Oxidation of KCNB1 Potassium Channels Causes Neurotoxicity and Cognitive Impairment in a Mouse Model of Traumatic Brain Injury

Wei Yu, Randika Parakramaweera, Shavonne Teng, Manasa Gowda, Yashsavi Sharad, Smita Thakker-Varia, Janet Alder, and Federico Sesti
Department of Neuroscience and Cell Biology, Rutgers University, Robert Wood Johnson Medical School, Piscataway, New Jersey 08854

The delayed rectifier potassium (K⁺) channel KCNB1 (Kv2.1), which conducts a major somatodendritic current in cortex and hippocampus, is known to undergo oxidation in the brain, but whether this can cause neurodegeneration and cognitive impairment is not known. Here, we used transgenic mice harboring human KCNB1 wild-type (Tg-WT) or a nonoxidable C73A mutant (Tg-C73A) in cortex and hippocampus to determine whether oxidized KCNB1 channels affect brain function. Animals were subjected to moderate traumatic brain injury (TBI), a condition characterized by extensive oxidative stress. Dasatinib, a Food and Drug Administration–approved inhibitor of Src tyrosine kinases, was used to impinge on the proapoptotic signaling pathway activated by oxidized KCNB1 channels. Thus, typical lesions of brain injury, namely, inflammation (astrocytosis), neurodegeneration, and cell death, were markedly reduced in Tg-C73A and dasatinib-treated non-Tg animals. Accordingly, Tg-C73A mice and non-Tg mice treated with dasatinib exhibited improved behavioral outcomes in motor (rotarod) and cognitive (Morris water maze) assays compared to controls. Moreover, the activity of Src kinases, along with oxidative stress, were significantly diminished in Tg-C73A brains. Together, these data demonstrate that oxidation of KCNB1 channels is a contributing mechanism to cellular and behavioral deficits in vertebrates and suggest a new therapeutic approach to TBI.

Key words: aging; dasatinib; Kv2.1; oxidative stress; ROS; Src kinases

Significance Statement
This study provides the first experimental evidence that oxidation of a K⁺ channel constitutes a mechanism of neuronal and cognitive impairment in vertebrates. Specifically, the interaction of KCNB1 channels with reactive oxygen species plays a major role in the etiology of mouse model of traumatic brain injury (TBI), a condition associated with extensive oxidative stress. In addition, a Food and Drug Administration–approved drug ameliorates the outcome of TBI in mice, by directly impinging on the toxic pathway activated in response to oxidation of the KCNB1 channel. These findings elucidate a basic mechanism of neurotoxicity in vertebrates and might lead to a new therapeutic approach to TBI in humans, which, despite significant efforts, is a condition that remains without effective pharmacological treatments.

Introduction
Ion channels are versatile proteins that generate and modulate electricity across biological membranes. Since electricity is an essential ingredient of life, ion channels are found in all organisms, from prokaryotes to eukaryotes to archaeans, in virtually any type of cell (Hille, 2001). A growing number of ion channels, including the potassium (K⁺) channels, are reported to interact with reactive oxygen species (ROS), either in cell signaling mechanisms or as a side effect of aging and disease (Patel and Sesti, 2016). Hence, oxidative modification of K⁺ channels has the potential to constitute a widespread mechanism of vulnerability but a strong causative link between these modifications and behavioral and functional impairment has still to be established for vertebrates. One channel known to undergo oxidation is the delayed rectifier voltage-gated K⁺ channel KCNB1 (Kv2.1), which carries a major somatodendritic current that regulates high-frequency firing of neurons in the cortex and hippocampus (Mu-
Figure 1. Predicted model of KCNBl-oxidation-mediated apoptosis. Oxidative stress induces the formation of KCNBl oligomers that accumulate in the plasma membrane. The presence of these protein aggregates leads to the activation of Src and downstream JNK kinases. The latter act to destabilize mitochondria, resulting in the release of more ROS (which may further sustain KCNBl oligomerization in a sort of autocatalytic process) and apoptosis.

Materials and Methods
Construction of Tg-WT and Tg-C73A transgenic mice. Transgenic mice were constructed by the Genomic Editing Core Facility at Rutgers using pronuclear injection. cDNA encoding human KCNBl tagged to the human influenza hemagglutinin (HA) tag in the C terminus (Cotella et al., 2012) was inserted in the mouse Thy1.2 cassette using a XhoI restriction site. Constructs were linearized at an EcoRI site, for injection. We obtained 3 Tg-WT and 4 Tg-C73A founders.

Biochemistry. The detailed biochemical procedure was described previously (Cotella et al., 2012). Briefly, frozen, half sagittal brains of either sex were homogenized with a glass tissue grinder in lysis buffer (0.52 mg sucrose, 5 mM Tris-Cl pH 6.8, 0.5 mM EDTA, 1 mM PMSF, and protease inhibitor cocktail set 1, Calibiochem). Samples were centrifuged at 2000 rpm for 10 min, and the supernatant used for biochemical analysis. Protein content was quantified with the Bradford method. Western blots were performed using 5% non-fat milk in 0.05% Tween-20/PBS (PBST) for 2 h at room temperature. After overnight incubation at 4°C with the primary antibody (anti-KCNB1 NeuroMab clone K89/34, UC Davis/NIH; anti-HA H6908, Sigma; anti-actin MAB1501, Millipore), the membrane was washed for 20 min and incubated at room temperature with the appropriate secondary antibody. To detect activated Src tyrosine kinases, brain lysates were incubated at 4°C overnight in the presence of anti-Src antibody (catalog #2108, Cell Signaling Technology). Then, protein A agarose beads (30 µl of 5096 bead slurry) were added and incubated for 2 h at 4°C. Samples were centrifuged for 1 min at 14,000 X g. The pellet was washed five times with cell lysis buffer. The pellet was resuspended with 50 µl 2X SDS PAGE gel and immunoblotted with either anti-Src antibody or anti-P-Src-antibody (catalog #2108, Cell Signaling Technology). The blots were washed in PBST for 20 min and incubated for 5 min with chemiluminescence substrates and exposed. Densitometric analysis was performed using ImageJ (NIH) software.

Lateral fluid percussion injury. All experimental protocols involving animals were approved by the Rutgers University Institutional Animal Care and Use Committee. LFP brain injury involves the displacement of neural tissue by a rapid fluid pulse to the brain and has been described in detail previously (Akker et al., 2011). For surgery, 3-month-old males were anesthetized with 4-5% isoflurane in 100% O2 and placed in a mouse stereotaxic frame. Mice were maintained at 2% isoflurane, and respiration was monitored throughout the procedure. The site of injury was located halfway between lambda and bregma, and between the sagittal suture and the lateral ridge over the right hemisphere. A 3 mm thin plastic disc was fixed with Locite glue (444 Tak Pak, Henkel) onto the skull. Using a trephine (3 mm outer diameter), a craniotomy was performed, keeping the dura intact. A rigid Luer lock needle hub (3 mm
inside diameter) was secured to the skull over the opening with cyanoacrylate adhesive and dental acrylic (Butler Schein). The skull sutures were sealed with the cyanoacrylate to ensure that the fluid bolus from the injury remained within cranial cavity and the hub was filled with saline. After a 60 min period of recovery, the animals were reanesthetized and connected to the fluid percussion injury device (Custom Design and Fabrication, Virginia Commonwealth University) through the Lucite loc fitting of the hub. Once a normal breathing pattern resumed, before sensitivity to stimulation, a ~1.5 atm pulse (~15 ms) was generated through the LFP device. Upon return of the righting reflex (4–10 min for moderate injury), the hub and dental acrylic were removed. The scalp incision was sealed with 3M Vetbond (Thermo Fisher Scientific) and the animals were returned to normal housing conditions.

At this moderate level of injury, ~10% of animals died as a result of the injury within the acute posttraumatic period (15 min), generally from respiratory failure and pulmonary edema (Lishitz, 2003). This is a normal and anticipated feature of the TBI model because it mimics human TBI (Domeniconi et al., 2007). Mice that undergo the surgical procedures but that were uninjured served as the sham controls. Assignment of mice to the LFP or sham group was done in a random manner.

**Drug administration.** Dasatinib (LC Laboratories) was given intraperitoneally at 25 mg/kg. Dasatinib was diluted in vehicle solution (50 mM NaAc, pH 5.0) from a 200 mg/ml stock in dimethyl sulfoxide. Each mouse was subjected to a daily dose of either vehicle or dasatinib solution via an intraperitoneal injection starting from the day of surgery (2 h after injury).

**Behavioral assays.** For the Morris water maze (MWM), mice were acclimated to the paradigm and tested for baseline response using a visible platform test 4 d before injury. The animals were placed in a circular pool of water containing nontoxic white paint and a clear platform for escape. To assess learning, mice were trained using a hidden platform fixed in one of four quadrants for 5 consecutive days starting at 2 d postinjury (dpi; four trials/day). Black and white distal cues were placed on the walls. The quadrant in which the mouse was placed was pseudorandomly varied throughout training, and the time to locate the platform was recorded. Maximum trial time was 60 s, and the mouse remained or was placed on the platform for 15 s and warned for 10 min between trials. To assess memory retention, the day after the last training session (8 dpi), the animals were subjected to a 60 s probe trial with the platform removed and the time spent in the target quadrant was measured (Longhi et al., 2005). Data were recorded using a video-tracking system (EthoVision XT, Noldus Information Technology).

**Vestibulometer rotarod test.** A separate set of mice was used for motor testing. Mice were acclimated to the rotarod device three times per day with 1 h intertrial intervals for the 2 d before the surgery. Balance and motor function were measured on a 36 mm outer diameter rotating rod whose velocity increased from 4 to 40 rpm over a maximum 180 s interval. Each trial ended when the animal fell off the rotarod. At 1, 7, and 21 dpi, each subject underwent three trials a day with 1 h intertrial intervals on the rotarod device. The same mice were used for each time point. The average latency to fall of injured mice was recorded and compared to that of sham mice.

**Immunohistochemistry.** Mice were perfused transcardially with 0.9% saline followed by 4% paraformaldehyde at either 7, 14 and 21 dpi. The brains were cryoprotected in 30% sucrose, and 20 µm frozen sections were prepared throughout the site of injury on the cortex and the hippocampus in a 1:20 series so that the same set of tissue samples could be used for expression of different makers. For activated Caspase-3, KCNB1, and HA immunohistochemistry, sections were pretreated with 0.01 M citrate buffer, and then anticleaved Caspase-3 antibody (1:1000, catalog #9661, Cell Signaling Technology) or anti-KCNB1 antibody (1: 100) or anti-HA antibody (8 µg/ml) was applied overnight, followed by application of the appropriate secondary conjugated antibody. For Fluoro-Jade C (FJC) staining, sections were pretreated with 1% NaOH and 0.06% KMnO4, and then 0.005% Fluoro-Jade C (catalog #AG325, Millipore)/0.0001% was applied for 10 min. For glial fibrillary acidic protein (GFAP), slides were incubated overnight with anti-GFAP antibody at 4°C (1:500, catalog #MAB3402, Millipore). Slides were then incubated in 2° goat anti-mouse antibody (1:500, Alexa Fluor 594). All slides were mounted in Vectashield AntiFade Mounting Medium with DAPI mounting buffer (Vector Laboratories) and stored at 4°C. Staining was visualized on a Zeiss Axiohot microscope at 40× or with a Zeiss Axioscope Image M1 at 100× (see Fig. 2C). Positive cells on the ipsilateral hemisphere were counted in coronal sections representing a 1:20 series throughout the site of injury at the cortex and inclusive of the entire length of the hippocampus. For the cortex, a total of six fields of view at 40× (three most dorsal along the surface of the cortex starting from midline and moving laterally and three just ventral to those fields) were counted on the side ipsilateral to the injury. The ipsilateral hippocampus including CA1 and CA3 as well as the dentate gyrus was used for quantitation of cells in the hippocampus.

**Preparation of hippocampal neural cultures.** The detailed procedure was described previously (Thakker-Varia et al., 2001). Briefly, hippocampi were obtained from time-mated embryonic day 16 (E16) mice killed by CO2 asphyxiation. Hippocampal tissue from individual embryos was mechanically triturated in Neurobasal medium containing B27 (Invitrogen) and glutamine and plated in two 35 mm poly-d-lysine-coated Petri dishes at ~350,000 cells/dish (1.5 ml medium/dish). Cultures were maintained in Neurobasal medium at 37°C in a 95% air/5% CO2 humidified incubator and contained virtually pure neurons. Tail samples from individual embryos were processed for genotyping.

**Electrophysiology.** Data were recorded with an Axopatch 200B amplifier (Molecular Devices), a personal computer (Dell), and Clampex soft-
labeling buffer solution according to the kit’s instructions (Annexin-V-FLUOS staining kit, Sigma). We showed previously that C73A truly acts for subsequent analysis. Experiments were performed blind. The fluorescence of Annexin-V in each image was calculated using ImageJ software (Molecular Devices). Data were filtered at $f = 1$ kHz and sampled at $2.5$ kHz. The bath solution contained the following (in mM): $5$ KCl; $100$ HEPES, pH $7.5$ with NaOH; $1.8$ CaCl$_2$; $1.0$ MgCl$_2$; and $1.0$ MgCl$_2$. Offset potentials due to series resistances (~5 mV) were not compensated for when generating current–voltage relationships. Macroscopic conductances were fitted to the Boltzmann function—voltage relationships (Eq. 1) for non-Tg (circles), Tg-WT (squares), and Tg-C73A (triangles). Data from $n = 15$ (non-Tg), $n = 16$ (Tg-WT), and $n = 16$ (Tg-C73A) neurons, $p = 0.019$ for longitudinal statistical comparisons at $+20$, $+60$, and $+80$ mV (one-way ANOVA). Pairwise comparisons at $+20$, $+60$, and $+80$ mV are all statistically significant except Tg-WT versus Tg-C73A (Tukey’s post hoc test). Differences between Tg-WT and Tg-C73A means are not statistically significant (two-tailed Student’s $t$-test). $N = 3$ experiments.

Representative whole-cell currents elicited in non-Tg, Tg-WT, and Tg-C73A hippocampal neurons by voltage jumps from $-80$ to $+80$ mV in $20$ mV increments. The holding voltage is $-80$ mV. $D$, Mean steady-state current–voltage relationships for non-Tg (circles), Tg-WT (squares), and Tg-C73A (triangles). Data are from $n = 15$ (non-Tg), $n = 18$ (Tg-WT), and $n = 16$ (Tg-C73A) neurons, $p = 0.019$ for longitudinal statistical comparisons at $+20$, $+60$, and $+80$ mV (one-way ANOVA). Pairwise comparisons at $+20$, $+60$, and $+80$ mV are all statistically significant except Tg-WT versus Tg-C73A (Tukey’s post hoc test).

Normalized macroscopic conductance–voltage relationships (Eq. 1) for non-Tg (circles, $n = 15$), Tg-WT (squares, $n = 18$), and Tg-C73A (triangles, $n = 16$). Data were fitted to the Boltzmann function (Eq. 2, solid lines) with $V_{1/2} = 2.1, 9.3$, and $5.4$ mV for non-Tg, Tg-WT, and Tg-C73A, respectively. $p < 0.05$; $**p < 0.01$.

**Results**

**Construction of transgenic mice expressing human KCNB1**

To study oxidation of KCNB1 channels in vivo, we engineered two transgenic mice, expressing respectively, a nonoxidizable variant of human KCNB1 (C73A) and the wild-type channel as a control, both tagged to the HA epitope tag in the C terminus in a B6XCBA background. We showed previously that the addition of the HA tag has no effect on oxidation or other properties of the channel (Cotella et al., 2012; Wu et al., 2013). We refer to these mice simply as Tg-C73A and Tg-WT. To direct expression of hKCNB1 mainly in cortex and hippocampus, we subcloned the channel in the thy1.2 cassette, which represents the standard for expression in these regions of the mouse brain (Aigner et al., 1995). Transgenic mice were constructed using pronuclear injection. We obtained 3 Tg-WT founders and 4 Tg-C73A founders. All transgenic animals heterozygous in hKCNB1 were normal in size and weight ($32.4 \pm 1.0$, $32.1 \pm 0.6$, and $32.6 \pm 1.5$ g, respectively, for 3-month-old non-Tg, Tg-WT, and Tg-C73A mice of either sex; $N = 30$ mice/group) and did not exhibit any apparent phenotype. In contrast, while homozygous Tg-C73A were normal, homozygous Tg-WT developed more slowly and exhibited high mortality rates after LFP injury (58 vs 12% of homozygous Tg-C73A). In what follows, we present results obtained with line #54 (Tg-WT) and line #18 (Tg-C73A), both heterozygous in hKCNB1 (mortality rates after LFP injury, 13 and 9%, respectively).

**Transgenic KCNB1 channels are expressed in cortex and hippocampus**

To confirm that hKCNB1 was expressed in the cortex and hippocampus, coronal frozen sections cut from the brains of 3–month-old animals of either sex were fixed and stained with an antibody against the HA tag (anti-HA) that recognizes only transgenic hKCNB1 and, separately, with an antibody against KCNB1 (anti-KCNB1) that detects both mouse (mKCNB1) and human KCNB1 (hKCNB1) through a conserved C-terminal epitope (HMLPGGAGHGSTRDQSI). As expected, hKCNB1 was detected only in the cortex and hippocampus of transgenic sections, whereas total KCNB1 was detected also in nontransgenic sections (Fig. 2A). Furthermore, we did not detect appreciable KCNB1
expression in other areas of the brain. Staining with either KCNB1 or HA antibody revealed typical cluster distribution of KCNB1 channels in the plasma membrane of both cortical (Fig. 2B) and hippocampal neurons (Fig. 2C; Rhodes et al., 1995; Lim et al., 2000; O'Connell et al., 2006, 2010; Sarmiere et al., 2008; Fox et al., 2013).

Transgenic hKCNB1 channels contribute to outward K⁺ current in hippocampal neurons

To assess the relative amounts of endogenous and transgenic KCNB1 protein and consequently to have a rough measure of the level of overexpression of the latter, half brains were lysed, immunoblotted, and analyzed by densitometry. Results of four experiments with the anti-KCNB1 antibody indicated that the amounts of transgenic channels were comparable in the two transgenic lines and were about half the amounts of endogenous channels (Fig. 3A). When similar experiments were performed with the anti-HA antibody, hKCNB1 was not detected in non-Tg brains, as expected, and its levels were similar in the two transgenic lines, in agreement with the results of the experiments with anti-KCNB1 (Fig. 3B). To assess the contribution of transgenic KCNB1 channels to total neuronal current, we recorded somatic whole-cell currents in primary hippocampal neurons obtained from nontransgenic and transgenic embryos. Voltage jumps ranging from −80 to +80 mV (from an holding voltage of −80 mV) evoked robust outward currents that reversed around −40 mV (Fig. 3C,D; data are shown without subtraction of leak currents). Steady-state current amplitudes at +80 mV were comparable in Tg-WT and Tg-C73A transgenic neurons and were roughly ~40% larger than in non-Tg neurons, consistent with biochemical results (Fig. 3A) and with previous studies that showed that C73A KCNB1 channels conduct like WT KCNB1 channels (Cottrella et al., 2012). Normalized macroscopic conductances (Fig. 3E) fitted to a Boltzmann function exhibited half-activation (Vₚ/₂) values around 0 mV in good agreement with the results of others (Frazzini et al., 2016). The Vₚ/₂ values of transgenic conductances were slightly more positive than those of nontransgenic conductances (Vₚ/₂ = 1.8 ± 0.3, 8.7 ± 2.1, and 5.7 ± 1.1 mV for non-Tg, Tg-WT, and Tg-C73A, respectively), but these differences were not statistically significant.

Overall, these data indicate that the transgenic channels are expressed in neurons of the cortex and hippocampus and that the cyst73 to alan replacement in KCNB1 does not affect the channels' functional attributes or their ability to cluster in the plasma membrane. We cannot rule out that the resolution of our analysis may have missed micro differences in cluster distributions of the transgenic channels. However, since clustering affects KCNB1 conductance (O’Connell et al., 2006, 2010; Fox et al., 2013), considering that there were no differences in total outward K⁺ currents expressed in Tg-WT and Tg-C73A hippocampal neurons, this possibility seems unlikely.

Oxidation of KCNB1 is negligible in Tg-C73A mice

The LFP injury enabled us to expose live animals to conditions of oxidative stress in a reproducible fashion. However, this ap-
proach was valid only if KCNB1 oxidation remained low in the injured Tg-C73A brain. To answer this crucial question we assessed the extent of KCNB1 oxidation in the brains of the various genotypes using Western blot analysis as done previously (Cottella et al., 2012). Oxidized KCNB1 channels in the mouse brain or heterologously expressed in mammalian cells form oligomers held together by disulfide bridges that run with multiple molecular masses ranging from 170 to 500 kDa and that are suppressed by reducing agents DTT and β-mercaptoethanol (Cottella et al., 2012). Indeed, a fraction of total KCNB1 channels from lysates of injured non-Tg brains were run as oligomers with a molecular mass of ~200 kDa in the absence of reducing or denaturing agents (Fig. 4A). This fraction of oligomers was enriched by ~40% following exposure to 1.0 mM hydrogen peroxide (H₂O₂) and was abolished by 20 mM DTT. KCNB1 oligomers were present in lysates of injured Tg-WT and non-Tg brains and in very low amounts in Tg-C73A lysates, but not in uninjured brains (Fig. 4A). Densitometric analysis indicated that the ratio between the oligomeric and monomeric bands (oxidation ratio; Cottella et al., 2012), which gives a measure of the level of KCNB1 oxidation, was moderately increased in Tg-WT mice compared to non-Tg mice (35% increase; Fig. 4C), and most importantly, this ratio was significantly decreased in the brains of Tg-C73A compared to non-Tg mice (70% decrease). Similar oxidation ratios were obtained using the anti-HA antibody, except that no protein was detected in lysates of non-Tg brains, as expected (Fig. 4C).
as done previously (Cotella et al., 2012). In cells transfected with an equal ratio of wild-type and C73A cDNA, the oxidation ratio was roughly one-third that of cells expressing the wild-type channel alone (Fig. 4D), a fraction comparable to that detected in injured Tg-C73A brains. The lack of oligomerization prevents C73A mutant channels heterologously expressed in mammalian cells to induce apoptosis in response to an oxidative challenge (Wu et al., 2013). Since KCNB1 oligomerization was low in Tg-C73A brains, their neurons should be resistant to oxidant-induced apoptosis. Therefore, we challenged cultures of primary hippocampal neurons with DTDP, a potent oxidant, and assessed apoptosis. Representative pictures of cells in control or oxidized and stained with apoptotic marker Annexin-V are shown in Figure 5. Thus, in agreement with previous studies (Redman et al., 2007), non-Tg neurons were susceptible to DTDP-induced apoptosis. However, while Tg-WT neurons exhibited levels of apoptosis comparable to those of non-Tg cells, Tg-C73A neurons were significantly less susceptible. Together, these data indicated that oxidation of KCNB1 and the toxic effects associated with it are low in the brains of Tg-C73A mice.

**Tg-C73A mice perform better than non-Tg and Tg-WT mice on the rotarod.**

To determine the impact of KCNB1 channel oxidation on the function of the brain, we tested the animals on the accelerating rotarod device (rotarod) and the MWM, which respectively provide sensitive measures of the function of the motor and sensory cortex and the hippocampus (Smith et al., 1995; Laurer and McIntosh, 1999), the areas of the brain most affected by the injury in our LFP model (the injury occurs at the motor and sensory cortex, which is overlying the hippocampus; Alver et al., 2011). Two days before undergoing surgery, the mice were exposed to the rotarod paradigm. This procedure served the double purpose of allowing the animals to acclimatize to the new protocol and allowing us to identify impaired animals or detect differences due to genotype and normalize data to baseline response. However, all mice behaved similarly during pretesting before injury, indicating that genotype did not affect baseline motor function (Fig. 6A). In contrast, mice subjected to the LFP injury had a shorter latency to fall relative to their shams, and most importantly, Tg-C73A injured mice could remain longer on the rotating cylinder than Tg-WT or non-Tg mice (Fig. 6B). There were no significant differences between the latencies of the shams (Fig. 6C), strengthening the notion that in the absence of events that trigger the release of ROS, the transgenic mice are normal at baseline. In all groups of mice, latencies reached a peak around the first week after injury and then either remained stable or, as in the case of Tg-C73A, moderately decreased. This trend probably reflects the fact that the animals were getting acquainted to the device (in fact, latencies were shorter presurgery, probably because the animals experienced the device for the first time; Fig. 6A). We and others have observed this behavior in previous stud-
Table 1. Cortical sections stainings

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<th>Injured</th>
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<tr>
<td></td>
<td>non-Tg</td>
<td>Tg-WT</td>
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<tr>
<td>FLJC</td>
<td>72.4 ± 3.3 (9)</td>
<td>88.5 ± 2.1 (9)</td>
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<tr>
<td>Caspase-3</td>
<td>24.4 ± 0.6 (18)</td>
<td>30.6 ± 0.9 (18)</td>
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<td>Vehicle</td>
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<tr>
<td>GfAP</td>
<td>74.6 ± 2.4 (18)</td>
<td>37.0 ± 1.1 (18)</td>
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<tr>
<td>FLJC</td>
<td>61.9 ± 16 (12)</td>
<td>31.8 ± 12 (12)</td>
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<td>Datasinib</td>
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Mean number of cells per cortical section, positive for the indicated markers at 7 dpi. The number of sections (maximum number of sections/brain, 6) is indicated in parentheses. For FLJC, top row, p < 0.0001, except for sham non-Tg versus sham Tg-WT and sham non-Tg versus sham Tg-C73A (not statistically significant). For caspase-3, p < 0.0001, except for sham non-Tg versus sham Tg-WT, sham non-Tg versus sham Tg-C73A, and sham Tg-WT versus sham Tg-C73A (not statistically significant). For GfAP, bottom row, p < 0.0001, and all pairwise comparisons are statistically significant (one-way ANOVA/Tukey’s post hoc test).

Tg-C73A mice perform better than non-Tg and Tg-WT mice in the Morris water maze

Spatial learning and memory were assessed in the MWM using independent groups of mice. As observed in the rotarod test, all mice behaved similarly in the presurgery trial using a visible platform, and therefore baseline adjustments were not performed (Fig. 6D). During the training period after surgery (Fig. 6E) and in the test of memory consolidation that was administered the day after the last training (Fig. 6G), injured Tg-C73A mice performed significantly better than non-Tg and Tg-WT, with the latter being the most cognitively impaired. Indeed, injured Tg-C73A animals did not exhibit any appreciable loss of cognitive function in the MWM (Fig. 6F, compare injured mice and shams). To exclude (or quantify) that impaired motor function could have affected the ability of the animals to reach the platform, we measured swimming speeds, which were comparable in the various groups of mice, suggesting that the improved outcome in the Tg-C73A mice was not due to increased activity (Fig. 6H).

Typical histological lesions of TBI are reduced in cortex and hippocampus of Tg-C73A mice

Inflammation, disruption of axonal transport followed by axonal swelling, and finally degeneration and neuronal death are three hallmark lesions of TBI that affect many processes including motor function, memory, and spatial learning (Smith et al., 1991; Miyazaki et al., 1992; Hamm et al., 1993; Hicks et al., 1993; Dixon et al., 1999; Royo et al., 2003; Johnson et al., 2013; DeKosky et al., 2013). Therefore, we compared the extent of these lesions in corona sections of the region of injury throughout the cortex and hippocampus that we stained against glial fibrillary acidic protein (cortex, Fig. 7; hippocampus, Fig. 8A), which, being a marker for glia (astrocytes) and since gliosis accompanies inflammation, provides indirect evidence for inflammation (Jacque et al., 1978; Vos et al., 2010; Flturo-Jade C (FLJC, hippocampus, Fig. 8B; cortex, Table 1), a marker for neurodegeneration (Schueme et al., 1997); and activated caspase-3 (hippocampus, Fig. 8C; cortex, Table 1), a marker for neuronal apoptosis (caspase-3 staining colocalizes with NeuN but not GFAP; Beer et al., 2000; Clark et al., 2000; McEwen and Springer, 2005; Alder et al., 2013). Representative images of GFAP staining in cortical sections of the various genotypes at 7 and 21 dpi along with quantitative analyses are shown in Figure 7. Thus, consistent with the course of a traumatic event, GFAP staining revealed the onset of an inflammatory process, remitting over time that was high in Tg-WT and

Figure 9. Datasinib improves behavioral outcome in LFP-injured mice. A, Latency to fall from the rotating rod of non-Tg injured mice injected intraperitoneally daily with 25 mg/kg dasatinib (diamonds) or vehicle (circles). p = 0.0019 for statistical comparisons at each individual dpi (one-taled Student’s t test). In five experiments, N = 8 animals for vehicle- and dasatinib-injected mice, respectively. B, Latency to fall from the rotating rod of sham non-Tg mice injected with dasatinib (diamonds) or vehicle (circles). Differences between means are not statistically significant (one-tailed Student’s t test). In five experiments, N = 8 and 7 animals for vehicle- and dasatinib-injected mice, respectively. C, Latency to reach the platform of injured non-Tg mice injected with dasatinib (diamonds) or vehicle (circles). Where indicated, p = 0.0045 for statistical comparisons at each individual dpi (one-tailed Student’s t test). In five experiments, N = 9 and 9 animals for vehicle- and dasatinib-injected mice, respectively. D, Latency to reach the platform of sham non-Tg mice treated with dasatinib (diamonds) or vehicle (circles). The dotted line indicates sham mice treated with dasatinib. Differences between means are not statistically significant (one-tailed Student’s t test). In five experiments, N = 8 and 7 animals for vehicle- and dasatinib-injected mice, respectively. E, Consolidated memory retention test for the indicated groups of mice. p = 0.0025 (one-tailed Student’s t test). In five experiments, N = 8 and 9 animals for vehicle- and dasatinib-injected mice, respectively, and N = 8 and 6 for their respective sham. F, Mean swimming speeds of injured (filled symbols) and sham (hollow symbols) mice treated with dasatinib (diamonds) or vehicle (circles). Differences between means are not statistically significant (one-way ANOVA). In five experiments, N = 9 and 9 animals for vehicle- and dasatinib-injected mice, respectively, and N = 8 and 6 for their respective sham. p < 0.05, ***p < 0.001.
low in Tg-C73A sections. Results of immunostaining in hippocampal sections at 7 dpi are summarized in Figure 8 and in cortical sections in Table 1. In all cases, damage was maximal in Tg-WT and minimal in Tg-C73A brains underscoring the existence of a causative relationship between KCNB1 oxidation, tissue destruction, and behavioral deficit following the LFP injury. Thus, where KCNB1 oxidation was exacerbated, neuronal damage was increased and the outcome of TBI was severe, whereas where KCNB1 oxidation was suppressed, the outcome of TBI, along with tissue damage, were improved. We further notice that inflammation, neurodegeneration, and apoptosis were moderately more pronounced in the sham brains of Tg-WT compared to non-Tg and Tg-C73A mice. This may indicate moderate oxidation in the shams, probably due to the surgery, or, alternatively, the existence of KCNB1-independent mechanisms that contribute to tissue damage. We conclude that oxidation of KCNB1 channels resulting from TBI is an event that contributes toward tissue damage and impacts behavioral outcomes.

Mice treated with dasatinib perform better than vehicle mice in the rotarod
Previous in vitro studies showed that oxidation of hKCNB1 channels favors the activation of Src tyrosine kinases, which in turn initiate an apoptotic cascade (Fig. 1; Wu et al., 2013). Therefore, we sought to determine whether pharmacological strategies that act to inhibit the activities of Src kinases could improve neuronal lesions and behavioral outcome of LFP-injured animals. To this end we used dasatinib, an FDA-approved Src kinases inhibitor that is blood–brain barrier permeable and pharmacologically active in the brain (Porkka et al., 2008; Agarwal et al., 2012). Mice injected with dasatinib (25 mg/kg daily starting 2 h after surgery; Luo et al., 2006; Hasima and Aggarwal, 2012; Katsumata et al., 2012) or vehicle and their respective shams were subjected to the rotarod and the MWM protocols, and the results of these experiments are illustrated in Figure 9. Thus, dasatinib significantly increased latency to fall at all days after injury relative to vehicle-treated mice (Fig. 9A) and had no effect on sham animals (Fig. 9B).

Mice treated with dasatinib performed better than vehicle mice in the Morris water maze
During the 6 d training period (Figs. 9C,D) and in the memory retention test the following day (Fig. 9E), injured mice injected with dasatinib performed significantly better than vehicle mice. In fact, there was no significant difference in the behavioral responses of injured mice treated with the drug and the shams (Fig. 9D). Mean swimming speeds, displayed in Figure 9F, were similar in all groups, excluding that differences in the fitness of the animals might have impacted the outcome of the tests.

Histological lesions of TBI are reduced in the brains of mice treated with dasatinib
Cortical and hippocampal sections were stained with anti-caspase-3 and GFAP and FLJC (Fig. 11, Table 1). As expected, dasatinib treatment resulted in a marked decrease of the number of cells positive for the various markers compared to vehicle in both cortical (Fig. 10, Table 1) and hippocampal sections (Fig. 11). The number of positive cells to any marker was moderately higher in sham animals injected with vehicle compared to those injected with dasatinib (albeit these differences were generally not statistically significant). The protective effect of the drug in sham animals may reflect baseline oxidation of KCNB1 channels or, alternatively, inflammatory processes associated with the surgery. We conclude that inhibition of the activity of Src tyrosine kinases significantly ameliorates typical lesions of TBI, leading to improved motor function and spatial memory during the critical phase of the LFP injury.

Oxidation of KCNB1 channels is associated with Src kinases activity
To determine whether oxidation of KCNB1 channels was responsible for the activation of the Src kinases following the LFP injury and therefore link the effect of dasatinib on recovery post TBI to KCNB1 oxidation, we biochemically assessed the fraction of activated Src kinases in the brains of our mice using an antiphosphorylated Src family antibody that detects phosphorylation status of tyr416, a residue conserved in all members of the Src kinase family (Konig et al., 2008). Representative immunoblots of total and activated (phosphorylated) Src proteins in the brains of mice injected with vehicle or dasatinib and separately in the brains of non-Tg, Tg-WT, and Tg-C73A animals, along with densitometric analyses, are illustrated in Figure 12. Src activity was
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Figure 11. Typical lesions of TBI are ameliorated in the hippocampi of mice treated with dasatinib. A, Mean number of cells positive for GFAP per hippocampal section of the indicated groups of mice at 7 dpi. Each single mean was calculated from 18 sections (3 brains, 2 fields of view/section). B, Mean number of cells positive to FLJ4 per hippocampal section of the indicated groups of mice at 7 dpi. Each single mean was calculated from 12–18 sections (2–3 brains, 2 fields of view/section). C, Mean number of cells positive to anti-caspase-3 per hippocampal section of the indicated groups of mice at 7 dpi. Each single mean was calculated from 12 sections (2 brains, 2 fields of view/section). In A–C, p < 0.0001 with pairwise comparisons all statistically significant except sham vehicle versus sham dasatinib (one-way ANOVA/Tukey’s post hoc test). **p < 0.01.

Discussion

To determine the role that oxidation of KCNB1 channels has for the function of the brain, we developed a transgenic mouse expressing a KCNB1 mutant (C73A) resistant to redox and pursued a pharmacological approach that directly impinges on the apoptotic pathway activated in response to oxidation of the channel. We found that decreasing the amount of oxidizable KCNB1 channels by genetic means is strongly protective in a mouse model of moderate TBI, a condition characterized by high oxidative stress. Thus, neuronal damage induced by the LFP injury was markedly reduced in Tg-C73A mice, and consequently, the animals exhibited improved behavioral outcome after TBI. Moreover, the detrimental effects of the neurotoxic pathway activated in response to KCNB1 oxidation could be neutralized by dasatinib, which ameliorated the devastating effects of TBI during both the primary and secondary injury processes. Furthermore, Src kinase activity was significantly depressed in the brains of Tg-C73A mice, and, vice versa, the drug was associated with small amounts of oxidized channels. Overall, these findings demonstrate that oxidation of KCNB1 channels represents a mechanism of cognitive and functional impairment in vertebrates and further validate a molecular model for its toxicity that emerged from in vitro evidence. Despite a large volume of research, no medication has been proven effective for the treatment of TBI in humans. Protein kinases are one of the most investigated drug targets by the pharmaceutical industry, but the development of kinase-based therapies for brain diseases remains a challenge (Chico et al., 2009). Thus these findings not only underscore the pathological nature of oxidation of KCNB1 channels in TBI, but they may further suggest a new...
therapeutic approach for a condition that affects millions worldwide.

One general problem with transgenic approaches is the potential side effects of protein overexpression. The mice used in this investigation were normal in size and weight, developed normally, did not present any tissue damage, did not show any apparent phenotype, and, limited to the cognitive abilities assessed in this study, they were normal. Thus, it appears that a moderate KCNB1 gain of function is well tolerated in the brain even though we cannot completely exclude that the transgenic mice may have developed some sort of cognitive decompensation that went undetected in our investigations. These findings also argue against the idea that augmented KCNB1 current is proapoptotic, as suggested by some in vitro studies (for review, see Sesti et al., 2014). Moreover, since the N terminus and the C terminus of the channel physically interact during the channel’s activation (Ju et al., 2003; Kobilinsky et al., 2006) and oligomerization probably links the two termini together through disulfide bridges, it is possible that the antiapoptotic effect of some KCNB1 inhibitors (Liu et al., 2013; Zhou et al., 2016) stems from their ability to prevent oligomerization rather than conduction, or, alternatively, as in the case of heme oxygenase-1 in in vitro models of Alzheimer’s disease, through interfering with KCNB1 regulatory pathways (Het-tiarachchi et al., 2014). In contrast, the effects of the LFP injury were markedly aggravated in Tg-WT mice compared to non-Tg mice, suggesting that the increased amount of oxidized KCNB1 channels in the former is proportional to the extent of the neuronal damage and behavioral deficits. Specia et al. (2014) have shown that in KCNB1 KO mice the absence of KCNB1 current causes hyperexcitability, which in turn correlates with cognitive impairment and susceptibility to seizure. Oxidized KCNB1 channels do not conduct current (Cotella et al., 2012; Frazzini et al., 2016), and this may explain the cognitive impairment of LFP-injured mice. On the other hand, the potent effect of dasatinib would argue that neuronal damage is the underlying cause, but it must be considered that inhibition of Src tyrosine kinases results in lower levels of oxidized—and therefore nonconducting— KCNB1 channels. Thus, it is likely that decreased current and apoptotic stimuli both contribute to the neuronal and behavioral deficits observed in injured mice, and future investigations will dissect the individual contribution of each to TBI.

Oxidation of a K+ channel as a mechanism of neuronal vulnerability was initially demonstrated in Caenorhabditis elegans (Cai and Sesti, 2009). The channel that undergoes oxidation in the worm, KVS-1, is a homolog of KCNB1 (Rojas et al., 2008), and in both channels the effects of oxidation are mediated by conserved cysteine residues, cys113 in KVS-1 and cys73 in KCNB1. The findings reported here that show that oxidation of K+ channels contribute to cognitive impairment underscore the high degree of conservation of this mechanism of neuronal vulnerability and further broaden its potential relevance. For example, it is well established that the aging hippocampus of rodents develops hyperexcitability (Landfield et al., 1986; Barnes et al., 1987; Barnes, 1994; Papanotopoulos and Kostopoulos, 1996), and it may not be coincidental that KCNB1 undergoes oxidation in the brains of naturally aging mice (Cotella et al., 2012). Moreover, KCNB1 channels are oxidized in hippocampal neurons of the 3x-Tg-AD mouse where they promote hyperexcitability (Frazzini et al., 2016), while inhibition of Src kinases attenuates microgliosis in BV2 murine cells incubated with β-amyloid oligomers (Dhawan and Combs, 2012). In summary, the evidence presented here would argue that oxidation of KCNB1 channels is a mechanism that contributes to cognitive deficit in normal aging as well as in Alzheimer’s disease—albeit to different extents—and by analogy, that dasatinib may represent a valid model of therapeutic intervention in neurodegenerative disease.

References


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