\text{R}$, since application of the $\text{A}_1\text{R}$ antagonist DPCPX completely abolished the protection, resulting in type IV seizure activity (Figure 4) and duration (318 ± 101 seconds), which was comparable to WT (443 ± 147 seconds) ($P > 0.05$; Table 1).

To investigate the influence of reduced levels of ADK on the development of acute seizure-induced hippocampal injury, 24 hours after KA injection all animals were analyzed histologically by cresyl violet and TUNEL staining (Figure 5 and Table 1). In contrast to WT control animals, fb-Adk-def mice were completely protected from any injury (Figure 5, D–F). In fb-Adk-def mice, not a single TUNEL-positive cell was detected (Figure 5F and Table 1), indicating a powerful neuroprotective effect as a consequence of ADK deficiency. The obvious neuroprotection in fb-Adk-def mice was reversed when animals were pretreated with DPCPX, including hippocampal lesions (Figure 5, G–I) reminiscent of the lesions seen in WT animals and a similar number of TUNEL-positive cells per section (Table 1). To demonstrate that the chosen dose of DPCPX (1 mg/kg, i.p.) acts more by blocking effects of increased adenosine (as expected in forebrains of fb-Adk-def mice) on the $\text{A}_1\text{R}$ rather then interfering with “normal” levels of adenosine in WT mice, an additional group of WT mice ($n = 6$) was injected with DPCPX 30 minutes prior to intraamygdaloid KA injection. Brains of these
animals were analyzed 24 hours later for the presence of TUNEL-positive cells in the CA3 region. We found 78.2 ± 20.3 TUNEL-positive cells per CA3 section, a value comparable with controls (P > 0.05) with KA-injected WT mice not treated with DPCPX (73.8 ± 17.1; Table 1), indicating that the chosen dose of DPCPX normalized the effects of increased forebrain adenosine on A₃Rs in fb-Adk-def mice and therefore recreates a WT-like acute injury in these mutant animals.

(fb-Adk-def mice are resistant to epileptogenesis) To demonstrate that ADK could, via adenosine, not only have these acute effects but also influence epileptogenesis chronically, we compared consequences of KA injection after 3 weeks in WT mice (acute KA-induced SE and CA3 cell loss), fb-Adk-def mice (no acute changes), and fb-Adk-def mice pretreated 30 minutes before KA injection with a single dose of DPCPX (restoration of acute WT-like SE and CA3 cell death; Figures 4 and 5). Due to the transient nature of A₃R blockade, DPCPX-pretreated fb-Adk-def mice (fb-Adk-def/DPCPX) developed an acute epileptogenesis-initiating injury (SE, CA3 cell loss), but thereafter were under the influence of elevated forebrain adenosine.

WT, fb-Adk-def, and fb-Adk-def/DPCPX mice (n = 5 per group) received intraamygdaloid KA injections as described above. As a consequence of KA injection, all animals except fb-Adk-def mice developed SE analogous to that shown in Figure 4. All animals survived a time span of 3 weeks, when electrodes were implanted into the ipsilateral CA3 region. While spontaneous focal seizures were recorded from the CA3 region of all WT mice (Figure 6A), indicating epileptogenesis, spontaneous seizures were completely absent in fb-Adk-def and fb-Adk-def/DPCPX mice (Figure 6, B and C), indicating that elevated forebrain adenosine prevents the development of spontaneous recurrent seizures. After completion of EEG recordings, all animals were sacrificed for histology. Analysis of cresyl violet–stained brain sections showed the absence of the ipsilateral CA3 lesion in WT (Figure 6A) and fb-Adk-def/DPCPX mice (Figure 6C) but not in fb-Adk-def mice (Figure 6B), indicating the presence of the epileptogenesis-initiating primary injury (Figure 6, B and C). Since astrogliosis and overexpression of endogenous ADK were linked to epileptogenesis (Figure 1), we analyzed expression of the astrocyte markers GFAP and ADK in the hippocampal formation in more detail. Whereas WT animals displayed a striking and highly localized astrogliotic response within the ipsilateral CA3 with colocalized overexpression of GFAP and ADK (Figure 6A), no signs of astrogliosis or upregulation of ADK were found in fb-Adk-def or fb-Adk-def/DPCPX mice, even though the latter showed SE-induced CA3 cell loss similar to that of WT (Figure 6, B and C). The lack of astrogliosis, and consequently the absence of upregulated ADK, resulting in the lack of spontaneous seizures in fb-Adk-def mice, demonstrate that elevated forebrain adenosine can prevent epileptogenesis. To our knowledge, this is the first demonstration of a transgenic animal with resistance to epileptogenesis.

Adk−/− ES cell–derived neural progenitor grafts prevent development of spontaneous seizures. To determine whether adenosine-releasing intrahippocampal stem cell–derived implants could prevent the development of epileptogenesis, 3 groups of mice received implants of adenosine-releasing Adk−/− (KA-KO mice) (27) or respective WT (KA-WT mice) ES cell–derived neural precursor cells (NPs) or corresponding sham treatments (KA-SHAM mice) 24 hours after intraamygdaloid KA injection (n = 6 per group), i.e., at a time point at which the initial acute CA3 injury was already present. All animals survived a time span of 3 weeks, when electrodes were implanted into the ipsilateral CA3 region as described above. Spontaneous focal seizures (Figure 7, A and B) were recorded from the CA3 region of all KA-SHAM and KA-WT mice, with 4.2 ± 1.4 and 4.0 ± 1.3 seizures per hour and average durations of 17.3 ± 5.9 and 16.8 ± 5.5 seconds, respectively, values comparable to seizure parameters in KA-injected control mice (Table 2; P > 0.05). In contrast, not a single spontaneous seizure was detected after recording KA-KO mice with Adk−/− NP implants (n = 6) for 8 hours each (Figure 7C), a time span equivalent to >192 seizures in a corresponding number of control animals (n = 6).

Table 2
Spontaneous electroencephalographic seizures recorded from the CA3 region

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Adk-Tg (n = 4)</th>
<th>KA-control (n = 6)</th>
<th>KA-SHAM (n = 6)</th>
<th>KA-WT (n = 6)</th>
<th>KA-KO (n = 6)</th>
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<tbody>
<tr>
<td>Average duration (s)</td>
<td>26.7 ± 13.2</td>
<td>17.5 ± 5.8</td>
<td>17.3 ± 5.9</td>
<td>16.8 ± 5.5</td>
<td>0</td>
</tr>
<tr>
<td>Frequency (/h)</td>
<td>4.8 ± 1.5</td>
<td>4.3 ± 1.5</td>
<td>4.2 ± 1.4</td>
<td>4.0 ± 1.3</td>
<td>0</td>
</tr>
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</table>

Seizures were recorded from electrodes implanted within the CA3 region. Untreated Adk-Tg mice displayed spontaneous CA3 seizures in the absence of KA administration. Spontaneous seizures were recorded 3 weeks after KA injection in C57BL/6 WT mice (KA-control), in animals subjected to a sham procedure to mimic cell transplantation (KA-SHAM), and in WT graft recipients (KA-WT). No significant difference of average duration or frequency of spontaneous seizures were found in KA-control, KA-SHAM, or KA-WT groups (P > 0.05, ANOVA). In contrast, recipients of adenosine-releasing Adk-KO NPs (KA-KO) were completely protected from any seizures.
**Adk**<sup>−/−</sup> ES cell–derived neural progenitor grafts reduce astrogliosis and upregulation of ADK. After completion of the EEG recordings, i.e., 3 weeks after KA injection, all animals were processed for histoch­emistry. To verify the existence of the acute epileptogenesis triggering brain injury as becoming evident 24 hours after KA, i.e., before cell transplantation, we subjected coronal brain sections from KA-SHAM, KA-WT, or KA-KO animals and naive control animals (no KA injection) to cresyl violet staining. CA3-selective cell loss was only evident in the KA-injected animals (Figure 8, A–C), while the CA3 region remained intact in the control animals (Figure 8D). To investigate whether **Adk**<sup>−/−</sup> NPs prevented spontaneous seizures via reduction of astrogliosis and reduced upregulation of ADK, we analyzed expression of the astrocyte markers GFAP and ADK in the hippocampal formation in the 4 groups of mice. KA-WT and KA-SHAM animals displayed a highly localized astrogliotic response within the ipsilateral CA3 (Figure 8, E and F), with colo­calized overexpression of GFAP and ADK (Figure 8, M, N, Q, and R). In contrast, astrogliosis was markedly reduced (Figure 8, G, O, and S) and no obvious upregulation of ADK became evident within the ipsilateral CA3 of the KA-KO group representing the **Adk**<sup>−/−</sup> graft recipients (Figure 8, K, O, and S).

To generate quantitative data on GFAP and ADK expression, we determined the density of ADK immunoreactivity within the CA3 field and counted the number of GFAP-positive cells in ipsilateral and contralateral CA3 fields of KA-injected control animals and in ipsilateral hippocampus of KA-SHAM, KA-WT, and KA-KO animals (Figure 9). ANOVA analysis revealed a significant increase in the density of ADK immunoreactivity as well as the number of GFAP-positive astrocytes (P < 0.01 and P < 0.001, respectively) within the ipsilateral CA3 of the KA control animals (KA-SHAM and KA-WT). In contrast, the number of GFAP-positive astrocytes in the KA-KO animals was only moderately increased (P < 0.01), while changes in ADK immunoreactivity densities failed to attain statistical significance (P > 0.05). These data demonstrate that **Adk**<sup>−/−</sup>-derived brain implants prevent the upregulation of host ADK and thus eliminate a major factor contributing to seizure generation.

**Adk**<sup>−/−</sup> ES cell–derived neural progenitor cells display intrahippocampal location. In coronal brain sections from animals sacrificed 3 weeks after grafting, we detected dense cellular transplants in the vicinity of the surgical injection tract, located in the infrahippocampal cleft according to our previous studies (23) (Figure 10, A and B). Likewise, in all animals, we detected individual graft-derived cells in the CA1 region of the ipsilateral hippocampus (Figure 10, C and D). Most of these graft-derived cells showed a morphology and orientation comparable with mature hippocampal neurons, including an elaborate dendritic tree and a principal axon. We did not find any differences in graft location between WT and **Adk**<sup>−/−</sup> graft recipients. In all ani­mals (n = 12), the location of the graft was confined to the ipsilateral hippocampus, while the contralateral hippocampus as well as all other brain areas analyzed were devoid of graft-derived cells.
Discussion
As a long-term consequence of brain injury, astrogliosis is considered to be a hallmark of the epileptic brain. Autopsy and surgical resection specimens have demonstrated that chronic temporal lobe epilepsy and posttraumatic seizures may originate from gliotic scars (28, 29). The involvement of astrocytes in the pathogenesis of epileptogenesis can no longer be ignored since recent findings that astrocytes control synaptic transmission by the vesicular release of glutamate, the NMDA receptor coagonist D-serine, and ATP as a source of synaptic adenosine (2).

As a response to astrogliosis and chronic seizures, the adenosine system undergoes changes, which can promote further seizures. Thus downregulation of A1Rs has been described as a consequence of chronic seizures (30, 31). Likewise, a combined decrease in the density of A1Rs and metabolic changes leading to lower levels of adenosine were described in the hippocampus of epileptic rats (32). Indeed, overexpression of ADK within an epileptic hippocampus was associated with astrogliosis and focal seizures (18). Likewise, deficits in adenosinergic neuromodulation promote seizures in hippocampal slices (10, 33). Here we propose the ADK hypothesis of epileptogenesis and provide experimental evidence on 4 different levels:

Mouse model of focal epileptogenesis. We previously suggested that upregulation of ADK during astroglisosis might cause spontaneous seizures (18, 19). However, these studies were based on intrahippocampal injection of KA causing widespread hippocampal astrogliosis and upregulation of ADK. In this model of TLE, it was not possible to distinguish direct effects on hippocampal integrity caused by the excitotoxin kainate or secondary effects caused by the resulting SE. Widespread astrogliosis and upregulation of ADK in this model precluded the delineation of seizure generation at high spatial resolution. In the present investigation we therefore decided to study the adenosine system in a model of focal epileptogenesis, in which we used unilateral intraamygdaloid injections of KA. This model has the advantage of inducing an acute injury restricted to the ipsilateral hippocampus without direct manipulation of the hippocampus (24).

We believe that the chronic epileptogenic response to an acute intraamygdaloid KA-induced hippocampal injury as reported in this study (Figure 1) is a novel finding and involves a precipitating event (i.e., KA-induced SE), injury-induced cell death, highly localized astrogliosis, upregulation of glial-based ADK, and the development of focal spontaneous seizures at a frequency of 4.3 ± 1.5 seizures per hour. Remarkably, spontaneous seizures develop exclusively within the astrogliotic CA3 region but not elsewhere within the hippocampus; likewise, neither generalized seizures nor any convulsive seizures were detected. Since seizure recordings were performed during 8 hours of continuous monitoring in each of 6 animals, a total of >200 spontaneous seizures were recorded. However, the limitation to 8 hours of recording does not allow us to draw any conclusions on possible day/night fluctuations of seizure activity.

It is important to note that seizures originate in a focal area, with prominent neuronal cell death (Figure 1D). Nevertheless, neuronal processes originating from adjoining hippocampal areas

Figure 9
Quantitative assessment of ADK immunoreactivity and GFAP expression in the CA3 region. Quantitative data were generated from KA-control, KA-SHAM, KA-WT, and KA-KO animals 3 weeks after KA injection/cell transplantation. (A) The relative density of ADK immunoreactivity was determined on DAB-stained sections by scanning CA3a fields of 300 × 200 μm on 2 sections each from n = 6 animals per treatment group. Data were normalized to the contralateral hippocampus of KA-injected control animals. (B) The total number of GFAP-positive cells in corresponding CA3a fields of 300 × 200 μm was determined by counting GFAP-positive cells on 2 sections each from n = 6 animals per treatment group. Data analysis was done by ANOVA; mean ± SD. **P < 0.01; ***P < 0.001.

Figure 10
Adk–/– NP-derived cell implant 3 weeks after grafting. (A) Cresyl violet-stained overview showing a dense cluster of implanted cells within the infrahippocampal cleft (box). (B) Green fluorescence staining of graft-derived cells within the infrahippocampal cleft. (C and D) Graft-derived cells located within the ipsilateral CA1 region and displaying neuron-like morphology. Scale bars: 150 μm (A); 20 μm (B); 10 μm (C and D).
penetrate the injured CA3 (Figure 1H), are present within an area of upregulated ADK (Figure 1L), and are likely the source for the highly localized CA3 seizures (Figure 1). The restriction of both upregulated ADK and focal seizures to the injured CA3 suggests the existence of an endogenous mechanism preventing the spread of seizures into regions with normal ADK expression. This endogenous protective mechanism may rely on the release of adenosine during seizures and the activation of A1Rs, which have previously been implicated in keeping an epileptic focus localized (34, 35). These findings suggest that astrogliotic upregulation of ADK is an integral part of epileptogenesis and a cause for seizures. Indeed, previous studies in the mouse intrahippocampal KA model suggest that upregulation of ADK (first observed around 1 week after SE) precedes the development of spontaneous seizures (first observed around 2 weeks in the intrahippocampal KA model and around 12 days in this study) (18, 36). The results presented here document focal epileptogenesis at high spatial resolution and demonstrate that focal upregulation of ADK, which is not expected to alter the basal hippocampal adenosine-tone in other subregions, is sufficient to create a highly focused core of hyperexcitability.

Transgenic overexpression of ADK causes seizures. Adk-Tg mice display a striking overexpression of ADK in the hippocampal CA3 region (Figure 3D); however, in contrast to upregulated ADK as a component of epileptogenesis (Figure 1), transgenic overexpression is not associated with astroglialosis. Our finding of spontaneous seizures (4.8 ± 1.5 seizures per hour) in the CA3 of Adk-Tg mice at a similar frequency to seizures found in the KA-model 3 weeks after SE (4.3 ± 1.5 seizures per hour) suggests that overexpression of ADK in the absence of astroglialosis is sufficient to cause spontaneous seizures.

Reduction of ADK prevents epileptogenesis. By comparing the causal sequence of events triggered by the intraamygdaloid application of KA in Adk transgenic mice, which either overexpress ADK (Adk-Tg) or have reduced ADK (fb-Adk-def) in their forebrains (Figure 3), we demonstrate that the severity of KA-induced SE (Figure 4) and the degree of SE-induced cell death (Figure 5) are inversely related to the levels of ADK expression. While WT mice developed typical type IV seizures and a well-defined lesion in the ipsilateral CA3 of the hippocampus, this strong seizure response and selective CA3 damage were never observed in the fb-Adk-def mice. Most importantly, in fb-Adk-def mice pretreated with the A1R antagonist DPCPX, the initial KA-induced brain injury was comparable with WT mice, but due to elevated forebrain adenosine, subsequent epileptogenesis was prevented (Figure 6C). The efficacy of DPCPX in restoring WT-like brain injury in fb-Adk-def mice suggests that A1Rs are fully functional in fb-Adk-def mice, making secondary effects due to potential receptor desensitization unlikely. This DPCPX control experiment also demonstrates that fb-Adk-def mice are principally able to express KA-induced seizures, indicating full functionality of KA receptors in these animals. It is important to note that fb-Adk-def mice do not lack forebrain ADK. Such a complete lack of ADK would likely be lethal. Rather, these mutants are characterized by a subtle reduction (about 40%) and altered cellular distribution of ADK compared with WT. Similarly, ADK is only increased 1.5-fold in Adk-Tg mice, an increase that is completely sufficient to enhance brain vulnerability. The importance of subtle changes in ADK activity makes sense from a biochemical point of view. Adenosine levels are mainly regulated by an active substrate cycle between adenosine and AMP involving ADK and 5'-nucleotidase. Thus small changes in ADK can rapidly translate into major changes in adenosine (37–39). Because of difficulties in performing correct local adenosine measurements in living brain tissue and because of large variations according to the methods used (40), we previously chose an indirect electrophysiological approach to demonstrate ADK-dependent changes in ambient adenosine, confirming a reduction of endogenous adenosine in the brain of Adk-Tg mice (19). In line with reduced levels of brain adenosine, Adk-Tg mice have increased susceptibility to ischemic cell death (20) and have cognitive deficits and altered sensitivity to psychomimetic drugs, which are compatible with reduced brain adenosine (41). Conversely, the reversal of adenosine-mediated neuroprotection in forebrains of fb-Adk-def mice by a dose of the A1R antagonist DPCPX, which does not influence the activity of normal adenosine in WT mice, indicates increased levels of adenosine in the forebrain of fb-Adk-def mice. Since levels of ATP are about 100,000-fold higher than those of adenosine (7), since adenosine is the only substrate for ADK, and since the enzyme is not required for the de novo synthesis of purines, subtle changes in ADK are not likely to affect the pool of adenosine nucleotides nor any other metabolites. The fact that rather modest changes in ADK are needed to produce major effects has obvious implications for future therapy.

Preventive cell therapy. Previous therapeutic approaches have demonstrated that focal brain implants of fibroblasts, myoblasts, and stem cells engineered to release adenosine are effective in the suppression of kindled seizures (22). However, these approaches share a common caveat with currently used therapeutic strategies to treat epilepsy, in affecting merely the symptoms of epilepsy (i.e., seizures) rather than providing a true cure. The findings reported here suggest that upregulation of ADK is a key component of epileptogenesis. Consequently, therapeutic augmentation of the adenosine system should prevent epileptogenesis. After triggering epileptogenesis in mice by intraamygdaloid KA, we transplanted Adk−/− ES cell–derived NPs into the infrahippocampal cleft and demonstrate that these implants prevent epileptogenesis after acute brain injury. This becomes evident by the lack of spontaneous seizures 3 weeks after cell transplantation, by reduced astroglialosis, and most importantly, by lack of pathogenic upregulation of host ADK. To our knowledge, this is the first demonstration that epileptogenesis can be prevented therapeutically. Further studies are thus warranted to address issues of effective cell dosages, spatial requirements, and long-term effects of adenosine-releasing brain implants. Interestingly, the therapeutic efficacy of adenosine-releasing brain implants seems to depend on ADK expression levels. While the Adk−/− ES cell–derived NPs used here (0% ADK) provided complete protection from epileptogenesis, human mesenchymal stem cell implants with an RNAi-mediated knockdown of ADK (20%–28% of residual ADK) provided only partial protection from acute SE-induced hippocampal injury (42). These findings may suggest a quantitative relation between ADK expression levels, cellular adenosine release, and functional outcome.

The link between expression levels of ADK and susceptibility to epileptogenesis described here allows defining high levels of ADK expression as a risk factor for epileptogenesis. This is of importance, since following an acute brain injury only a small number of patients will develop epilepsy. Thus identification of high ADK expression, e.g., by suitable PET ligands, would allow us to determine those patients at risk for developing epilepsy and would indicate initiation of adenosine-based preventive therapies.

The results presented here define ADK as an attractive target to prevent epileptogenesis. Indeed, ADK inhibitors, such as ABT-702 or related compounds, are effective in the suppression of established or induced seizures; since they can potentiate a site- and event-specific
The present study has identified ADK as a diagnostic marker to prevent potentially novel principle to prevent epileptogenesis. In conclusion, the emergence of spontaneous seizures during epileptogenesis. Thus and EEG recordings commenced, and then a 31-gauge internal cannula was inserted into the lumen of the guide to inject 0.3 μg KA in a volume of 0.2 μl PBS, pH 7.4, into the amygdala. The EEG was monitored for 30 min using a Nervus video–EEG recording device until lorazepam (6 mg/kg) was administered i.v. to terminate seizures. The EEG was further monitored for up to 30 min to ensure seizure cessation. An observer unaware of the experimental treatment performed quantification of EEG records, and the duration of type IV seizure activity was calculated. Selected animals were injected once with 1 mg/kg i.p. 8-cyclopentyl-1,3-dipropylxanthine (DPCPX; Sigma-Aldrich) 30 min prior to intraamygdaloid KA injection. Three weeks after KA injection, a bipolar coated stainless steel electrode (0.20 mm in diameter; Plastics One Inc.) was implanted into the ipsolateral CA3 region and fixed with a pedestal of dental acrylic. Coordinates for the CA3 electrodes (with toothbar at 0) were 2.18 mm caudal to bregma, 2.5 mm lateral to midline, and 2.25 mm ventral to dura. Additional animals received CA1 or cortical recording electrodes. Twenty-four hours after electrode implantation, each animal was subjected to one 8-hour session (from 10 am to 6 pm) of continuous EEG monitoring performed during the light period. An observer unaware of the experimental treatment performed quantification of EEG records.

Cell culture. WT and genetically altered ES cells (Adk<sup>−/−</sup> and Adk<sup>+/+</sup>, respectively) used in these experiments have been described previously (23, 27). NPs were generated from Adk<sup>−/−</sup> and Adk<sup>+/+</sup> ES cells using a well-established stepwise differentiation protocol (46). NPs were routinely cultured on polyornithine-coated dishes in N3 medium as previously described (23). To keep the cells in a proliferative state, the growth factor FGF2 (10 ng/ml) was added daily. After neural differentiation, adenosine release from Adk<sup>−/−</sup> cells amounted to 9.00 ± 4.55 ng adenosine/10<sup>6</sup> cells, while the amount of adenosine released from WT cells (0.33 ± 0.20 ng adenosine/10<sup>6</sup> cells) was minimal (23). Prior to transplantation the cells were labeled with a commercial cell tracer kit (Vybrant CFDA SE Cell Tracker Kit; Invitrogen). The labeling efficiency was about 10%. Cell transplantation. All animals, including control animals, received daily immunosuppression with cyclosporine A (15 mg/kg, i.p.) initiated 2 days prior to cell transplantation, which was performed under anesthesia with 68.5% N<sub>O</sub>2, 30% O<sub>2</sub>, and 1.5% isoflurane 24 h after KA injection. Immediately before transplantation, cells were harvested and resuspended at a concentration of 50,000 cells per 2.5 μl in culture medium. Cell injections (50,000 cells per mouse) were performed using a glass capillary (inner diameter of tip, 70–90 μm). The cells were slowly injected in a volume of 2.5 μl using a drill hole above the left hippocampus and a single diagonal injection tract spanning from coordinate (AP, +1.6; ML, +1.2; DV, 0.0) to coordinate (AP, –2.8; ML, –1.75; DV, –4.0), thus depositing the cells within the infralimbicopamal cleft of the to-be-injured brain hemisphere as described previously (23). Cells were slowly injected (1 μl/min) while withdrawing the capillary (1 mm/min). The capillary was fully retracted 5 min after injection to avoid reflux of cells. Using an identical procedure, sham-treated control animals (n = 6) received culture medium instead of cells.

Histology. Brains obtained 24 h following the KA injection were immediately frozen in 2-methylbutane (–30°C) and sectioned at 12 μm on a cryostat. Coronal sections at the level of bregma –1.7 mm were air dried, then a single diagonal injection tract spanning from coordinate (AP, +1.6; ML, +1.2; DV, 0.0) to coordinate (AP, –2.8; ML, –1.75; DV, –4.0), thus depositing the cells within the infralimbicopamal cleft of the to-be-injured brain hemisphere as described previously (23). Cells were slowly injected (1 μl/min) while withdrawing the capillary (1 mm/min). The capillary was fully retracted 5 min after injection to avoid reflux of cells. Using an identical procedure, sham-treated control animals (n = 6) received culture medium instead of cells.

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hippocampal formation encompassing the complete rostrocaudal extent of the hippocampus, were mounted onto gelatin-coated slides and stained with cresyl violet. For the immunohistochemical detection of GFAP and ADK, we followed our previously published procedures and stained sections adjacent to the cresyl violet–stained sections (17). Further adjacent sections were stained with NFM (Santa Cruz Biotechnology Inc.) and DAPI (Invitrogen). Cellular implants were identified by tracer-derived green fluorescence. GFAP and ADK expression was quantified by analyzing fields of 200 × 300 μm encompassing the entire CA3a region. Corresponding fields from 2 sections from each animal (n = 6 animals per group) were analyzed by scanning ADK immunofluorescence on DAB-stained slices using a Kodak imaging device and by counting individual GFAP-positive cells. Data were analyzed by ANOVA with Student-Newman-Keuls test.

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