Role of microglial IKKβ in kainic acid-induced hippocampal neuronal cell death

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Microglial cells are activated during excitotoxin-induced neurodegeneration. However, the in vivo role of microglia activation in neurodegeneration has not yet been fully elucidated. To this end, we used Ikkβ conditional knockout mice (LysM-Cre/ikkbF/F) in which the Ikkβ gene is specifically deleted in cells of myeloid lineage, including microglia, in the CNS. This deletion reduced IxB kinase (IKK) activity in cultured primary microglia by up to 40% compared with wild-type (IkkβF/F), and lipopolysaccharide-induced proinflammatory gene expression was also compromised. Kainic acid (KA)-induced hippocampal neuronal cell death was reduced by 30% in LysM-Cre/ikkbF/F mice compared with wild-type mice. Reduced neuronal cell death was accompanied by decreased KA-induced glial cell activation and subsequent expression of proinflammatory genes such as tumour necrosis factor (TNF)-α and interleukin (IL)-1β. Similarly, neurons in organotypic hippocampal slice cultures (OHSCs) from LysM-Cre/ikkbF/F mouse brain were less susceptible to KA-induced excitotoxicity compared with wild-type OHSCs, due in part to decreased TNF-α and IL-1β expression. Based on these data, we concluded that IKK/nuclear factor-κB dependent microglia activation contributes to KA-induced hippocampal neuronal cell death in vivo through induction of inflammatory mediators.

Keywords: excitotoxicity; hippocampus; IKKβ; kainic acid; microglia

Abbreviations: CA1 = cornu ammonis 1; CA3 = cornu ammonis 3; CD11b = cluster of differentiation molecule 11b; GFAP = glial fibrillary acidic protein; HMGB-1 = high-mobility group box-1; Iba-1 = ionized calcium binding adaptor molecule-1; IL-1β = interleukin-1β; IKK = IκB kinase; IR = immunoreactive; KA = kainic acid; LPS = lipopolysaccharide; MCAO = middle cerebral artery occlusion; NF-κB = nuclear factor-kappa B; NG2 = neuron-glial antigen 2; OHSCs = organotypic hippocampal slice cultures; PI = propidium iodide; PMΦ = peritoneal macrophages; TLR = toll-like receptor; TNF-α = tumour necrosis factor-α; tPA = tissue plasminogen activator

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Introduction

Excitotoxicity is a mechanism that contributes to neuronal cell death following acute neuronal damage, including traumatic brain injury and stroke, and is implicated in chronic neurodegenerative diseases (Doble, 1999). It is well known that over-stimulation of the glutamate receptor is responsible for excitotoxic neuronal cell death (Doble, 1999). In this process, activated microglia are easily detected in and around the regions encompassing dying neurons (Andersson et al., 1991b). Several studies also
suggest that activation of microglia may contribute to excitotoxic-induced neuronal cell death. Upon excitotoxic brain injury, proinflammatory cytokines are expressed by activated microglia (Barone and Feuerstein, 1999). In another study, inhibition of microglial activation and proliferation by a chemical inhibitor reduced excitotoxic spinal cord neuronal cell death (Tikka et al., 2001). The critical role of microglial production of tissue plasminogen activator (tPA) in excitotoxic-induced hippocampal neuronal cell death was also documented in a study using tPA-deficient mice (Tsirka et al., 1995). These previous reports support a model in which excitotoxic brain damage induces microglial cell activation, thereby augmenting neuronal cell death. However, it has also been argued that microglia protect hippocampal neurons from excitotoxic cell death (Bruce et al., 1996; Neumann et al., 2006; Simard and Rivest, 2007). In these studies, inhibition of microglial activation by glucocorticoid-activated kainic acid (KA)-mediated excitotoxicity, and tumour necrosis factor (TNF)-α, secreted by activated glial cells during excitotoxic brain damage, was found to protect hippocampal neurons from oxidative stress. Additionally, activated microglia have been shown to release neurotrophic factors, which promote neuronal survival against excitotoxic neuronal damage (Elkabes et al., 1996; Young et al., 1999). Therefore, it is conceivable that microglia are activated to protect neurons during excitotoxic brain damage. Thus, the in vivo role of microglial cell activation in excitotoxic neuronal cell death is still debatable. Nonetheless, in light of the in vitro neurotoxic effects of activated microglial cells, inflammatory microglial activation is regarded as an attractive therapeutic target for the treatment of various neurological disorders that accompany excitotoxic neuronal cell death (Block et al., 2007). In this regard, it is of critical importance to elucidate the in vivo role of microglial activation, as inhibitors of microglial activation could turn out to exacerbate brain damage if microglia activation in fact plays a neuroprotective role in vivo.

It is well known that nuclear factor-κB (NF-κB) activation plays a critical role in the microglial production of proinflammatory genes including TNF-α, interleukin-1β (IL-1β) and inducible nitric oxide synthase (iNOS) (Jana et al., 2001; Rasley et al., 2002; Moriyama et al., 2006). Upon stimulation, NF-κB is activated by IκB kinase (IKK) complex, in a manner dependent mainly on the IκB catalytic subunit (Karín, 1999). It was previously documented that NF-κB is activated in microglia in excitotoxic brain injury (Matsuoka et al., 1999; Acarin et al., 2000). This may account for inflammatory cytokine expression by microglia during excitotoxic neurodegeneration. Therefore, we reasoned that by deleting the IκB gene in microglial cells, we might be able to inhibit inflammatory microglia activation. Further, by using these mice in an excitotoxic brain injury model, we can address the in vivo role of microglia activation in excitotoxin-induced neuronal cell death. We tested this hypothesis by using LysM-Cre/IκBβ-floxed (LysM-Cre/IκBβΔ/Δ) mice, in which the IκB gene was specifically deleted in cells of myeloid origin, including the microglia in the CNS (Greten et al., 2004).

Materials and Methods

Animals and genotyping

Myeloid cell type-specific IκBβ-deficient (LysM-Cre/IκBβΔ/Δ) mice were generated by breeding IκBβ-floxed (IκBβΔ/+) mice and LysM-Cre knock-in mice expressing Cre under the control of endogenous lysozyme M promoter as previously described (Claussen et al., 1999; Li et al., 2003). PCR genotyping was performed using primers, 5′-TGA CCC GGG AAT GAA TAG GA-3′ and 5′-GTC TTC AAC CTC CCA AGC CTT-3′, which amplify both the IκBβ(+) (220 bp) and IκBβΔ (310 bp) alleles. LysM-Cre mice were genotyped by PCR using the primer pair NLS-Cre (5′-CCC AGA AAT GCC AGA GTG TCC-3′) and Cre8 (5′-CCC AGA AAT GCC AGA TTA CG-3′). Mice were housed at 23 ± 2°C with a 12 h light–dark cycle and food and water ad libitum. All surgical and experimental procedures were reviewed and approved by the Institutional Animal Care and Use Committee (IACUC) of Seoul National University.

Primary glial culture from neonates and cortical neuron culture

Primary microglia cultures were prepared as previously described (Lee et al., 2000). Briefly, mixed glial cultures were prepared from postnatal day 1–3 wild-type and LysM-Cre/IκBβΔ/Δ mice. After removing meninges from the cerebral hemispheres, tissue was dissociated into a single-cell suspension by gentle trituration. Cells were cultured in glial culture media (DMEM supplemented with 10 mM HEPES, 10% FBS, 2 mM l-glutamine and 1× antibiotic/antimyocytic) in 75 cm2 flasks at 37°C in a 5% CO2 incubator and the medium was changed every 5 days. Microglia were harvested from mixed glial cultures on day 14. After shaking at 200 r.p.m. for 4 h on an orbital shaker, the media from the cultures was collected and centrifuged at 800g for 10 min. Microglia were plated in glial culture media. After 30 min, dishes were washed with medium to remove unattached astrocytes. The purity of microglia was routinely monitored and was >98% as determined by histochemical staining with cluster of differentiation molecule 4 staining for 4 h on an orbital shaker, the media from the cultures was collected and centrifuged at 800g for 10 min. Microglia were plated in glial culture media. After 30 min, dishes were washed with medium to remove unattached astrocytes. The purity of microglia was routinely monitored and was >98% as determined by histochemical staining with cluster of differentiation molecule 4 staining. Purity of microglia was routinely monitored and was >98% as determined by histochemical staining with cluster of differentiation molecule 4 staining. The purity of microglia was routinely monitored and was >98% as determined by histochemical staining with cluster of differentiation molecule 4 staining. Purity of microglia was routinely monitored and was >98% as determined by histochemical staining with cluster of differentiation molecule 4 staining.

Determination of loss of the IκBβ allele at the genomic level by real-time PCR

Genomic DNA (100 ng in 4 μl) was prepared from each sample, and mixed with SYBR Green PCR Master Mix (10 μl, Applied Biosystems, Foster City, CA, USA), primers (1 μl at 10 μM each) and H2O (5 μl). Real-time PCR was performed for 40 cycles of 95°C for 15 s and 60°C for 1 min using an ABI 7500 Real Time
PCR System (Applied Biosystems, CA, USA). Primers, 5′-AAG
ATG GGC AAA CTG TGA TGT G-3′ and 5′-CAT ACA GGC ATC
CTG CAG AAC A-3′, were used to amplify the Ikkβ allele, and
primers, 5′-GGT GCA TGG TGT GTG AAG AC-3′ and 5′-CAT
GCA TAC TAG CGC CAC AC-3′, were used to amplify the Tnfr1
gene as a control. The ratio of Ikkβ and Ikkβ signal was
calculated after normalization to the Tnfr1 signal.

Real-time RT–PCR

Real-time RT–PCR was performed using SYBR Green PCR Master
Mix as previously described (Lee et al., 2004). Reactions were
performed in duplicate in a total volume of 10 μl, each containing
10 pM primer, 4 μl cDNA and 5 μl SYBR Green PCR Master Mix.
The mRNA levels of each target gene were normalized to that of
GAPDH mRNA. Fold-induction was calculated using the 2−ΔΔCt
method as previously described (Livak and Schmittgen, 2001). All
real-time RT–PCR experiments were performed at least three
times, and the mean ± SEM values have been presented unless
otherwise noted. The primer sequence information can be found
in the Supplementary materials.

Stereotaxic injection and tissue processing

For intracerebroventricular (i.c.v.) injection of KA, 8- to 12-week-
old male wild-type and LysM-Cre/IkkβF/F mice (22–25 g) were
anesthetized by pentobarbital sodium (30 mg/kg, body weight, i.p.)
and placed on a stereotaxic apparatus (Mneyurolab, MO, USA).
Animals were injected with PBS or KA (0.2 μg in 4.0 μl of PBS)
at the speed of 0.5 μl/min into the right ventricle using a 26-G needle
(stereotaxic coordinates in millimetre with reference to the
bregma: AP, −2.0; ML, −2.9; DV, −3.8). After 5 min, the
needle was removed with three intermediate steps over 3 min to
minimize backflow, and the incision was cleaned with saline and
sutured. Animals were kept on a warm pad until recovery. On
either day 1 or day 3 after surgery, brains were removed from the
mice after perfusion, immersed for 12 h in 4% PFA fixative at 4°C
and serially cryoprotected in 10, 20 and 30% sucrose in PBS for
48 h at 4°C. Serial coronal sections (30 μm thickness) were cut on
a cryostat and collected as free-floating sections in PBS. Sections
were stored at −20°C until needed for histochemical studies.

Evaluating neuronal damage

For Nissl staining, hippocampal tissue sections were mounted on
gelatin-coated slides, dried for 1 day at RT and stained with 0.5%
cresyl violet. The numbers of cornu ammonis (CA) 1 and 3
neurons were counted at three levels of the dorsal hippocampus.
Specifically, alternate sections were obtained at 1.6, 1.9 and
2.2 mm posterior to the bregma, and two regions from each level
(six regions for each animal) were used to count cells in the CA1
region. The number of intact neurons within the CA1 layer was
counted using a light microscope (BX51, Olympus, Japan) at
400× magnification and expressed as the number of CA1 neurons
per millimeter of linear length as described previously (Choi et al.,
2005). To maintain consistency across animals, a rectangular box
(1.0 × 0.25 mm) was centred over the CA1 cell layer beginning
1.0 mm lateral to the midline and 0.5 mm medial to the CA2
subfield. Only neurons with normal visible nuclei were counted.
The mean number of CA1 neurons per millimeter of linear length
of the ipsilateral hemispheres was calculated for each treatment
group. The number of CA3 pyramidal neurons was also counted
under light microscope as described previously (Hernandez-
Sanchez et al., 2001). Cell counts were made in a defined area
(1.0 mm × 0.75 mm) centred over the CA3 region using three
sections per brain. All assessments of histological sections were
blindly performed.

Immunohistochemistry and quantitative analysis

Immunohistochemical analysis was performed as previously
described (Park et al., 2006), and detailed methods can be found
in the Supplementary materials. To perform quantitative
analysis of CD11b and GFAP immunostaining, 3–4 sections per
animal were selected and images were captured and analysed using
MetaMorph software (Universal Imaging, PA, USA). One or two
fields (200 μm × 200 μm or 100 μm × 100 μm) in each slide within
the midpoint of hippocampal CA1 and CA3 regions encompassing
all layers were selected for quantification, and the intensity of
CD11b and GFAP immunoreactivity was evaluated by means of
a relative optical density value. To quantify CD11b+/NG2+ cells,
images were captured under a confocal laser scanning microscopy.
The entire quantifying procedure was blindly performed.

Organotypic hippocampal slice cultures

Organotypic hippocampal slice cultures (OHSCs) were prepared
and maintained as described previously (Jung et al., 2004). After
14 days in culture, slices were exposed to 50 μM KA following the
previously reported protocol (Kristensen et al., 2001). Neuronal
degeneration was quantified by the uptake of propidium iodide
(P1) into the damaged cells (Macklis and Madison, 1990). To
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Since microglia belong to the myeloid lineage, have characteristics of macrophages/monocytes and constitutively express the lysozyme gene (Perry et al., 1985; Zucker-Franklin et al., 1987; Hao et al., 1991), we tested whether Ikkβ is deleted in microglial cells of LysM-Cre/IkkβF/F mice. First, we cultured primary microglial cells from neonatal LysM-Cre/IkkβF/F mice and measured the Ikkβ deletion rate at the genomic level with real-time PCR using a specific primer set designed to detect undeleted-Ikkβ alleles, as previously described (Li et al., 2003). In cultured primary microglia of LysM-Cre/IkkβF/F mice, 36 ± 3% of the Ikkβ alleles in the population were deleted, while deletion of Ikkβ was not detected in primary cortical neurons (Fig. 1B). In primary cultured astrocytes, the deletion frequency was only 3 ± 1%, probably due to contamination of the astrocyte culture with microglia (Fig. 1B). We then tested whether deletion of Ikkβ in microglia correlates with reduced IKK activity using an in vitro kinase assay. Lipopolysaccharide (LPS)-stimulated IKK activity in primary microglial cells from LysM-Cre/IkkβF/F mice was decreased by 40% compared with IKK activity in wild-type (IkkβF/F) microglia (Fig. 1C). These results suggest that ~40% of the microglial cells lost Ikkβ alleles. Similarly, LPS-induced expression of proinflammatory genes such as TNF-α, IL-1β, iNOS, intercellular adhesion molecule-1 (ICAM-1) and vascular cell adhesion molecule-1 (VCAM-1) was decreased by 30–60% (Fig. 1D). These data demonstrate that cultured microglial cells from LysM-Cre/IkkβF/F mice are less responsive to LPS stimulation. Since microglial cell cultures are derived from neonates, it is possible that the deletion rate may not represent the Ikkβ deletion rate in adult mice in vivo. To test this, we harvested microglial cells directly from adult mice using a previously reported procedure (Slepko and Levi, 1996) and assessed the Ikkβ deletion rate. In microglia directly isolated from 8-week-old adult LysM-Cre/IkkβF/F cerebella, only 4% of the Ikkβ alleles were deleted, although we observed 71 ± 2% Ikkβ deletion in PMΦ (Fig. 1E), which is consistent with a previous study (Greten et al., 2004). The Ikkβ deletion rate in the microglia population directly isolated from neonate mice cerebella was ~20% (data not shown), which is higher than the deletion rate in adult mouse microglia in vivo, but still lower than the rate in primary cultured microglia. These results show that Ikkβ is more easily deleted in primary cultured microglia, which are in a more activated state than microglia in vivo (Eder et al., 1999; Hurley et al., 1999), and implies that the in vivo Ikkβ deletion rate may be increased upon microglia activation. To test this, we stimulated LysM-Cre/IkkβF/F microglia in vivo by i.c.v. LPS injection. One day after LPS injection, in microglia isolated from the whole cerebrum, the Ikkβ deletion rate increased to 31 ± 2% and injection of KA increased the Ikkβ deletion rate to 20 ± 3% (Fig. 1E). In microglia isolated from the hippocampus, where KA-induced microglial activation is most prominent, the deletion rate increased to 73 ± 2% (Fig. 1E). Taken together, these data show that KA-injection enhances microglia-specific Ikkβ deletion in LysM-Cre/IkkβF/F mice, and suggest that microglia in LysM-Cre/IkkβF/F mice are less responsive to inflammatory or stress stimulation.

**KA-induced death of hippocampal neurons is reduced in LysM-Cre/IkkβF/F mice**

To determine the in vivo role of Ikkβ-mediated microglial cell activation in excitotoxin-induced hippocampal neuronal cell death, we introduced KA directly into the brains of wild-type and LysM-Cre/IkkβF/F mice. The i.c.v. KA introduction is a well-established excitotoxicity model that induces behavioural manifestations of seizures in mice and selective hippocampal cell death (Cho et al., 2003). We did not observe any discernible differences between wild-type and LysM-Cre/IkkβF/F mice in terms of seizure-like behaviour or hippocampal EEG analysis [power spectrum main frequency range, 1 day after KA injection: 0.5–3 Hz in wild-type mice (n = 5) versus 0.7–1.7 Hz in LysM-Cre/IkkβF/F mice (n = 4); 3 days after KA injection: 0.5–2 Hz in wild-type mice (n = 5) versus 0.5–2 Hz in LysM-Cre/IkkβF/F mice (n = 4)]. Hippocampal neuronal cell death was measured by cresyl violet staining. After KA administration, a characteristic loss of pyramidal neurons in the CA1 and CA3 subfields of the ipsilateral hippocampus was observed in wild-type mice (Fig. 2C and D), whereas no obvious neuronal loss was observed in the ipsilateral hippocampus of PBS-injected wild-type mice (Fig. 2A and B). The extent of hippocampal neuronal loss was significantly reduced in LysM-Cre/IkkβF/F mice (Fig. 2E and F). Three days after KA injection, the number of live neuronal cells in the CA1 and CA3 subfields of wild-type mice decreased by 58% and 76%, respectively, whereas in LysM-Cre/IkkβF/F mice, live neuronal cells in the CA1 and CA3 areas were reduced by only 38% and 60%, respectively (Fig. 2G). Comparable levels of neuronal loss were obtained by counting neuronal cells by immunohistochemistry using anti-NeuN antibody (data not shown). Interestingly, however, the reduction in neuronal loss in LysM-Cre/IkkβF/F mice compared with wild-type mice was not prominent 1 day after KA injection (Fig. 2G).

**KA-induced glial cell activation is reduced in LysM-Cre/IkkβF/F mice**

To assess the effects of Ikkβ deletion on KA-treated hippocampal glial cells, activation of microglia and astrocytes was analysed by immunohistochemistry using anti-Iba-1 and anti-GFAP antibodies, respectively (Figs 3 and 4). In PBS-treated wild-type mice, ionized calcium binding adaptor molecule-1 (Iba-1)-immunoreactive (IR) microglia were in their resting form (Fig. 3A–C). In KA-injected wild-type mice, along with hippocampal neuronal loss, the number of Iba-1-IR cells was remarkably increased in both the CA1 and CA3 regions of the ipsilateral hippocampus (Fig. 3D–F). Iba-1-IR microglia showed activated cell morphology, with enlarged cell bodies and thicker processes. In KA-injected LysM-Cre/IkkβF/F mice, however,
microglia activation was notably suppressed. There were fewer Iba-1-IR cells, and their morphology was more ramified (Fig. 3G–I). Quantitatively, Iba-1 expression was decreased by 30% (Fig. 3J). The suppression of microglial activation in LysM-Cre/Ikk$\beta^{-/-}$ mice was also confirmed by measuring the CD11b mRNA levels (Fig. 3K). The extent of microglial activation in the hippocampus was well correlated with the rate of neuronal loss in the adjacent
region (data not shown). Similarly, astrocyte activation in the ipsilateral hippocampus after KA-treatment was also attenuated in LysM-Cre/IlkβF/F mice compared with wild-type mice (Fig. 4A–K). These results show that microglia-specific Ilkβ deletion suppresses KA-induced microglia activation in the hippocampus in vivo and that microglia activation may influence astrocyte activation.

**Fig. 2.** KA-induced hippocampal neuronal cell death is decreased in LysM-Cre/IlkβF/F mice. Wild-type (IlkβF/F) (A–D) and LysM-Cre/IlkβF/F (E and F) mice were i.c.v. injected with either PBS (A and B) or KA (0.2 μg in 4 μl PBS) (C–F). Cryosections (30 μm thick) were stained with cresyl violet. Arrows indicate dying neurons and arrow heads show live cells. Scale bars: 50 μm. (G) The rate of neuronal loss in the CA1 or CA3 area of ipsilateral side hippocampus was evaluated by counting live cells. Data are presented as mean ± SEM. (Student’s t-test, *P < 0.05, **P < 0.01; versus wild-type mice.)

**Microglial IKKβ deletion is responsible for the attenuation of KA-induced hippocampal neuronal cell death**

Our results from LysM-Cre/IlkβF/F mice suggest that microglia-specific Ilkβ deletion decreases KA-induced hippocampal neuronal cell death. However, they do not
rule out the possibility that other myeloid cells such as macrophages or neutrophils from the periphery are involved, since the Ikkβ gene is also deleted in these cell types (Greten et al., 2004). To test this possibility, we examined macrophage and neutrophil infiltration of the brain parenchyma following KA treatment. For macrophage detection, we used antibodies for neuron-glial antigen 2 (NG2) and CD11b. It has been previously reported that blood-derived macrophages are double-positive for NG2 and CD11b, while oligodendrocyte precursor cells are single-positive for NG2 (Bu et al., 2001; Jones et al., 2002).

A subpopulation of CD11b-IR cells co-expressing NG2 (CD11b+/NG2+) was detected in the ipsilateral hippocampus of KA-injected wild-type mice (Fig. 5A–C), whereas no obvious CD11b+/NG2+ cells were observed in the PBS-injected ipsilateral hippocampus or KA-injected contralateral hippocampus (data not shown). Interestingly, the number of CD11b+/NG2+ cells was slightly reduced in the LysM-Cre/IkkβΔ/Δ mice (3.8 ± 0.7/field) compared with the wild-type mice (5.3 ± 0.5/field), though this was not statistically significant (Fig. 5D). In addition, we did not observe significant neutrophil infiltration in hippocampal parenchyma in either control or LysM-Cre/IkkβΔ/Δ mice after KA injection (data not shown), which is consistent...
with findings from a previous report (Andersson et al., 1991a). These data suggest that it is not likely that reduced KA-induced neuronal loss in LysM-Cre/Ilkβ^{f/f} mice is due to a reduction in myeloid cell infiltration in these mice. To further exclude the contribution of blood-derived macrophages or neutrophils, we adopted an OHSC system, and tested the effects of Ilkβ^{f/f} deletion in microglia on excitotoxicity ex vivo. Hippocampal slices from wild-type and LysM-Cre/Ilkβ^{f/f} mice were maintained in culture medium for 2 weeks prior to KA stimulation to eliminate any blood-derived macrophages or neutrophils (Fig. 6A). Stimulated cultures were evaluated using cellular uptake of PI as a measure of excitotoxic neuronal damage. Immediately after a 3 h exposure to KA (0 h of recovery), the fluorescence values of PI uptake in the CA1 and CA3 areas were slightly increased to 21.1 ± 3.2% and 11.5 ± 2.7%, respectively, in wild-type OHSCs, and similar levels of PI uptake were detected in the LysM-Cre/Ilkβ^{f/f} mice (Fig. 6D, E, H and I). Upon 24 h recovery after KA exposure, PI uptake in the CA1 and CA3 regions of wild-type OHSCs was further increased to 42.5 ± 5.5% and 25.5 ± 4.7%, respectively. However, in OHSCs from LysM-Cre/Ilkβ^{f/f} mice, the increase in PI uptake during the recovery period was significantly attenuated compared...
with wild-type: it increased to only 25.6 ± 3.6% and 17.1 ± 2.6%, respectively (Fig. 6F–I). In control OHSCs, however, a slightly higher level of basal PI uptake was detected in slices from LysM-Cre/IKK\(b\)/F/F mice. Taken together, these data argue that IKK\(b\) deletion in microglia is responsible for the attenuation of KA-induced hippocampal neuronal cell death and that the IKK\(b\)/NF-\(\kappa B\) signalling pathway in microglia may play an important role in KA-induced excitotoxicity in the hippocampus.

**KA-induced proinflammatory gene expression is reduced in LysM-Cre/IKK\(b\)/F/F mice**

In an attempt to elucidate the mechanisms underlying the difference in excitotoxic susceptibility of wild-type and LysM-Cre/IKK\(b\)/F/F mice, we measured the mRNA expression levels of proinflammatory NF-\(\kappa B\)-target genes such as TNF-\(\alpha\), IL-1\(\beta\) and iNOS in the hippocampus. These genes have been implicated in excitotoxic hippocampal neuronal cell death (De Simoni et al., 2000; Rizzi et al., 2003). The mRNA levels of TNF-\(\alpha\), IL-1\(\beta\) and iNOS in the hippocampi of KA-stimulated wild-type mice increased 12-, 35- and 3-fold, respectively (Fig. 7). KA-induced expression of these proinflammatory genes in hippocampi of LysM-Cre/IKK\(b\)/F/F mice, however, was attenuated by 30–50%. These data demonstrate that KA-induced inflammatory gene expression is reduced in LysM-Cre/IKK\(b\)/F/F mice.

**TNF-\(\alpha\) and IL-1\(\beta\) contribute to KA-induced hippocampal cell death in OHSCs**

We then tested the effects of proinflammatory cytokines on excitotoxicity in OHSCs (Fig. 8). Treatment of KA-stimulated LysM-Cre/IKK\(b\)/F/F OHSCs with TNF-\(\alpha\) (10–40 ng/ml in CA1; 20 ng/ml in CA3) completely elevated the cell death rate to the level seen in wild-type OHSCs (Fig. 8B). Likewise, treatment with IL-1\(\beta\) (0.1–10 ng/ml in CA1; 5–10 ng/ml in CA3) enhanced the KA-mediated excitotoxicity in LysM-Cre/IKK\(b\)/F/F OHSCs (Fig. 8C). The specificity of the cytokines was confirmed using blocking antibodies against TNF-\(\alpha\) and IL-1\(\beta\) in this experiment (data not shown). Furthermore, the addition of anti-TNF-\(\alpha\) or anti-IL-1\(\beta\) blocking antibodies in the wild-type OHSCs reduced KA-mediated excitotoxicity by 30–60% (Fig. 8B and C). Taken together, these data suggest that TNF-\(\alpha\) and IL-1\(\beta\) expression in wild-type OHSCs potentiates KA excitotoxicity, and that decreased expression of these cytokines in LysM-Cre/IKK\(b\)/F/F OHSCs partly accounts for the decreased KA excitotoxicity.

**Ischaemic brain damage and microglia activation after transient MCAO is reduced in LysM-Cre/IKK\(b\)/F/F mice**

To verify the neuroprotective effects of microglial IKK\(b\) deletion in a more physiologically relevant disease model, we induced ischaemic brain damage by MCAO in
LysM-Cre/Ikk^{+/−}/F/F mice. It is well known that excitotoxicity is one of the underlying mechanisms of ischaemic neurodegeneration (Doble, 1999). A 1-h MCAO followed by a 3-day reperfusion period induced a 40% degeneration of the ipsilateral brain, as calculated by infarct volume (Fig. 9A and B). In LysM-Cre/Ikk^{+/−}/F/F mice, however, the infarct size decreased to 10%. We then tested microglia activation after MCAO by Iba-1 immunostaining. As previously reported (Schilling et al., 2003), ischaemic injury induced microglia activation around the infarct region in wild-type mice (Fig. 9C). The levels of microglia activation were variable depending on the distance to the infarct region: the closer to the infarct region, the stronger the microglia activation (Fig. 9C, compare A and B). Microglia activation was also detected in LysM-Cre/Ikk^{+/−}/F/F mice. However, the activation levels were much weaker than those of wild-type mice (Fig. 9C, compare upper panels versus lower panels). These data demonstrate that LysM-Cre/Ikk^{+/−}/F/F mice. It is well known that excitotoxicity is one of the underlying mechanisms of ischaemic neurodegeneration (Doble, 1999). A 1-h MCAO followed by a 3-day reperfusion period induced a 40% degeneration of the ipsilateral brain, as calculated by infarct volume (Fig. 9A and B). In LysM-Cre/Ikk^{+/−}/F/F mice, however, the infarct size decreased to <10%. We then tested microglia activation after MCAO by Iba-1 immunostaining. As previously reported (Schilling et al., 2003), ischaemic injury induced microglia activation around the infarct region in wild-type mice (Fig. 9C). The levels of microglia activation were variable depending on the distance to the infarct region: the closer to the infarct region, the stronger the microglia activation (Fig. 9C, compare A and B). Microglia activation was also detected in LysM-Cre/Ikk^{+/−}/F/F mice. However, the activation levels were much weaker than those of wild-type mice (Fig. 9C, compare upper panels versus lower panels). These data demonstrate that...
**Discussion**

To address the *in vivo* role of inflammatory microglia activation in excitotoxicity, we employed myeloid cell type-specific *Ikkβ* conditional knockout (*LysM-Cre/*Ikkβ*F/F) mice. In primary cultured microglia from neonate *LysM-Cre/*Ikkβ*F/F mice, we found that the *Ikkβ* deletion frequency was ~36%, but the deletion frequency in microglia isolated directly from adult or neonate *LysM-Cre/*Ikkβ*F/F mice was much lower. This difference may be due to the fact that microglia are in their resting state *in vivo* and then become spontaneously activated during *in vitro* culture, resulting in upregulation of the lysozyme M gene (Ohmi et al., 2003). This explanation accounts for the increase in the *Ikkβ* deletion rate to ~73% in microglia from KA-treated *LysM-Cre/*Ikkβ*F/F ipsilateral hippocampus (Fig. 1E). Interestingly, we detected a higher microglial *Ikkβ* deletion frequency in neonate mice compared with adult mice. This implies that lysozyme M gene expression is developmentally regulated in microglia, which may also contribute to the enhanced *Ikkβ* deletion frequency observed in the primary cultured microglia. In addition, our *in vitro* data demonstrate that the *Ikkβ* gene is deleted in microglia, but not in astrocytes or in neurons of the *LysM-Cre/*Ikkβ*F/F brain and, in the absence of IKKβ in microglia, IKKβ/NF-κB-dependent inflammatory gene expression is attenuated. These data argue that *LysM-Cre/*Ikkβ*F/F mice can be used to investigate the *in vivo* role of microglia activation in excitotoxic neurodegeneration.
It should be noted that, in these mice, only partial deletion (73% maximum) of the \(Ikk^\beta\) allele in the KA-activated hippocampal microglia population was achieved. The incomplete deletion of the \(Ikk^\beta\) allele in the entire population of microglial cells resembles the situation in macrophages, where the \(Ikk^\beta\) deletion rate rarely exceeded 75% in bone marrow-derived macrophages from \(LysM-Cre/Ikk^{\beta/\beta}\) mice (Greten et al., 2004). The incomplete deletion of a target gene is often a drawback of using tissue-specific conditional knockout mice. Although a study using conventional knockout mice does not face such problems, it does not provide researchers with cell type-specific information either. Indeed, it was reported that p50 knockout mice are more vulnerable to KA-induced excitotoxicity, indicating a beneficial role of NF-\(\kappa\)B activation in these mice (Yu et al., 1999). However, such effects were attributed mainly to NF-\(\kappa\)B activation in neurons, but not in microglia. Similarly, the neuroprotective function of microglial activation was suggested in a recent study using MyD88 knockout mice (Simard and Rivest, 2007), in which the effects of MyD88 deletion in microglia versus non-microglial cells could not be differentiated. Our in vivo data, however, indicate that IKK\(\beta\) deletion in microglia exerts protective effects against KA-induced excitotoxicity. Thus far, several reports have suggested a neurotoxic role of microglia in excitotoxicity. However, most of these studies are based on circumstantial evidence supported by in vitro experiments using cultured microglia. In this regard, our study using \(Ikk^\beta\) conditional knockout mice, conclusively demonstrates the in vivo role of the IKK/NF-\(\kappa\)B-mediated microglia activation in excitotoxicity. Interestingly, the attenuation of excitotoxicity in \(LysM-Cre/Ikk^{\beta/\beta}\) mice was not statistically significant 1 day after injection, but was substantial 3 days after i.c.v.

**Fig. 9** Infarct size and microglial activation following MCAO are reduced in \(LysM-Cre/Ikk^{\beta/\beta}\) mice. Wild-type and \(LysM-Cre/Ikk^{\beta/\beta}\) mice were subjected to transient MCAO for 1 h and reperfused. (A) After 71 h, the brains were removed, cut into 2-mm thick blocks and stained with triphenyl tetrazolium chloride. (B) The infarct area was measured and expressed as the percentage of the ipsilateral hemisphere. Data are presented as mean ± SEM. (**P < 0.001 by Student's t-test; versus wild-type mice; \(n = 4\)). (C) Cryosections of the second blocks were stained with anti-Iba-1 antibody. Representative images of five different regions were captured and are presented (ipsilateral: a–d, contralateral: e). Scale bars: 50 \(\mu\)m.
administration of KA (Fig. 2). It has been reported that introduction of KA into the brain induces excitotoxicity through two different mechanisms. Primary hippocampal damage is induced within 24 h by KA-mediated seizure activity, while further delayed neuronal cell death follows this initial damage after 2–3 days (Doble, 1999). The absence of any significant difference in terms of hippocampal EEG activity argues against that reduced neuronal death in the knockout mice is due to reduced seizures in these mice. Rather, our data suggest that IKKβ-mediated microglia activation contributes to delayed excitotoxic neurodegeneration in the later stage. These results are consistent with those of previous reports showing that inflammatory mediators contribute to excitotoxic neurodegeneration at later stages (Giulian and Vaca, 1993). Notably, we observed attenuation of glial cell activation not only in microglia, but also in astrocytes of LysM-Cre/IKKβ F/F mice (Fig. 4A–K). Considering that IKKβ is deleted only in microglia, it is likely that KA-induced astrocyte activation is secondary to microglia activation.

Thus far, it is not clear how microglia become activated upon KA stimulation. Although direct microglial activation by KA has been reported (Noda et al., 2000), we were not able to detect any proinflammatory gene expression after KA treatment of cultured hippocampal glial cells (data not shown). Therefore, it is more likely that microglia are indirectly activated by KA-damaged neurons. In this regard, it is of interest that high-mobility group box-1 (HMGB-1), a non-histone DNA-binding protein, was recently reported to be released by damaged neurons in the ischemic brain, thus activating microglia (Kim et al., 2006). It has also been documented that HMGB-1 exerts its cytokine-like function by activating toll-like receptor (TLR) 2 and 4 on innate immune cells (Park et al., 2004). In the CNS, TLR2 and 4 are constitutively expressed on microglia (Olson and Miller, 2004). Considering that IKK/NF-κB activation is a major downstream signal of TLR, it is tempting to speculate that, in our excitotoxicity model, microglia are activated by TLR binding to HMGB-1 released from KA-damaged hippocampal neurons. This can be addressed in future studies using TLR2- or 4-deficient mice.

It should be noted that, in this study, we did not find direct in vivo evidence that the reduction in neuronal loss in the knockout mice was due to IKKβ deletion in microglia, since IKKβ in these mice was also deleted in other myeloid lineage cells. However, indirect evidence suggests microglia-specific effects on excitotoxicity. First, in immunohistochemistry tests, we did not observe a statistically significant reduction in macrophage infiltration in LysM-Cre/IKKβ F/F mice after KA administration. More importantly, neurons in OHSCs from LysM-Cre/IKKβ F/F mice were relatively resistant to the KA-induced excitotoxicity. In such an OHSC model, the effects of blood-derived myeloid cells were minimized, since hippocampal slices were cultured in vitro without blood supply for 2 weeks before the experiment. Considering these data, we concluded that microglia-specific IKKβ deletion plays a major role in the attenuation of excitotoxicity.

In ex vivo experiments, excitotoxic hippocampal cell death was reduced by 30–40% in the OHSCs of LysM-Cre/IKKβ F/F mice, which is reminiscent of the reduction rate in vivo. However, the kinetics of the cell death were dissimilar. In the in vivo system, we did not observe a statistically significant difference in the cell death rate between wild-type and LysM-Cre/IKKβ F/F mice 24 h after KA injection, whereas in OHSCs, the decrease in cell death was prominent after 24 h of recovery (Figs 2 and 6). This can be simply attributed to the temporal difference in KA accessibility to hippocampal neurons in vivo versus ex vivo. Alternatively, this can be explained by the difference in the deletion rate of IKKβ at the time of stimulation. Since hippocampal slices were prepared from neonate mice, it is likely that the microglial IKKβ deletion rate of OHSCs is higher than the in vivo rate of adult mice, which may account for the difference.

To elucidate the basis of neurotoxic effects of IKKβ activation, we monitored the expression of several putative neurotoxic mediators that can be induced upon IKKβ activation in microglia. KA-induced TNF-α, IL-1β and iNOS gene expression was reduced in the ipsilateral hippocampus of LysM-Cre/IKKβ F/F mice (Fig. 7). In addition, exogenous addition of TNF-α and IL-1β to LysM-Cre/IKKβ F/F OHSCs enhanced their excitotoxic susceptibility (Fig. 8). These data imply that IKKβ-dependent expression of these inflammatory cytokines may be, at least in part, responsible for the delayed excitotoxicity. The neurotoxic effects of IL-1β and iNOS in excitotoxicity are well documented (Hara et al., 1997). Likewise, TNF-α has been implicated as a critical mediator of neuronal cell death in cerebral ischaemia (Meistrell et al., 1997). However, it has also been reported that TNF-α expression during excitotoxic damage plays a neuroprotective role (Cheng et al., 1994). Furthermore, TNF receptor-deficient mice are more susceptible to excitotoxic brain injury, which also suggests a neuroprotective role of TNF-α in vivo (Bruce et al., 1996). Thus far, there is no clear explanation for these discrepancies. It should be noted, however, that the TNF receptor gene is deleted in all brain cells of the knockout mice, including neurons and glia, from early development. In LysM-Cre/IKKβ F/F mice, however, TNF-α production is altered only in microglia and only after excitotoxic brain injury, which may account for the different results. Moreover, deletion of microglial IKKβ reduced the expression of other inflammatory mediators. Therefore, the neurotoxic effects of microglial activation might be due to the concerted effects of various IKKβ target genes. In our study, we measured expression of proinflammatory cytokines in vivo, but confirmed their neurotoxic effects ex vivo using OHSCs. Therefore, it is formally possible that another IKKβ-dependent gene not tested in this study contributes to delayed neuronal cell death in vivo. For instance, microglial production of tPA
Supplementary material
Supplementary material is available at Brain online.

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