Seizures and Disturbed Brain Potassium Dynamics in the Leukodystrophy Megalencephalic Leukoencephalopathy with Subcortical Cysts

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Objective: Loss of function of the astrocyte-specific protein MLC1 leads to the childhood-onset leukodystrophy “megalencephalic leukoencephalopathy with subcortical cysts” (MLC). Studies on isolated cells show a role for MLC1 in astrocyte volume regulation and suggest that disturbed brain ion and water homeostasis is central to the disease. Excitability of neuronal networks is particularly sensitive to ion and water homeostasis. In line with this, reports of seizures and epilepsy in MLC patients exist. However, systematic assessment and mechanistic understanding of seizures in MLC are lacking.

Methods: We analyzed an MLC patient inventory to study occurrence of seizures in MLC. We used two distinct genetic mouse models of MLC to further study epileptiform activity and seizure threshold through wireless extracellular field potential recordings. Whole-cell patch-clamp recordings and K⁺-sensitive electrode recordings in mouse brain slices were used to explore the underlying mechanisms of epilepsy in MLC.

Results: An early onset of seizures is common in MLC. Similarly, in MLC mice, we uncovered spontaneous epileptiform brain activity and a lowered threshold for induced seizures. At the cellular level, we found that although passive and active properties of individual pyramidal neurons are unchanged, extracellular K⁺ dynamics and neuronal network activity are abnormal in MLC mice.

Interpretation: Disturbed astrocyte regulation of ion and water homeostasis in MLC causes hyperexcitability of neuronal networks and seizures. These findings suggest a role for defective astrocyte volume regulation in epilepsy.

ANN NEUROL 2018;83:636–649

View this article online at wileyonlinelibrary.com. DOI: 10.1002/ana.25190

Received May 26, 2017, and in revised form Jan 12, 2018. Accepted for publication Feb 18, 2018.

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Additional supporting information can be found in the online version of this article.

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Megencephalic leukoencephalopathy with subcortical cysts (MLC; MIM 604004) is a rare childhood-onset leukodystrophy,1,2 caused by homozygous recessive mutations in MLC13 or GLIALCAM.4 MLC1 is a membrane protein almost exclusively expressed in brain astrocytes.5 GlialCAM is a chaperone of MLC1, ensuring its localization in the membrane of astrocyte endfeet.4,6,7 Patients show very high water content in the brain white matter. Biopsies reveal countless fluid-filled vacuoles within the outer lamellae of myelin sheaths and, to a lesser degree, in perivascular astrocyte endfeet.8 Mutations affecting MLC1 protein negatively impact volume-regulated anion channel (VRAC) activity in astrocytes, leading to impaired regulatory volume decrease9 and resulting in chronically swollen astrocytes.10

Because the pathophysiology of MLC has mainly been studied in isolated cells, consequences of astrocyte dysfunction for neuronal network functioning are unknown. The disease is characterized by infantile-onset macrocephaly, slow deterioration of motor functions, and later cognitive decline. Minor head trauma can lead to temporary worsening, often with seizures, prolonged unconsciousness, and motor deterioration.11 Mild epilepsy, which typically responds well to treatment, has been described in many MLC patients, but status epilepticus has also been described.12 Occurrence of epilepsy has not been studied in a large cohort of genetically confirmed patients, and its cellular basis is not understood.

The panglial syncytium, the glial network consisting of gap-junction coupled astrocytes and oligodendrocytes, is essential for the uptake and dispersal of extracellular K\(^+\) ([K\(^+\)]\(_o\)) released from neurons during periods of high neuronal activity.13,14 Disruption of proteins important for this process, such as Kir4.1 K\(^+\) channels,15,16 glial gap-junction proteins,17 and the astrocytic water channel aquaporin-4 (AQP4),18 lead to impaired [K\(^+\)]\(_o\), clearance and to epileptiform brain activity or an altered seizure threshold. MLC1 and GlialCAM undergo molecular interactions with several of these proteins.19,20 Therefore, we hypothesize that defective astrocyte volume regulation in MLC leads to impaired control of [K\(^+\)]\(_o\),8 resulting in hyperexcitability of neuronal networks and epilepsy.

We recently developed an Mlc1-null mouse recapitulating major pathological features of MLC.10 Additionally, Glialcam-null mice21 show similar pathological features.7,22 Mlc1- and Glialcam-null mice (collectively referred to here as MLC mice) are excellent tools to investigate the pathophysiology of MLC. Here, we combine a clinical inventory of MLC patients with in vivo and in vitro electrophysiology in MLC mice to study the cellular pathophysiology of epilepsy in MLC. We confirm that early seizure onset is common in MLC patients. Similarly, MLC mice show spontaneous epileptiform activity and a reduced threshold for seizure induction. At the cellular level, we find that excitability of single neurons is unchanged, whereas stimulation-induced increases in [K\(^+\)]\(_o\), and network excitability are increased. Taken together, we demonstrate that loss of function of the astrocytic protein MLC1 disturbs neuronal network activity while leaving intrinsic neuronal properties intact.

Materials and Methods

Study Approval

Experimental procedures involving mice were in strict compliance with animal welfare policies of the Dutch government and were approved by the Institutional Animal Care and Use Committee of the VU University, Amsterdam. The clinical inventory was approved by the Medical Ethics Committee of the VU University Medical Center.

MLC Patients

We executed a cross-sectional observational study among 205 genetically proven patients with autosomal-recessive MLC present in the Amsterdam Leukodystrophy database, containing patients referred to the VUmc Center for Childhood White Matter Disorders for genetic testing. As part of a larger study on the clinical phenotype of MLC,23 standardized clinical questionnaires including items on epilepsy were completed primarily by the patient’s physician. If this source was unavailable, information was derived from medical records, supplemented by information provided by families.

Mice

Transgenic mice and their wild-type littermates had a C57BL/6J background. Generation of both Mlc1 deficient mice (Mlc1-null)21 and Glialcam deficient mice (Glialcam-null)21,22 was previously described. Null animals were compared with wild-type littermate controls, with the exception of extracellular field potential/seizure threshold recordings. Here, littermate data for both MLC lines were grouped because no differences between littermates were observed.

Behavioral Assays

In vivo experiments were performed on adult (8–12 months old) mice. Forty-eight-hour recordings of basal activity were performed on both male and female mice. Kainate-induced seizures were only studied in male mice because sex strongly influences sensitivity to kainate.24 To score hindlimb clasping, each mouse was suspended for 30 seconds by its tail and kept 10cm above a stable platform. Seizure threshold and severity were assessed using an intraperitoneal injection of kainate (kainic acid; 10mg/kg). Behavioral response was scored using a revised Racine’s seizure scale.25 Maximal score during
subsequent 10-minute time bins was assessed by a trained observer unaware of genotype. Mice were sacrificed if seizure severity reached score 5c.

**Telemetric Extracellular Field Potential Recordings**

Electrocorticogram (ECoG) recordings and hippocampal local field potential (LFP) recordings were performed with wireless ETA-F10 transmitters (Data Science International, St. Paul, MN). For electrode placement, animals were anesthetized with isoﬂurane (induction 3%, ﬂow rate 0.8l/min; maintenance 1.5–1.8%, ﬂow rate 0.6l/min). For ECoG recordings, a recording electrode (stereotaxic coordinates relative to bregma: 2.22mm anterior and 1mm lateral, corresponding to motor cortex) was implanted subdurally through a small hole drilled in the skull and held in place with a screw (A2-70; Jevaka, Almere-Poort, The Netherlands) made of stainless steel. For hippocampal LFP recordings, an insulated depth-electrode (W-electrode; Open Source Instruments, Watertown, MA) was lowered into the brain (stereotaxic coordinates relative to bregma: 2.2mm posterior, 2.0mm lateral, 1.7mm depth, corresponding to the CA1 region of hippocampus). A ground electrode was placed subdurally 6mm posterior and 1mm lateral relative to bregma. All electrodes were sealed with dental cement. The calibrated transmitter was placed subcutaneously along the dorsal flank of the animal. Mice received analgesia (Temgesic; 0.1mg/kg subcutaneously) before surgery and recovered for 7 days before recordings.

ECoG/LFP signal was collected through a radiofrequency receiver (RLA1020; Data Science International) and analog output adapter (Option RO8, Data Science International) using Powerlab 4/10 (AD-Instruments, Austin, TX; sampling: 1kHz; filtering: 200Hz low pass). Infrequent periods of severe electrical noise were manually removed from the recording before further analysis. Recordings were analyzed using the Neuroarchiver tool (Open Source Instruments; http://www.opensourceinstruments.com/Electronics/A3018/Seizure_Detection.html). For more details on analysis, see the website or publication.26 In brief, data were split into 1-second epochs. A power threshold was set (5 times baseline power), and putative events were epochs in which this threshold was crossed. Values of six metrics (power in the 0–200Hz band, coastline, voltage asymmetry, coherence, intermittency, and rhythm) were determined for each putative event, and these were compared with a library of events manually classified as seizure-like. Events were classified according to their Euclidian distance to events in a manually constructed event library. The library was constructed by an operator who classified events as “no event/baseless” (no obvious electrographic or behavioral event), “interictal event,” or “artifact” (grooming related noise or antenna glitches). In recordings from kainate-injected animals, additional event types appeared: large single negative peak spike waves (simple spike-waves; SS); spike waves having one or more shoulders (complex spike-waves; CS); and runs of intermediate frequency (IF) discharges (5–12Hz) lasting 0.2 to 12 seconds with typically smaller amplitude (Fig 4B and Supplementary Movie 1). Following library construction, all data were analyzed without further manual interference.

**Slice Preparation**

Acute brain slices were prepared from 3- to 5-month-old mice. After decapitation, the brain was removed in ice-cold solution containing (mM): 70 sucrose, 70 NaCl, 25 D-glucose, 25 NaHCO3, 2.5 KCl, 1.25 NaH2PO4, 1H2O, 5.0 MgSO4.7H2O, 1 CaCl2, 3 sodium pyruvate, and 1 sodium ascorbate (carboxygenated with 5%CO2/95%O2, 300–310mOs). Coronal hippocampal slices (350µm) were cut on a microtome (Thermo Fisher Scientific, Waltham, MA), kept in slicing solution at 32°C for 20 minutes, followed by 20-minute recovery at room temperature. Afterward, slices were stored for >1 hour at room temperature in ACSF containing (mM): 125 NaCl, 3 KCl, 1.2 NaH2PO4, 1 MgCl2, 2 CaCl2, 26 NaHCO3, and 10 glucose (carboxygenated with 5%CO2/95%O2, 300–305mOs).

**Patch-Clamp Recording**

Slices were transferred to the setup and perfused with ACSF (5ml/min at 30–31°C). Recording ACSF contained CNQX (10µM), AP-5 (50µM), and Gabazine (3µM). Patch pipettes (3–5MΩ) were filled with intracellular solution containing (mM): 111 K-glucurate, 8 KCl, 10 HEKES, 4 Mg-ATP, 0.4 Na-GTP, 10 K-phosphocreatine, 0.2 EGTA, and 0.37% biocytin (7.4 pH; 290 ± 10mOs). Recordings were made using an Axopatch 700B amplifier and Clampex software (sampling: 50kHz; filtering: 30kHz; Axon Instruments, Union City, CA). After determination of resting membrane potential, cells were kept in current clamp at −65mV through steady current injection. Data were analyzed using a custom-written script in Matlab (version R2012a; The MathWorks, Inc., Natick, MA). Action potentials (APs) were required to exceed a membrane potential threshold of −20mV and a speed threshold of 10mV/ms to be included in analysis. AP threshold was deﬁned by determining the peak of the AP derivative, and measuring the membrane potential at which 10% of this maximum was reached. AP half-width was calculated as the time difference between the two time points corresponding to 50% of the AP amplitude (AP peak/AP threshold). For most neurons, pyramidal identity was conﬁrmed after recording by biocytin staining, using the avidin-biotin-peroxidase method.

**K+-Sensitive Electrode Recordings**

K+-sensitive microelectrodes were prepared from thin walled nonfilamented glass capillaries (GC150T-10; Harvard Apparatus, Holliston, MA) pulled to a tip inner diameter in the range of 2µm.28 Microelectrodes were silanized with gaseous N,N-dimethyltrimethylsilylamine (41716; Sigma-Aldrich, St. Louis, MO) at 200°C for 8 to 10 hours before being filled with a short column (120–150µm) of liquid membrane solution (99373, K+ Ionophore I Cocktail B; Sigma-Aldrich) and 150mM of NaCl, 3mM of KCl as backﬁlling solution (150–180MΩ). Tips of a K+-sensitive and a field-potential electrode (2MΩ, ﬁlled with 150mM of NaCl, 3mM of KCl) were placed ~5 µm (±2 µm) from each other. Voltage signal from both channels to the same reference electrode were recorded using a MultiClamp 700B double amplifier (Axon Instruments), and differential voltage between the two electrodes was calculated by...
subtraction. K⁺-sensitive electrodes were calibrated before and after the experiment (calibration solutions: 3, 6, 12, 24, or 48 mM of KCl, 150 mM of NaCl). Data obtained with a K⁺-sensitive electrode were excluded when calibrations before and after differed by more than 2.5 mV for 3, 6, or 12 mM of KCl.

For recording [K⁺]o dynamics in hippocampus, both K⁺-sensitive electrode and field-potential electrode were placed in CA1 stratum radiatum (depth ~80–100 µm). A concentric bipolar stimulation electrode was placed ~550 to 600 µm lateral from the recording site onto stratum radiatum. Input-output curves were constructed by recording field excitatory postsynaptic potentials (fEPSPs) following single stimulation pulses of increasing strength (0–950 µA; 50-µA steps). Trains of 10-second stimulation at 5, 10, or 20 Hz, 450 µA, were applied twice and the averaged response parameters were calculated. Subsequently, recording electrodes were moved into stratum pyramidale (stimulation electrode remained in the same location). Data were analyzed using custom-written procedures in IGOR-pro (Wavemetrics, Lake Oswego, OR). Population spike area in stratum pyramidale was defined as the area between large negative deflections in the measured fEPSP and a fitted diagonal connecting the two neighboring local maxima (Fig. 8B; shaded area).

**Statistical Analysis**

Statistical analysis was performed with Prism (version 7; GraphPad Software Inc., La Jolla, CA). For behavioral and EEG recordings, nonparametric analysis of variance (ANOVA; Kruskal-Wallis), followed by Dunn’s multiple comparisons (wild-type versus Mlc1-null and Glialcam-null) was performed. For patch-clamp data and basal [K⁺]o parameters, we tested for normality using the Shapiro-Wilk test. Data were compared using regular t test or, in case of deviant normal distribution, Mann-Whitney U test. For input-output analysis and analysis of stimulation-induced [K⁺]o, two-way ANOVA followed by Sidak’s multiple comparison test was performed. Kaplan-Meier survival curves were compared using log-rank statistics.

**Results**

**Clinical Seizures in MLC Patients**

Information on seizure characteristics was obtained from 141 patients with recessive MLC1 mutations and 17 patients with recessive GLIALCAM mutations (137 families; Supplementary Table 1). Survival analysis showed that 75.8% of patients with MLC1 mutations and 73.4% of patients with GLIALCAM mutations experienced at least one seizure before 25 years of age (Fig 1A,B). Curves were not different for patients with an MLC1 defect and patients with a GLIALCAM defect.

Of all patients with seizures, 13% had experienced only one seizure at the time of clinical inventory, often provoked by mild head trauma (8 of 12 patients) or by febrile illness (2 of 12 patients). Seventy-three percent had epilepsy that was well controlled with medication. Of all patients ≥6 years of age, 63% met the International League against Epilepsy criteria for epilepsy. Clinically, generalized tonic-clonic seizures and focal seizures (sometimes with impaired awareness and/or focal to bilateral tonic-clonic seizures) were most common. Mild head trauma was an important provoking factor, causing one or more seizures in 55% of patients with seizures. One or more status epilepticus had occurred in 17%, of which most (58%) occurred for the first time a few years after the first seizure. 6 of the 158 patients were deceased. Of these 6, 3 deaths were related to epilepsy.

Taken together, the patient inventory shows that an early onset of epileptic seizures is an integral part of the MLC phenotype. Mild head trauma is an important provoking factor and status epilepticus occurs relatively frequently and early in the disease.

**Hindlimb Clasping and Spontaneous Interictal Activity in MLC Mice**

To understand the cellular pathophysiology of epilepsy in MLC, we studied two MLC mouse models: the Mlc1-null mouse and the Glialcam-null mouse. Previous automated behavioral analysis of Mlc1-null mice did not uncover behavioral abnormalities. For Glialcam-null
mice, behavioral analysis has not been reported. We observed that both Mlc1-null and Glialcam-null mice showed hindlimb clasping when suspended from the tail (Fig 2). This is a sign of neurological abnormality, co-occurring with seizures or increased seizure susceptibility in several mouse models for neurological diseases.30,31

To test whether abnormal electrical activity is present in brains of MLC mice, we performed wireless ECoG recordings with an electrode placed above the motor cortex in both MLC mouse models (Fig 3A). Long-term recordings (48 hours) from mice in their home cage revealed that both MLC lines show a significantly higher occurrence of abnormal discharges than wild-type mice (Fig 3B–E). These discharges bear resemblance to interictal spikes.32 They did not have a clear behavioral correlate, as assessed from simultaneous video recordings of mouse behavior.

To assess whether interictal spikes also occurred in the hippocampus, we performed wireless LFP recordings with a depth-electrode placed in the hippocampal CA1 area. Similar to what we observed in motor cortex, MLC mice showed a significantly higher occurrence of interictal discharges compared to wild-type mice (wild-type: 15.7 ± 7.1 events/day, n = 3; MLC: 331.2 ± 95.9 events/day, n = 3 Glialcam-null and 1 Mlc1-null; p = 0.039).

Taken together, MLC mice show an overt hindlimb clasping phenotype. Although major behavioral seizures did not occur during our recording sessions, and were not observed during previous phenotyping,10 unprovoked interictal spikes are present in MLC mice.

Lowered Seizure Threshold and Increased Seizure Severity in MLC Mice

The presence of unprovoked interictal spikes in MLC mice suggests that they might be more sensitive to evoked seizure activity. To test this, mice received a single intraperitoneal injection with the chemoconvulsant kainate, at a dose which does not induce severe seizures in C57BL/6J mice (10mg/kg),33 followed by ECoG recording and behavioral scoring using the modified Racine score.25 While wild-type mice only reached Racine score 2 or 3, with time Mlc1-null and Glialcam-null mice developed more severe seizures with multiple generalized tonic-clonic seizure events (Racine score 5a–5c; Fig 4A).

Analysis of ECoG recordings following kainate injection revealed three main types of epileptic events: spike-wave complexes either with a single negative peak (SS) or featuring one or more shoulders (CS), interspersed with runs of intermediate frequency (IF) discharges (5–12Hz) lasting 0.2 to 12 seconds, which typically had a small amplitude (Fig 4B). IF discharges correlated with severe motor seizures (Supplementary Movie 1), in line with previous data.34 In the 60-minute period following kainate injection, wild-type mice showed few simple SS and CS events, with very rare IF discharges (Fig 4C–E). In contrast, both Mlc1-null and Glialcam-null mice showed a significantly higher number of IF discharges than wild-type animals (Fig 4E). No significant differences were observed in the occurrence of SS and CS events, although a trend toward more events in MLC mice was present. Taken together, these findings highlight that MLC mice have a lower threshold for kainate induced seizures than wild-type mice.

Neuronal Excitability in MLC Mice

Spontaneous interictal activity in motor cortex and hippocampus and a lowered threshold for evoked seizures suggest neuronal hyperexcitability in MLC mice. To test whether this was alternatively to increased intrinsic neuronal excitability, we performed whole-cell patch-clamp recordings from CA1 pyramidal neurons in acute hippocampal brain slices (Fig 5 and Supplementary Table 2). Passive electrophysiological properties (membrane potential,
input resistance) did not differ between MLC mice and their wild-type littermates (Fig 5D, E). Furthermore, AP waveforms were indistinguishable between Mlc1-null mice and littermate controls (Fig 5C, D). Glialcam-null mice also showed similar AP waveform properties as their wild-type littermates, apart from a slight increase in AP peak and half-width (Fig 5E). Importantly, input-output characteristics and maximal firing frequency were identical in both MLC mouse lines compared to their wild-type littermate controls (Fig 5F, G).

We hypothesized that stimulating single neurons to fire APs for a prolonged period might uncover more subtle differences in intrinsic properties. Sustained AP firing was evoked by applying prolonged current injections with step-wise increments (Fig 6A). Prolonged stimulation altered the AP waveform (Fig 6B), but again this did not differ between MLC mice and their wild-type littermate controls (Fig 6C–E).

To check whether intrinsic neuronal properties in other brain regions are similarly unaffected in MLC mice, we performed an additional set of patch-clamp recordings from layer 5 pyramidal neurons in primary motor cortex (M1). Like in hippocampus, we observed no differences in passive and active neuronal properties, nor in responses to prolonged current injections, that could explain hyperexcitability in MLC mice (data not shown).

Taken together, these results show that intrinsic excitability of principal neurons in hippocampus and neocortex is largely unchanged in MLC mice, and cannot account for the observed interictal activity and decreased seizure threshold.

**Increased Stimulation-Induced \([K^+]_o\) Around CA1 Synapses of MLC Mice**

Impaired astrocyte functioning can lead to disturbed \([K^+]_o\) dynamics. Therefore, we measured \([K^+]_o\) dynamics using...
FIGURE 4: Lowered seizure threshold in MLC mice. (A) Behavioral scoring of seizure activity over time in wild-type (blue) and MLC mice (Mlc1-null: red; Gliacam-null: green; thin lines show individual mouse responses; thick lines show averages) following kainate injection (10mg/kg intraperitoneally). For each 10-minute interval, the highest level of epileptic activity was scored using the modified Racine seizure scale. The bar graph shows maximum Racine score during 60-minute trial (open circles indicate individual mice, bars show average). MLC mice reached significantly higher behavioral seizure scores (wildtype: n = 8; Mlc1-null: n = 8; Gliacam-null: n = 7; p = 0.0014; wildtype vs Mlc1-null, p = 0.0060; wildtype vs Gliacam-null, p = 0.0023). (B) Top: Morlet-wavelet ECoG spectra from kainate-injected mice. Bottom: ECoG traces showing simple spike waves (SS), complex spike waves (CS) and runs of intermediate frequency (IF) discharges (expanded below). (C–E) Temporal progression of kainate-induced SS, CS, and IF events, respectively. Bar graphs show total number of SS, CS, and IF events in 60-minute trial. A significantly higher number of IF discharges was observed in MLC mice (wildtype: 4.1 ± 2.9 discharges/h; Mlc1-null: 39.0 ± 24.8; Gliacam-null: 35.4 ± 15.8; p = 0.0101; wildtype vs Mlc1-null, p = 0.028; wildtype vs Gliacam-null, p = 0.012). Error bars and shaded regions indicate SEM. *p < 0.05; **p < 0.01; ***p < 0.001. ECoG = electrocorticogram; MLC = megalencephalic leukoencephalopathy with subcortical cysts.
calibrated K⁺-sensitive electrodes²⁸ in acute brain slices of hippocampal CA1 area (stratum radiatum; Fig 7A). We did not observe differences in basal [K⁺]o between MLC mice and wild-type littermates at rest (Fig 7B).

Next, we simultaneously recorded [K⁺]o and field potential responses in stratum radiatum upon extracellular stimulation of Schaffer collaterals (Fig 7A). First, we determined the input-output curve of the fEPSP...
amplitude by single stimulations with step-wise increases in stimulation strength. Notably, no genotype differences were observed in input-output curves, showing that basal synaptic strength of the Schaffer collaterals is unchanged in MLC mice (Fig 7C).

Increases in $[K^+]_o$ were induced by sustained repetitive Schaffer collateral stimulation at a fixed stimulation strength for 10 seconds, either at 5, 10, or 20Hz. A stimulation frequency-dependent increase in $[K^+]_o$ was observed in all experiments (Fig 7D,E). However, significantly higher stimulation-induced increases in $[K^+]_o$ were measured both in Mlc1-null (Fig 7F) and in Glial-cam-null mice (Fig 7G). In contrast, the sum of all fEPSP amplitudes measured during repetitive stimulation was not different between genotypes (Fig 7F,G). Observed differences in peak $[K^+]_o$ were stimulation frequency-dependent, with 20-Hz stimulation showing the strongest increase in peak $[K^+]_o$ (Fig 7F,G). Despite the fact that MLC mice showed a significantly higher increase in $[K^+]_o$ during the stimulation, the poststimulation $[K^+]_o$ recovery (80–20% decay time) was not different between MLC mice and wild-type littermates (data not shown).

These findings show that while the strength and efficacy of Schaffer collateral synapses are unchanged in MLC, prolonged high-frequency stimulation of these synapses leads to a larger increase in $[K^+]_o$ in MLC mice than in wild-type littermates.

**Increased Network Excitability in CA1 Somatic Layer of MLC Mice**

Increased $[K^+]_o$ in the dendritic layer upon repetitive stimulation could lead to stronger and more sustained depolarization of the dendritic tree and may therefore result in increased network excitability. After finishing recording in stratum radiatum, we moved recording electrodes into the CA1 cell body layer (stratum pyramidale) while keeping the stimulation electrode at the same position (Fig 8A). When determining the input-output curve in stratum pyramidale with single stimulations, both MLC mice showed a trend toward an increase in population spikes (pop-spikes; Fig 8B), although this did not reach significance (Fig 8B,C). Similar to what was observed in stratum radiatum, prolonged Schaffer collateral stimulation led to larger increases in $[K^+]_o$ in both MLC mouse models (Fig 8C), indicating increased network excitability in MLC mice (Fig 8F,G). Strikingly, the sum of all pop-spike areas evoked during the train stimulation was also increased in both MLC mouse models (Fig 8C), indicating increased network excitability in MLC mice (Fig 8F,G). Similar to what we observed in stratum radiatum, the poststimulation $[K^+]_o$ recovery (80–20% decay time) in stratum pyramidale was not different between MLC mice and wild-type littermates (data not shown).

In conclusion, $[K^+]_o$ recordings show that, while Schaffer collateral input strength is unchanged,
prolonged synaptic stimulation leads to higher rises in \([K^+]_o\) in CA1, both in stratum radiatum and in stratum pyramidale. Furthermore, there is an increase in stimulation-induced pop-spikes in CA1, suggesting increased excitability of neuronal networks in MLC mice.

FIGURE 7: Larger stimulation-induced rises in \([K^+]_o\) in CA1 stratum radiatum of MLC mice. (A) Schematic drawing of the hippocampus showing recording and stimulating electrodes. (B) Basal \([K^+]_o\) in stratum radiatum (wildtype [blue]: 3.0 ± 0.06mM; n = 10 vs Mlc1-null [red]: 3.1 ± 0.04; n = 10; p = 0.63; wildtype [blue]: 3.1 ± 0.08; n = 11 vs Glialcam-null [green]: 3.1 ± 0.10; n = 11; p > 0.99). (C) Input-output curve showing fEPSP amplitude vs stimulation current. Curves did not differ between MLC mice and littermate controls (wildtype vs Mlc1-null: p = 0.47; wildtype vs Glialcam-null: p = 0.14) (D,E) Top: averaged trace of \([K^+]_o\) upon 10-second stimulation (gray area) at 5, 10, and 20Hz. Bottom: averaged fEPSP amplitude for each stimulation pulse during 5, 10, and 20Hz stimulation. (F,G) Significant stimulation frequency-dependent increase in \([K^+]_o\) in MLC mice (wildtype vs Mlc1-null: p = 0.034; wildtype vs Glialcam-null: p = 0.013) with no significant change in fEPSP amplitude sum (wildtype vs Mlc1-null: p = 0.087; wildtype vs Glialcam-null: p = 0.29). Thin lines show individual experiment. Error bars and shaded regions indicate SEM. *p < 0.05; **p < 0.01. MLC = megalencephalic leukoencephalopathy with subcortical cysts.
FIGURE 8: Increased stimulation-induced \([K^+]_o\) rises and network excitability in CA1 stratum pyramidale of MLC mice. (A) Schematic drawing of the hippocampus showing recording and stimulating electrodes. (B) Representative field recording upon single Schaffer collateral stimulation (450 µA) in a wildtype mouse (blue) and an Mlc1-null mouse (red). The pop-spike area was defined as the shaded region in the traces. (C) Pop-spike area versus stimulation current. Although a trend towards larger pop-spike area was observed this did not reach significance (wildtype [blue; n = 10] vs Mlc1-null [red; n = 10]: \(p = 0.053\); wildtype [blue; n = 11] vs Glialcam-null [green; n = 11]: \(p = 0.076\)). (D,E) Top: averaged trace of \([K^+]_o\) upon 10-second stimulation (gray area) at 5, 10, and 20Hz. Bottom: averaged pop-spike area for each stimulation pulse during 5, 10, and 20Hz stimulation. (F,G) Significant stimulation frequency dependent increase in \([K^+]_o\) in MLC mice (wildtype vs Mlc1-null: \(p = 0.021\); wildtype vs Glialcam-null: \(p = 0.035\)). Total summed pop-spike area was also significantly increased in MLC mice (wildtype vs Mlc1-null: \(p = 0.015\); wildtype vs Glialcam-null: \(p = 0.033\)). Thin lines show individual experiment. Error bars and shaded regions indicate SEM. *\(p<0.05\); **\(p<0.01\). MLC = megalencephalic leukoencephalopathy with subcortical cysts.
Discussion
We have shown that an early onset of epileptic seizures is common in MLC patients. We studied the cellular pathophysiology of seizures in two different genetic mouse models for MLC. Both Mlc1-null and Glialcam-null mice show hindlimb claspings, unprovoked interictal brain activity, and a lowered threshold for kainate-induced seizures. While intrinsic excitability of principal neurons is unchanged in MLC mice, increases in $[K^+]_o$ upon network activation and network excitability are higher in both MLC mouse models. Thus, astrocyte dysfunction in MLC leads to disturbed $[K^+]_o$ dynamics and network hyperexcitability, and lowers the threshold for seizures.

Seizures in MLC Patients and MLC Mouse Models
This is the first report on seizure characteristics in a large cohort of genetically confirmed patients with recessive MLC. An estimated 75% of all MLC patients experience at least one seizure, and 63% of patients older than 6 years has been diagnosed with epilepsy, which is in line with previous reports, and confirms that epilepsy is common in MLC. Epilepsy in MLC patients is mostly well controlled with medication compared to the general epilepsy population (only 14% had moderately to poorly controlled epilepsy versus 20–30% of general epilepsy patients). However, occurrence of status epilepticus is high (~17%) compared to the overall occurrence reported for children and adults with epilepsy during follow-up (~10%), indicating that once seizures are initiated, restoring neuronal network activity to its normal state is hampered in MLC.

A striking feature of MLC is the high susceptibility to develop seizures immediately following mild head injury (54% of patients with seizures). In patients with epilepsy in general, mild head trauma is only rarely reported as a seizure-precipitating factor. Interestingly, animal studies show that closed head injury leads to a rise in brain $[K^+]_o$. The high sensitivity to mild head injury of MLC patients could therefore reflect an increased sensitivity to rises in $[K^+]_o$, which would be in line with our observations in mouse models for MLC (see below).

Our results in Mlc1-null and Glialcam-null mice confirm that epileptiform brain activity and a lowered seizure threshold are an integral part of MLC. However, because overt spontaneous behavioral seizures are absent in both mouse models for MLC, the phenotype of MLC mice is milder than that of MLC patients. This discrepancy between MLC mice and patients has been discussed before and might be related to different compensation by mice to loss of MLC1 function compared to humans. Another likely contributing factor is the much shorter life-span of mice than that of humans, given that MLC patients develop major neurological dysfunction only after a delay of several years to decades.

Cellular Basis of Epilepsy in MLC
The use of mouse models allowed us to investigate the cellular basis of epilepsy in MLC. Decreased seizure threshold is often associated with increased neuronal excitability. However, we did not observe any change in intrinsic neuronal excitability of either hippocampal CA1 pyramidal neurons or cortical L5 pyramidal neurons in MLC mice that could explain the decreased seizure threshold. We conclude that intrinsic hyperexcitability of principal neurons is not the cause of seizures in MLC.

$K^+$-sensitive electrode recordings revealed increased peak $[K^+]_o$ rises in hippocampal CA1 upon trains of Schaffer collateral stimulation in MLC mice. This was not attributed to increased synaptic strength, given that fEPSP amplitudes were unchanged. When recording population responses to the same Schaffer collateral stimulation in stratum pyramidale, we observed an increase in CA1 pyramidal neuron population spiking, indicative of increased network excitability. In line with this, it was previously shown that modest rises in $[K^+]_o$ increase hippocampal population spiking while leaving levels of synaptic transmission unaffected. Therefore, we conclude that although intrinsic excitability of individual neurons is unchanged, MLC mice show altered $[K^+]_o$ dynamics and increased network excitability.

How could loss of function of the astrocyte-specific protein MLC1 lead to disturbed $[K^+]_o$ dynamics and seizures? It is well established that astrocytes are crucial for $[K^+]_o$ homeostasis. Mathematical models show that disturbing glial $K^+$ uptake can lead to seizures, and dysfunctional astrocyte $[K^+]_o$ homeostasis is implicated in epilepsy. $K^+$ homeostasis is tightly linked to the homeostasis of other ions, and MLC1 and GlialCAM colocalize and/or interact with several proteins involved in ion and water homeostasis. This includes the dystrophin glycoprotein complex, the swelling-sensitive cation channel, TRPV4, and the gap-junction protein, connexin 43. MLC1 and GlialCAM may therefore play a central role in organizing these components necessary for astrocyte ion and water homeostasis.

In the healthy brain, astrocytes counteract increases in $[K^+]_o$ during periods of high neuronal activity through transporter-mediated $K^+$ uptake and spatial $K^+$ buffering. This leads to astrocyte swelling and depolarization, which can be counteracted by activation of VRACs. We and others have previously shown that...
mutations in MLC1 or GLIALCAM lead to dysfunction of astrocytic VRACs,\(^7\)–\(^10\) resulting in chronic astrocyte swelling. We previously hypothesized that this could limit the K\(^+\) uptake capacity of astrocytes, thereby leading to activity-dependent white matter vacuolization.\(^8\) In line with this hypothesis, our study confirms that [K\(^+\)]\(_o\) is disturbed in MLC. We show that this is accompanied by network hyperexcitability and seizures. The fact that MLC patients show a high occurrence of status epilepticus fits in this framework, given that a disturbed uptake of K\(^+\) likely hampers restoration of disturbed ion gradients following a seizure.

MLC1 and GlialCAM share their localization in astrocyte endfeet abutting blood vessels with inwardly rectifying K\(_r\)/4.1 K\(^+\) channels and AQP4 water channels. Both these proteins are necessary for [K\(^+\)]\(_o\) regulation and have been linked to seizures. K\(_r\)/4.1 is crucial for K\(^+\) spatial buffering\(^49–51\) and for efflux of K\(^+\) from astrocytes following active transport.\(^48\) Astrocyte-specific K\(_r\)/4.1 conditional knockout results in reduced astrocyte function and seizures.\(^43,53\) In line with this, our study demonstrates that loss of function of the astrocyte-specific protein MLC1 leads to dysfunction of astrocytes for neuronal network functioning and form an important next step in our understanding of the cellular pathophysiology of MLC.

Acknowledgment
This study was financially supported by E-Rare (11-330-1024), the Dutch Hersenstichting (2009(2)-14), an NWO Spinoza award (2008), and an Amsterdam Neuroscience proof-of-concept grant (2014).

We thank all families and physicians who contributed to the inventory of clinical characteristics. We thank Prof Juha Voipio for advice on potassium-sensitive electrode recordings, Dr Thijs Verhoog for help with action potential analysis, Dr Kevan Hashemi for ECoG analysis software, and Anton Pieneman, Dr Tim Heistek, and Hans Lodder for technical assistance.

Author Contributions
M.D., E.B., H.D.M., M.S.v.d.K., and R.M. contributed to the conception and design of the study. M.D., E.B., E.M.C.H., O.S., M.B., H.K., M.H.P.K., U.B., R.C.W., H.D.M., M.S.v.d.K., and R.M. contributed to the acquisition and analysis of data. M.D., E.B., E.M.C.H., M.S.v.d.K., and R.M. contributed to drafting the text and preparing the figures.

Potential Conflicts of Interest
Nothing to report.

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