High-frequency voltage oscillations in cultured astrocytes

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Abstract

Because of their close interaction with neuronal physiology, astrocytes can modulate brain function in multiple ways. Here, we demonstrate a yet unknown astrocytic phenomenon: Astrocytes cultured on microelectrode arrays (MEAs) exhibited extracellular voltage fluctuations in a broad frequency spectrum (100–600 Hz) after electrical stimulation. These aperiodic high-frequency oscillations (HFOs) could last several seconds and did not spread across the MEA. The voltage-gated calcium channel antagonist cilnidipine dose-dependently decreased the power of the oscillations. While intracellular calcium was pivotal, incubation with bafilomycin A1 showed that vesicular release of transmitters played only a minor role in the emergence of HFOs. Gap junctions and volume-regulated anionic channels had just as little functional impact, which was demonstrated by the addition of carbenoxolone (100 µmol/L) and NPPB (100 µmol/L). Hyperpolarization with low potassium in the extracellular solution (2 mmol/L) dramatically raised oscillation power. A similar effect was seen when we added extra sodium (+50 mmol/L) or if we replaced it with NMDG⁺ (50 mmol/L). The purinergic receptor antagonist PPADS suppressed the oscillation power, while the agonist ATP (100 µmol/L) had only an increasing effect when the bath solution pH was slightly lowered to pH 7.2. From these observations, we conclude that astrocytic voltage oscillations are triggered by activation of voltage-gated calcium channels and driven by a downstream influx of cations through channels that are permeable for large ions such as NMDG⁺. Most likely candidates are subtypes of pore-forming P2X channels with a low affinity for ATP.

Introduction

Regarded as merely passive glue to stick neurons together a hundred years ago, glial cells are nowadays appreciated for their diverse and sophisticated functions within the central nervous system. Especially, astrocytes – the most abundant cell type in the mammalian CNS – gain more and more attention for maintenance but also strong interaction and modification of neuronal function. The term “tripartite synapse” was established when it became clear that the flow of information between pre- and postsynapse is modulated by astrocytes as a third partner. Astrocytes take up potassium, purge the synaptic cleft from the neurotransmitter glutamate, replenish neurotransmitter pools by providing neurons with the precursor glutamine, and affect the synaptic plasticity of neurons via diverse mechanisms (Barker and Ullian 2010).

Astrocytes are regarded as electrically passive in comparison to neurons, and it has been widely accepted that they process information mainly via changes in intracellular calcium: They react to certain stimuli with calcium oscillations that trigger a multitude of downstream signaling events. These observations merged into a concept of astrocytic calcium excitability. Many excellent reviews have been published that summarize signaling pathways of astrocytes and their interaction with neurons (Araque and Navarrete 2010; Halassa and Haydon 2010; Santello et al. 2012; Zorec et al. 2012; Ota et al. 2013).

Increased cytosolic Ca²⁺ levels in astrocytes can be triggered by electric (Hassinger et al. 1996) or mechanical...
stimulation, (Guthrie et al. 1999; Stout et al. 2002) by glutamate (Cornell-Bell et al. 1990) or ATP (Jeremic et al. 2001, 2001), by bradykinin (Parpura et al. 1994; Liu et al. 2009), by hyperosmolarity (Reetz et al. 1997; Pangrsic et al. 2006) or by low-extracellular Ca\(^{2+}\) (Zanotti and Charles 1997). A transient increase in intracellular Ca\(^{2+}\) can lead in return to the release of neurotransmitters like ATP, adenosine, or glutamate and of neuromodulators like taurine and D-serine. These substances are thus summarized under the term "gliotransmitters".

A variety of possible release sites for these gliotransmitters has been discussed. There are indications that astrocytes possess an exocytotic vesicular release machinery similar to neurons, but release of transmitters was also demonstrated through hemichannels and through voltage-regulated anionic channels (VRAC) after cell swelling. There is also evidence that ATP and glutamate leave the cells through membrane channels that open upon prolonged activation as large diameter pores: Some members of the ATP-gated P2X receptors are able to form pores that permit passage of molecules as large as fluorescent dyes.

Calcium signaling is not restricted to single astrocytes but covers smaller cell assemblies by propagating calcium waves. The presence of calcium-waves originating from one astrocyte to surrounding cells was first reported by Cornell-Bell et al. in 1990; where they were elicited by extracellular glutamate (Cornell-Bell et al. 1990). The observation that ATP induces Ca\(^{2+}\) influx into astrocytes was already made in the 1980s but it was Guthrie et al. 1999 who claimed that ATP serves as an extracellular messenger released from astrocytes to mediate calcium waves (Guthrie et al. 1999). These calcium waves cover a distance from 100 to 250 \(\mu\)m and are either propagating from cell to cell via gap junctions or regenerated in each cells after extracellular stimulation with ATP (Bennett et al. 1991; Finkbeiner 1992; Bowser and Khakh 2007). Cell swelling of astrocytes in connection with brain oedema also triggers Ca\(^{2+}\) signaling, at least partly dependent on activation of purinergic receptors by ATP (Darby et al. 2003; Nase et al. 2008; Akita et al. 2011; Thran et al. 2011).

Astrocytes participating in a Ca\(^{2+}\) wave show a distinct oscillatory pattern in intracellular Ca\(^{2+}\) with decreasing frequency (Charles et al. 1991; Pasti et al. 1997). But compared to neuronal signaling, astrocystic Ca\(^{2+}\) oscillations are very slow. Commonly used sampling frequencies used for calcium imaging are in the range from 0.5 to 10 Hz.

High-frequency voltage oscillations (250–400 Hz) have been observed in cortical and hippocampal cell cultures upon electrical stimulation on microelectrode arrays (MEAs), but were not assigned to a certain cell type (Hales et al. 2012). This represented a new application field for MEAs that were so far used to detect action potentials in dissociated neuronal or cardiomyocyte cell cultures or to measure summated postsynaptic potentials in the form of local field potentials in slices.

In this work, we show that astrocytes are the cellular source of very fast and long-lasting extracellular voltage oscillations in reaction to an electrical stimulus. Not strictly periodic, these oscillations cover a broad frequency spectrum up to 600 Hz with decreasing frequencies over time. The oscillations proved to be very susceptible to changes in the extracellular environment.

**Methods**

**Ethical approval**

All animal procedures were performed in accordance with the German animal care and ethics legislation and were reported to the local government authorities.

**Cell cultures**

Mixed cortical cultures where prepared from newborn (P0) Wistar rats of both genders provided by the local animal laboratories. Pups were anaesthetized with isoflurane and killed by decapitation. Cortices were removed and cut into three coronal slices in phosphate-buffered saline (PBS). Cortex tissue was dissected from the middle slice which contained the striata and the anterior part of the thalamus. Cortical tissue was collected and cut into small pieces with a scalpel and then transferred into trypsin (0.05 %; 37°C for 5 min). Cells were washed several times in PBS and medium (Dulbecco’s Modified Eagle’s medium [DMEM] with 25 mmol/L glucose and 25 mmol/L HEPES [4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid]), supplemented with 10 % fetal calf serum and 25 \(\mu\)g/mL insulin) and triturated several times with a pipette. Dissociated cells were counted and one drop of the cell suspension with a density of 1 \(\times\) 10^6/mL was plated on poly-D-lysine- and laminin-coated microelectrode arrays (Multi Channel Systems, Reutlingen, Germany). MEAs were placed into a 5 % CO\(_2\) incubator with 37°C and 90 % humidity. The next day, 1.5 mL of B27-neurobasal medium with the mixture of penicillin and streptomycin (Pen/Strep), glutamine (500 \(\mu\)mol/L), and Basic fibroblast growth factor (2 ng/mL) was applied onto the MEAs.

Cultured cortical astrocytes were also prepared from P0 Wistar rats. After a 10 day proliferation phase in T75 flasks, astrocytes had grown confluent and were detached from the flasks with trypsin (0.05 %; 37°C for 2 min) and after several washing steps plated on poly-D-lysine-coated MEAs or coverslips. Astrocytes were maintained in DMEM with high glucose and GlutaMAX (61965, Life Sciences).
Technologies, Carlsbad, CA) with 10 % horse serum, 1 x Pen/Strep, 1 mmol/L sodium pyruvate in an incubator with 10 % CO₂ (37°C and 90 % humidity).

Mouse embryonic fibroblasts (MEF) were obtained from E13 embryos. Briefly, head and inner organs were removed; the remaining tissue was minced, trypsinized, and triturated with a pipette. Cells were then plated on gelatine coated flasks, cultured until confluence, and then frozen in liquid nitrogen. Upon thawing and seeding on MEAs, fibroblasts were maintained in the same nutrition medium as the astrocyte cultures. MEF were not tested for their purity with immunostainings against the marker protein vimentin.

**Immunostainings**

Cultures were fixed with 4 % paraformaldehyde for 1 h at time points when parallel cultures on MEAs exhibited high-frequency oscillations. We used the following antibodies: monoclonal Anti-Glia Fibbrillary Acidic Protein (GFAP) antibody produced in mouse (G3893, Sigma-Aldrich, St. Louis, MO) and Neuronal Class III β-Tubulin Rabbit Monoclonal Antibody (MRB-435P, Covance, Princeton, NJ). Secondary antibodies were labeled with the fluorescent dyes Alexa 488 and Cy3, respectively.

**Recording and stimulation**

Extracellular recording of electric activity in dissociated cell cultures was performed with the MEA2100-System (Multi Channel Systems, Reutlingen, Germany) which provides an integrated stimulus generator and temperature control. We used microelectrode arrays (MEAs) with a square grid of 60 titanium/titanium nitride alloy (Ti/TiN) electrodes (30 μm diameter, 200 μm spacing) and an input impedance of <50 kΩ according to the manufacturer (Multi Channel Systems). The standard data acquisition software MC_Rack was used for visualizing and storing the extracellular potentials measured. Signals were sampled simultaneously at 25 kHz from all 60 electrodes. To attenuate stimulation artifacts, a second order high-pass filter with a cut off frequency of 100 Hz was used in the MEA2100 prior to signal amplification. As we used the same electrodes for stimulation and recording, this procedure enabled us to record a stable extracellular signal just about 50 ms after stimulus delivery. In standard measurements, we applied biphasic square wave voltage pulses (800 mV; 200 μs/phase) with the negative phase first to each of the MEA electrodes (except the ground electrode). Prior to stimulation experiments, cultures were left at rest in the amplifier headstage to equilibrate for 10 min. In our experimental paradigm, each recording started with a stimulus-free baseline period of 30 sec, followed by 10 voltage pulses delivered every 20 sec, with a total recording length of 240 sec.

Recordings were made at 37°C in buffered salt solution, containing 100 mmol/L NaCl, 10 mmol/L glucose, 10 mmol/L HEPES hemisodium salt, 5 mmol/L KCl, 500 μmol/L MgCl₂, and 1 mmol/L CaCl₂ (unless otherwise indicated). The rather low osmolarity of the buffer was initially chosen to resemble that of the cell culture medium for neurons (Neurobasal Medium). Different pH values were adjusted by adding several microliters of 1 mol/L NaOH or HCl, respectively, to the recording solution.

Cell culture ingredients were purchased from Life Technologies and all drugs from Sigma-Aldrich, except (3S)-3-[(3-[4-(Trifluoromethyl)benzoyl]amino)phenyl]methoxy]-1-aspartic acid (TFB-TBOA) (Tocris Bioscience, Bristol, UK), gabazine, D-(-)-2-Amino-5-phosphonopentanoic acid (D-APV), and 6-cyano-7-nitroquinoxaline-2,3-dione, D-APV; D-(-)-2-Amino-5-phosphonopentanoic acid (CNQX) (Abcam, Cambridge, UK).

**Data analysis**

Spikes and bursts were detected by custom-built software (RESULT Medical GmbH, Düsseldorf, Germany). The threshold for spike detection was set to 6.2 standard deviations of the average noise amplitude for each electrode. The actual temporal clustering of spikes in each channel was compared with a Poisson process of independently occurring spikes at the same firing rate. If the Poisson probability of an observed cluster with \( n \) > 3 spikes within a time interval \( \Delta t \) was below 0.005, this cluster was considered as burst.

For the analysis of oscillations, continuous raw data and trigger time stamps of the recordings were imported from Multichannel Systems MCD files into MATLAB using the 64 bit Neuroshare library (version 3.7b). Individual electrodes were selected for analysis based on a clear response to stimuli in form of prolonged increased signal amplitude and controlled by visual inspection (Fig. 1B, lower panel).

In order to accurately retain the temporal relation between signal and stimuli, data were subsequently down-sampled to 2500 Hz after using a zero-delay linear phase finite impulse response low-pass filter of order 400 with 1125 Hz cut-off frequency. The actual time-frequency analysis of oscillatory activity was performed using the ft_freqanalysis function of the Field-trip toolbox (version 20130512; Oostenveld et al. 2011). To suppress stimulus artifacts, the raw signal was set to zero during 50 ms after each stimulus prior to spectral analysis. Each recording contained responses to ten stimuli occurring at 30, 50, . . . , 210 sec.
The time-frequency representation of the signal after each stimulus was calculated by the convolution with wavelets for time points every 50 ms and at 300 equidistant frequencies from 2 to 600 Hz. The 2 Hz frequency interval required a corresponding 0.5 sec-wide time window for the spectral power estimation. A power value calculated at a given time thus contained signal information of the surrounding 0.5 sec. Spectral analysis used a sliding temporal Hanning window centered at each time of interest and power spectra were obtained separately for each stimulus trial.

The 30 sec long baseline period was split into six nonoverlapping pseudo-trials of 5 sec each, starting at 0, 5, ... , 25 sec. Baseline power spectra were estimated in the same way as for the actual stimulus trials, and were subsequently averaged over the six pseudo-trials and over time, yielding only frequency-dependent baseline power spectra.

Finally, time- and frequency-dependent power spectra obtained from ft_freqanalysis were averaged over all ten stimulus trials, and the corresponding time-independent pseudo-trial averaged baseline power spectra were subtracted from these averaged spectra. The resulting “power over baseline” spectra were plotted as color coded maps over time and frequency.

To calculate the duration of an oscillation, spectral power was summed over all frequencies from 2 to 600 Hz, and a double exponential decay function was fitted to these total power over baseline data which are only time-dependent. The duration was then calculated as the time elapsed until the function fitted to the power had dropped to $\exp(-\frac{4}{4}) \approx 1.8\%$ of its amplitude over background noise, which was typically at least two orders of magnitude smaller.

In order to quantify high-frequency oscillations (HFOs) by a single power estimator, we defined the power of a HFO as the sum of the trial-averaged total power over baseline for all points in time between zero and the duration of the oscillation.

In order to locate HFOs on the frequency axis, we also calculated the power-weighted median frequency in the first 0.5 sec where the oscillations were most pronounced. We used the median frequency instead of the mean, since the power distribution was heavily right-tailed and skewed towards high frequencies.

**Statistics**

For all drug-related experiments, only those MEA electrodes were chosen for evaluation that showed a strong response to the applied electrical stimulus under control conditions (HFO amplitude $> 3\times$ root mean square noise), clearly distinct from the average noise amplitude. This assessment was performed by visual inspection, with no regard to the response these specific electrodes showed after pharmacological treatment. In 3.3, we compared stimulus-induced HFOs of astrocyte cultures with those of fibroblast cultures or MEAs in nutrition medium but without any attached cells. Here, we averaged power over all stimulated electrodes irrespective of HFO amplitude.

Because HFO power distribution was not normally distributed (see Fig. 3), we presented the data graphically as median with upper and lower quartiles. Statistical significance in pharmacological experiments was calculated with the Wilcoxon-matched-pairs signed rank test over indi-
individual electrodes from the utilized MEAs, before and after treatment, resp. In one case where we compared two substance effects with each other [Na+- vs. NMDG+-treatment], we used the Mann–Whitney test because these observations were not paired events.

For better comprehensibility absolute HFO power was normalized to both the median value under control conditions and the mean of control. Generally, substance effects in the result section are presented as “normalized median power over baseline in %; normalized mean/C6SD; P value of the Wilcoxon matched-pairs signed rank test; number of evaluated electrodes”.

Results

Mixed cortical cell cultures exhibit high-frequency voltage oscillations

Our recordings in cortical cells cultured on microelectrode arrays (MEAs) confirmed the data presented by Hales et al. (2012): Electrical stimulation evoked voltage oscillations in a broad and continuous frequency spectrum up to 600 Hz. The amplitudes of these oscillations were comparable to extracellularly measured neuronal action potentials. Considerably, fewer electrodes showed these oscillations than spontaneous bursts of action potentials. However, we could not assign certain electrodes to one of the phenomena: Some electrodes showed oscillations but no bursts, others had bursts but no oscillations and a third group showed oscillations that were superimposed by spike bursts. A stimulus applied to one specific electrode caused a network burst on all electrodes, but HFOs were restricted to the electrode that had been stimulated (Fig. 1).

The duration and amplitude of these oscillations clearly depended on the power of the stimulus: oscillations grew stronger when either stimulus duration or amplitude was increased. When the stimulus was gradually reduced, oscillations became weaker until they were undistinguishable from the noise level. No threshold stimulus for the HFOs could be determined.

In accordance with Hales et al. (2012), we saw that oscillations developed over cultivation time, which might imply that single cells are not able to generate HFOs and maturing interconnections between cells are necessary. Interestingly, we observed synchronous network bursts covering the whole MEA some time before we were able to elicit HFOs on selected electrodes. On the other hand, oscillations persisted when spike activity started to diminish due to neuronal death. HFOs also survived during treatment with TTX and the synaptic blockers D-APV (20 μmol/L), CNQX (10 μmol/L) and gabazine (10 μmol/L) that blocked all spike activity and fast synaptic transmission. All these observations raised the suspicion that the cellular source of HFOs might not be neurons but a different cell type that is tightly connected with neuronal function, probably astrocytes. All further experiments within this work were therefore performed in astrocyte-enriched cultures with a negligible amount of neurons.

High-frequency oscillations are generated by astrocytes

In such astrocyte-enriched cultures, a small amount of neurons usually survives the purification process that is performed to isolate astrocytes. Thus, astrocyte cultures are rarely 100 % “pure”. But immunostainings with the neuronal marker protein β-tubulin III confirmed that the astrocyte cultures used for our studies showed only a negligible amount of “contaminating” neurons (Fig. 2) but a dense layer of GFAP (glial fibrillary acidic protein)-positive cells. GFAP is highly upregulated in astrocytes in culture. In addition, we never observed any spontaneous spike activity in cultured astrocytes on MEA. These observations make it highly unlikely that neurons are the cellular source of the observed voltage oscillations.

When biphasic voltage pulses were administered through all MEA electrodes (the grounding electrode excluded), nearly every electrode that was covered with cells exhibited high-frequency oscillations (HFOs) in return (Fig. 3). Accordingly, we conclude that astrocytes generated the electrically triggered HFOs that had been observed in cortical and hippocampal cell cultures before.

In contrast to Hales et al. (2012), our power spectral analysis of the oscillations revealed no distinct frequency band. We used a hardware high-pass filter (>100 Hz) in our measurements and saw a rather wide and smooth fre-
frequency distribution with power decreasing at higher frequencies. Thus, the oscillations were rather aperiodic, and oscillation power correlated with the oscillation duration.

In 53 independent control recordings containing a total of 905 electrodes with HFOs, the median duration was 2.4 s (mean ± SD: 2.7 ± 1.3 sec), and the median oscillation frequency was 142 ± 28 Hz. HFO power summed for all points in time until total power over baseline had dropped to 1.8 % was $2.9 \times 10^{-10}$ V² (mean ± SD: $4.4 \pm 4.0 \times 10^{-10}$ V²; Fig. 4).

The measured HFOs were constrained to a small area around the stimulated area. Electrodes could not detect voltage oscillations when a neighboring electrode in a distance from 200 μm was used for the stimulation. Oscilla-

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**Figure 3.** Astrocytic HFOs are locally restricted to the source of the stimulus. (A) When a stimulus was applied to the astrocyte culture via every MEA electrode (except ground), virtually all electrodes showed HFOs with varying amplitudes and durations in response. (B) When the stimulus was applied to a subset of electrodes (green frames), only cells in close proximity of the stimulated electrodes responded with HFOs. This indicates that (1) the stimulus was not passively conducted through the recording solution, and that – unlike in neuronal cultures – (2) there was no active transfer of the signal over longer distances within the cellular layer. The MEAs we used in this study featured 200 μm distance between neighboring electrodes.
tions were only detected by the same electrode that was used for the stimulation (Fig. 3).

**High-frequency oscillations are not electrical artifacts**

We performed several control experiments to rule out the possibility that the observed HFOs were merely electrical artifacts caused by amplifier electronics, electrodes, or medium.

Five MEAs with nutrition medium but without any attached cells were stored in the incubator for 3 weeks and regularly checked for HFOs. At no time point and on no electrode could an electrical stimulus induce visible voltage perturbations that exceeded the immediate but short-lived stimulus artifact (> 30 ms). In these experiments, the stimulus trigger-averaged signal power summed up in a 9 sec time window represents the combined physicochemical and electrical noise of the ionic solution, Ti/TiN electrodes and MEA amplifier.

To test for cell-type specificity of HFOs we seeded mouse embryonic fibroblasts (MEF) on eleven MEAs – a cell type usually used as feeder layer for embryonic stem cells. The fibroblasts were seeded in high density to ensure a closed cell layer across the electrode array, cultured for 3 weeks, and regularly checked for the presence of HFOs after electric stimulation. Small amplitude HFOs could be observed on few electrodes of some MEAs. As a positive control we seeded astrocytes on ten MEAs. HFO responses to electric stimuli were clearly visible only 3 days after seeding.

For these three conditions – medium, MEF, astrocytes – HFO power for each stimulated electrode (59 per MEA) was summed up over a time period of 9 sec after the stimulus. The median stimulus-averaged power of responses was $5.3 \times 10^{-13} \text{V}^2$ for nutrition medium.
without cells (531 electrodes; 9 MEAs), $1.4 \times 10^{-12}$ $V^2$ for MEF cultures (1298 electrodes, 11 MEAs at two points in time) and $2.5 \times 10^{-14}$ $V^2$ for astrocyte cultures (590 electrodes, 10 MEAs). Thus, the power of HFOs generated by astrocytes was 4800 times higher than the physico-chemical noise (recordings without cells) and 1800 times higher than voltage oscillations generated by fibroblasts (Fig. 5A).

In a next set of control experiments, we measured HFO responses of astrocyte cultures before and after treatments that should kill the cells but leave an intact layer of biological material covering the MEA electrodes. In a first approach, we added 10% of Dimethyl sulfoxide (DMSO) to the nutrition medium. HFOs were recorded before and 2 h after addition of DMSO. In a second approach, we induced a hyperosmotic shock to harm astrocyte integrity: Control

Figure 5. High-frequency oscillations are not electrical artifacts but generated by vital astrocytes. (A) Ten single rectangular voltage stimuli (800 mV, 200 $\mu$s/phase) were applied to all 59 recording electrodes of the MEA. Signal power was summed over a time window of 9 sec after the stimulus and averaged over all 10 stimuli. The left-hand graph depicts median power and interquartile range of the voltage oscillations elicited on all stimulated electrodes. The combined physicochemical and electronic noise of the ionic solution, Ti/TiN electrodes and amplifier was determined with MEAs that were covered with nutrition medium but without any attached cells. Only very slight oscillations were observed when we stimulated mouse embryonic fibroblasts cultured on MEAs. Stimulation of astrocytes at 3 div caused strong voltage oscillations on many electrodes that exceeded the noise level (“no cells”) nearly 5000 times. Average power spectra of the respective measurements are given on the right hand side. The logarithm (base 10) of the time- and frequency-dependent power over baseline is color-coded in the range $10^{-14}$ to $10^{-12}$ $V^2$. The horizontal time axis covers 9 sec after the stimulus. Frequency (0–600 Hz) is plotted on the vertical axis. (B) Astrocyte cultures were recorded before and 2 h after the addition of 10% DMSO – a concentration that is supposed to kill cells. The median power of the stimulus-induced oscillations decreased dramatically, presumably to noise level. The respective power spectra in a 9 sec window after the stimulus are depicted on the right hand side of the panel (0–600 Hz; $10^{-14}$ to $10^{-12}$ $V^2$). (C) Astrocytes were exposed to hyperosmotic shock by transferring them from recording solution with 100 mmol/L NaCl to 300 mmol/L NaCl. Recordings were performed before and 2 h after the presumably deadly osmotic shock. No HFOs could be elicited after the treatment. Power spectra on the right hand side clearly demonstrate the collapse in oscillation power after the death of the astrocytes (0–600 Hz; $10^{-14}$ to $10^{-12}$ $V^2$).
recordings of HFOs were performed in standard recording solution with 100 mmol/L NaCl. We then applied a recording solution with 300 mmol/L NaCl to the cells. HFOs were measured again after 2 h of osmotic shock.

For the analysis, we preselected electrodes that presented a high signal-to-noise ratio under control conditions. No HFOs were visible after both treatments. DMSO addition caused a decrease in HFO power from a median value of $3.4 \times 10^{-9}$ to $2.7 \times 10^{-12} \text{ V}^2$, i.e., 0.08% of the initial value (178 electrodes, 4 MEAs; Fig. 5B). HFOs did not reappear after washout indicating that cells were actually dead and loss of HFO power was not due to the physicochemical properties of DMSO. Hyperosmotic shock reduced HFO power to only 0.02% of initial power, from a median value of 1.9 to $2.7 \times 10^{-13} \text{ V}^2$ (109 electrodes, 4 MEAs; Fig. 5C). These experiments clearly show that HFOs were not electrical artifacts but are generated by living cells, especially by astrocytes.

Oscillations are triggered by voltage-dependent calcium channels

As electrical stimulation reliably caused HFOs in astrocyte cultures, we next investigated the mechanism of voltage stimulus sensing in these cells. In a first attempt we used a cocktail of blockers against voltage-gated sodium channels, ionotropic glutamate receptors and GABA$_A$-receptors. This combination of tetrodotoxin (1 mmol/L), CNQX (10 mmol/L), D-APV (20 mmol/L), and gabazine (10 mmol/L) did not block the emergence of HFOs (112 %; 118 ± 22 %; $P = 0.15$; n = 28). This result precludes voltage-gated Na$^+$ channels from being mediators of the electric stimulus. Parallel experiments with neuronal cultures proved the efficacy of the substance mixture. No spontaneous spike activity was seen after exposure to the mix. Incubation with the voltage-gated potassium channel antagonist 4-aminopyridine (4-AP; 50 mmol/L) did not block the appearance of HFOs after both treatments. DMSO addition caused a decrease in HFO power from a median value of 1.9 to $2.7 \times 10^{-13} \text{ V}^2$ (109 electrodes, 4 MEAs; Fig. 5C). These experiments clearly show that HFOs were not electrical artifacts but are generated by living cells, especially by astrocytes.

Oscillations are calcium dependent

In order to investigate the role of intracellular calcium for HFOs, we incubated astrocyte cultures with the membrane-permeable chelator 1,2-Bis(2-aminophenoxy)ethane-N,N,N',N'-tetraacetic acid tetrakis(acetoxymethyl ester) (BAPTA-AM). BAPTA-AM moves into the cells and “captures” intracellular Ca$^{2+}$ ions thereby suppressing any downstream-signaling events triggered by this second messenger. To rule out that all binding sites of the chelator were already fully occupied before it entered the cell we incubated the cultures with Ca$^{2+}$-free solution and added 50 mmol/L BAPTA-AM. After 1 h incubation and washout we were not able to elicit any HFOs, neither in Ca$^{2+}$-free nor in standard bath solution, which indicates a strong dependency of HFOs on intracellular Ca$^{2+}$ (HFO power: 1 %; 0.8 %; 0.1 %; $P \leq 0.0001$; n = 29). However, as all cultures that underwent this treatment were dead on the following day we cannot be completely sure whether the complete suppression of HFOs is a proof of its calcium dependency or simply a symptom of dying astrocytes (Fig. 7A).

To get more convincing evidence for an intracellular Ca$^{2+}$-dependent signaling cascade we used BaCl$_2$ combined with or instead of CaCl$_2$ in the extracellular solution. Under both conditions, HFOs were strongly suppressed: The power of the oscillations decreased to a median value of 14 % (14 ± 3 %; $P \leq 0.0001$; n = 29) when equimolar
amounts of CaCl₂ and BaCl₂ were used (1 mmol/L each) and to 15 % (12 ± 1 %; P ≤ 0.0001; n = 74) when only Ba²⁺ but no Ca²⁺ was added to the bath solution (Fig. 7A). This inhibiting effect of BaCl₂ is probably due to the “Ca²⁺-imitating” qualities of barium: Both are divalent cations from the group of alkaline earth metals and are able to enter cells via Ca²⁺ channels. But Ba²⁺ does not act as a secondary messenger-like Ca²⁺ does.
Figure 7. High-frequency oscillations are strongly dependent on calcium. (A) HFOs completely vanished when astrocytes were pre-incubated with the membrane-permeable Ca²⁺-chelator BAPTA-AM (50 μmol/L). Substitution of extracellular Ca²⁺ with Ba²⁺ (1 mmol/L) strongly reduced HFO power. Apparently, influx of Ca²⁺ is mandatory to elicit HFOs. HFOs were as strongly suppressed when equimolar amounts of Ca²⁺ and Ba²⁺ were present in the extracellular solution. Ba²⁺ seems to be more preferably transferred across the membrane when both ions compete for it. Glutamate (100 μmol/L) intensified HFO power more strongly than DHPG, an agonist of group I metabotropic glutamate receptors. Histamine (50 μmol/L) and the muscarinic agonist carbachol (10 μmol/L) only slightly increased HFO power (**P < 0.01; ***P < 0.001). (B) Lowering the extracellular calcium concentration resulted in a dramatic increase in HFO power: 500 μmol/L tripled the power and 200 μmol/L actually increased power 17-fold compared to standard conditions with 1 mmol/L CaCl₂. The opposite effect was seen with higher calcium concentrations. Data in A and B is depicted as normalized medians with inner and outer quartiles. (C) The left hand side depicts signals of four selected MEA electrodes over a time period of 20 sec after electrical stimulation (red arrow) under control conditions (1 mmol/L CaCl₂, upper panel) and under extracellular calcium deficiency (200 μmol/L CaCl₂, lower panel). The right hand side shows the power spectra of the same MEA electrodes averaged over ten stimuli applied at 20 sec intervals. HFOs under control conditions were small in amplitude and lasted approximately 2 sec. With only 200 μmol/L Ca²⁺, the HFOs elicited by the voltage pulse on the same electrodes persisted even after the inter-stimulus interval of 20 sec. Note that HFOs on most electrodes showed no gradual decrease in amplitude but rather spontaneous rekindling. Power was maximal at frequencies below 200 Hz.

From these experiments, we conclude that HFOs can only be generated, if Ca²⁺ enters the cells and mediates internal signaling routes. Voltage-gated Ca²⁺ channels that open only after strong depolarization (high-threshold or high-voltage-activated Ca²⁺ channels) show a higher permeability for Ba²⁺ than for Ca²⁺ (Yatani et al. 1987). The strong inhibition in experiments where Ba²⁺ did not replace Ca²⁺ but was added in equimolar amounts (1 mmol/L) to the recording solution suggests that Ca²⁺ influx via such high-threshold channels is essential for HFO generation. In agreement with these results, the calcium-channel antagonist cilnidipine blocks both L- and N-subtypes of high-voltage-gated calcium channels.

To test if a rise in intracellular Ca²⁺ was sufficient to elicit HFOs we applied several agonists to the cultures that are known to raise Ca²⁺ within the cell via activation of G protein-coupled receptors: (S)-3,5-Dihydroxyphenylglycine (DHPG) is an agonist of group I metabotropic glutamate receptors, histamine increases intracellular Ca²⁺ via activation of H1 receptors and carbachol (carbamoylcholine chloride) is an agonist for muscarinic acetylcholine receptors. None of the tested substances caused spontaneous HFOs when no electric stimulus was applied. DHPG (50 μmol/L), histamine (50 μmol/L) and carbachol (10 μmol/L) slightly changed the power of the stimulus-induced HFOs (DHPG: 98 %; 149 ± 56 %, P ≤ 0.0001; n = 49) (histamine: 120 %; 127 ± 55 %, P ≤ 0.0001; n = 38) (carbachol: 104 %; 109 ± 52 %, P = 0.0081; n = 38) (Fig. 7A). The addition of glutamate caused a dose-dependent increase in HFO power (10 μmol/L: 133 %; 123 ± 16%; P ≤ 0.0001; n = 52) (25 μmol/L: 263 %; 198 ± 26 %, P ≤ 0.0001; n = 52) (50 μmol/L: 667 %; 435 ± 56 %, P ≤ 0.0001; n = 52) (100 μmol/L: 376 %; 378 ± 44 %, P ≤ 0.0001; n = 61). All these effects were highly significant. The strong effect of glutamate on HFO power even at low doses is surprising, as block of ionotropic glutamate receptors had no effect and the group I mGlur agonist DHPG only a very small effect on HFOs. Glutamate might cause this strong effect by activation and interaction of several glutamate receptor subtypes.

From these experiments, we conclude that an increase in intracellular Ca²⁺ is necessary but not sufficient for the generation of HFOs. Ca²⁺ entry via voltage-gated channels seems to play a pivotal role while other signaling components – like for instance metabotropic glutamate receptors – might modulate the shape of the elicited HFOs.

Astrocytic HFOs are blocked by extracellular Ca²⁺

Not only intracellular but also extracellular Ca²⁺ modulates numerous physiological processes. To investigate its impact we varied the CaCl₂ concentration in the recording solution. For standard control condition we used a HEPES-buffered salt solution with 1 mmol/L CaCl₂. Higher Ca²⁺ concentrations attenuated HFOs, while lower concentrations strongly augmented them: 3 mmol/L CaCl₂ in the bath solution suppressed the total power of the oscillations to a median value of 16 % (18 ± 2 %; P ≤ 0.0001; n = 91) compared to standard conditions and a reduction in the CaCl₂ concentration to 500 μmol/L tripled the HFO power (300 %; 301 ± 31 %; P ≤ 0.0001; n = 91). The duration and power of HFOs dramatically increased when external Ca²⁺ was reduced to only 200 μmol/L (1650 %; 1271 ± 139 %; P ≤ 0.0001; n = 91) (Fig. 7B). Another aspect was also striking: While the highest median frequency and power of HFOs were measured directly after the stimulus and monotonically decreased afterwards, HFOs in low Ca²⁺ conditions showed a more chaotic pattern with both increasing and decreasing median frequencies and amplitudes over time. Overall, HFOs under Ca²⁺ deficiency seemed to lack the regulation and coordination seen under conditions with higher Ca²⁺ concentrations (Fig. 7B).

With even less than 200 μmol/L Ca²⁺ in the extracellular solution, HFOs on some electrodes lasted for several minutes, so that we could not use our standard stimula-
tion paradigm with stimulus pulses every 20 s. Here we noted that only a minority of the electrodes displayed such long oscillations and that the electrodes showing the longest oscillations changed from stimulus to stimulus. This suggests that HFOs are not only locally restricted, monomorphic and inevitable reactions of one independent cell to an electrical stimulus but that there is some kind of plasticity within the astrocyte layer that shapes the characteristics of the response.

**HFOs are slightly affected by blockade of vesicular transmitter release**

The strong dependency on intracellular Ca\(^{2+}\) and the phasic HFO pattern raised the suspicion that HFOs were a correlate of synaptic-like vesicular transmitter release. Release of great amounts of anionic transmitters-like ATP or glutamate might cause voltage oscillations in the extracellular space. In astrocytes, vesicular release of the transmitters glutamate and ATP has already been demonstrated in numerous studies and was shown to be inhibited by addition of bafilomycin (Araque et al. 2000; Coco et al. 2003). Bafilomycin A1 is a selective inhibitor of the vacuolar-type H\(^+\)-ATPase (V-ATPase), thereby inhibiting the filling of vesicles with transmitters.

Incubation with bafilomycin A1 (1 \(\mu\)mol/L for 1 h) had a minor effect on astrocytic HFOs elicited by electrical stimulation: They were not completely blocked, but the oscillation power was reduced to 75 % (65 ± 8 %; \(P \leq 0.0001\); \(n = 59\)) compared to standard conditions (Fig. 8A). This result points to synaptic release as a relevant factor in HFO generation, but not as its sole driving force. Control experiments with neuronal cortical cultures were performed to prove the treatment efficacy: Incubation with bafilomycin led in all cases to a complete inhibition of spontaneous spike activity.

**HFOs are augmented by membrane hyperpolarization**

To narrow down factors that contribute to the emergence of HFOs we tested, if the oscillations were susceptible to changes in the membrane potential. Since astrocytes have a big leak conductance for potassium, their membrane potential can be efficiently manipulated by varying the extracellular K\(^+\) concentration. We tested depolarizing and hyperpolarizing concentrations. Depolarizing the membrane potential with 10 mmol/L KCl in the recording solution instead of 5 mmol/L under control conditions led to a decrease in total oscillation power to 48 % (43 ± 5 %; \(P \leq 0.0001\); \(n = 44\)). After lowering the extracellular K\(^+\) concentration from 5 to 2 mmol/L and
thus hyperpolarizing the membrane potential we saw a
dramatic increase in oscillation power to 852 %
(869 ± 193 %; \( P \leq 0.0001; n = 44; \) Fig. 8B).

This effect might seem counterintuitive because depo-
larization above a certain threshold level causes neurons
to fire action potentials by opening voltage-dependent ion
channels. The membrane potential, however, primarily
determines the driving force of ions to move into or out
of the cell. An enhancement of HFOs by a very negative
membrane potential implied that the moving ions have
equilibrium potentials at the opposite side of the scale,
probably in the high-positive range. This would include
Na\(^+\), Ca\(^{2+}\), and intracellular anions like ATP or glutamate.
To exclude the possibility that HFOs were triggered by
hyperpolarization-activated channels we administered the
blocker CsCl in a concentration of 1 mmol/L. Under
these conditions, however, HFO power only slightly
decreased to 78 % (79 ± 22 %; \( P = 0.0683; n = 18; \) Fig. 8B).

Are hemichannels involved in HFO
generation?

One putative way of ions across the membrane is via
open hemichannels. Since gap junctions are known to
be blocked by Ca\(^{2+}\), an involvement of hemichannels in
HFO generation would fit our observations of the strong
dependency of HFOs to extracellular Ca\(^{2+}\). Carbenoxo-
lone is a widely used blocker of gap junctions and
hemichannels. Under carbenoxolone (100 \( \mu \)mol/L) treat-
ment, HFOs were significantly reduced in power but not
completely abolished 63 % (58 ± 12 %; \( P \leq 0.0001; n = 62; \) Fig. 8A). Thus, hemichannels might play a role
in HFO generation by permitting ion flux across the
membrane. However, since the specificity of carbenoxo-
lone has recently been questioned, we are advised to
interpret these results with caution (Beaumont and Mac-
caferri 2011). When cortical neuronal cultures were trea-
ted with carbenoxolone, spontaneous spike activity
nearly vanished. This cannot be explained by a specific
action of carbenoxolone, since neuronal network activity
is clearly driven by synaptic activity and not by gap
junctions.

Volume-regulated anion channels are not
involved in HFO generation

Another way of releasing anions from the cell into the
extracellular space is via volume-regulated anion channels
(VRACs). Glutamate and ATP were already shown to be
released from astrocytes via VRACs (Darby et al. 2003;
Anderson et al. 2004; Takano et al. 2005; Kimelberg et al.
2006). VRACs are activated by changes in the cell
morphology, for instance by cell swelling, and are perme-
able for chloride and small organic anions. Astrocytes
show fast changes in cell volume and have elaborate
mechanisms for its control. Cell volume is not Ca\(^{2+}\)-
dependent per se, but increases in intracellular Ca\(^{2+}\)
have been shown to cause transient astrocyte swelling, thereby
activating VRACs. We used 100 \( \mu \)mol/L 5-nitro-2-(3-phe-
nylpropylamino)benzoic acid (NPPB) to block VRACs,
but no effect on HFO power of was observed (110 %;
107 ± 13%; \( P = 0.7296; n = 57; \) Fig. 8A). Thus, volume-
regulated channels are not likely to play a role in HFO
generation.

An increase in extracellular sodium
concentration boosts HFOs

We saw a very strong increase in HFO power when we
added extra 50 mmol/L NaCl to the bath solution
(150 mmol/L instead of 100 mmol/L NaCl: 1289 %;
1037 ± 218 %; \( P \leq 0.0001; n = 44; \) Fig. 9A). If sodium
was indeed the “moving ion” that evoked the HFO, it would
be reasonable that the bigger driving force for Na\(^+\) due to
the more positive equilibrium potential with increased Na\(^+\)
concentration would augment HFOs. But by raising the
concentration of NaCl we unintentionally shifted the Cl\(^-\)
equilibrium potential as well. However, if Cl\(^-\) was the
“moving ion”, we would have expected increased HFO
power after depolarization of the membrane potential. This
was not the case. Thus, we conclude that influx of Na\(^+\) and
not of Cl\(^-\) causes the extracellular voltage oscillations.

For electrophysiological studies Na\(^+\) can be replaced by
the ion N-methyl-D-glucamine (NMDG\(^+\)). NMDG\(^+\) shares
some chemical properties with Na\(^+\), but it is a much bigger
molecule and cannot enter the cell via sodium-specific
channels. When we added 50 mmol/L NMDG/HCl to the
recording solution we expected HFOs to have about the
same power as under control conditions. But interest-
ingly, HFOs were boosted to nearly the same level as
under 50 mmol/L NaCl extra (1106 %; 944 ± 259 %;
\( P \leq 0.0001; n = 46; \) Fig. 9A). Mann–Whitney test revealed
no significant difference between the NaCl- and the
NMDG/HCl-induced effect on HFO power \( (P = 0.2222). \)

This might suggest a rather unspecific effect due to the
increased osmolarity. But it might also be a hint that the
electrical stimulation opens channels in the membrane
with pores wide enough to let Na\(^+\) and big molecules like
NMDG\(^+\) pass in equal measure.

Indications for the involvement of pore-
forming P2X receptors

One proposed mechanism how big molecules like
NMDG\(^+\) enter astrocytes are membrane pores associated
with ionotropic purinergic P2X receptors. P2X2, P2X4, and P2X7-receptors have been shown to form pores after several seconds of ATP activation that are permeable to large cations (Jarvis and Khakh 2009). Although the mechanism is not completely understood, it is presumed that either the receptor itself or recruited pannexins form pores that are even wide enough to let fluorescent dyes enter the cell (Suadicani et al. 2006; Hamilton et al. 2008). We used pyridoxalphosphate-6-azophenyl-2',4'-disulfonic acid tetrasodium salt (PPADS) to block P2 receptors. HFO power dose-dependently decreased when PPADS was admitted to the astrocyte culture (1 μmol/L: 115%; 10 ± 13%; P = 0.2534; n = 61; 10 μmol/L: 48%; 42 ± 5%; P ≤ 0.0001; n = 61) (50 μmol/L: 23%; 20 ± 2%; P ≤ 0.0001; n = 61) (100 μmol/L: 16%; 15 ± 2%; P ≤ 0.0001; n = 61). These results point to an involvement of P2 receptors, but do not clarify which subtypes are relevant.

PPADS blocks the rat variant of the P2X4 receptor inefficiently with an IC50 of >100 μmol/L (Jones et al. 2000), the P2X7 receptor shows an intermediate sensitivity (53 μmol/L; Watano et al. 2002) and the P2X2 receptor the highest sensitivity (~IC50 = 6 μmol/L; García-Lecea et al. 1999). These values highlight P2X7 receptors as most likely candidates involved in HFO generation as 10 μmol/L PPADS in our experiments already induced a moderate inhibition.

Since pore-forming is thought to be induced by a permanent activation of the receptor with ATP, we elicited HFOs when ATP was applied to the recording solution. We tested increasing concentration of ATP (10 μmol/L, 25 μmol/L, 50 μmol/L, 100 μmol/L; 1 mmol/L), but only the highest concentrations showed a significant effect on HFO power (100 μmol/L: 153%; 150 ± 144%; P = 0.0002; n = 89; 1 mmol/L: 379%; 1076 ± 1858%; P ≤ 0.0001; Fig. 9B). These concentrations seem rather high, but in comparison to other P2X receptors, P2X7 receptors have a considerably lower affinity to ATP (EC50 = 100 μmol/L; Jarvis and Khakh 2009; North 2002; Sun et al. 2013). It is known that low concentrations of divalent cations facilitate the transport through P2X7 pores, which very well fits our observations in low-Ca2+ solution where HFOs were extremely powerful (Virginio et al. 1997; Nörenberg et al. 2011). P2X2 receptors are not blocked by Ca2+ and P2X4 receptors only at higher concentrations (IC50 = 87 μmol/L; Jarvis and Khakh 2009). Thus, the P2X7 receptor might be the best candidate for mediating this ion flux that manifests itself as HFOs in extracellular recordings.

**Acidic pH increases the power of HFOs**

A striking effect on HFOs was seen when the pH of the extracellular solution was changed: Acidification to a minimum pH of 6.6 strongly enhanced HFO power compared to a solution with a pH of 7.2 (279%; 194 ± 34%...
While an increased pH (up to 8.4) had the opposite effect (16 %; 19 ± 6 %; \( P \leq 0.0001; n = 51 \)), it seemed as if the low pH had sensitized the astrocytes for the agonist ATP.

The strengthening effect of ATP (100 \( \mu \text{mol/L} \)) on oscillation power was larger at an pH of 7.2 (169 %; 157 ± 33%; \( P \leq 0.0001; n = 43 \); Fig. 10B) compared to the ATP effect at an more basic pH of 7.7 (148%; 117 ± 21%; \( P = 0.0056; n = 43 \)).

**Discussion**

**High-frequency oscillations**

High-frequency oscillations of field potentials above the gamma band (100—200 Hz) have been recorded in hippocampal and cortical slices but also in the intact brain. These so-called ripples are thought to be field correlates of simultaneous population spike bursts synchronized by recurrent inhibition and have been associated with memory formation during sleep (Buzsáki and Silva 2012). Transient field potential oscillations at even higher frequencies (200–600 Hz) were attributed to pathological conditions, especially in seizure initiation. In a recent study, it was shown that incidence, frequency, and form of the ripples strongly depend on the extracellular calcium concentration. Hippocampal slices bathed in 3 mmol/L \( \text{Ca}^{2+} \) showed HFOs in the ripple range but turned to fast ripples when the \( \text{Ca}^{2+} \) was lowered to 1 mmol/L (Aivar et al. 2014). The authors claim that low \( \text{Ca}^{2+} \) enhances the intrinsic excitability of neurons by shifting the voltage-dependence of \( \text{Na}^{+} \) channels. This would increase the probability of neuronal bursting, which is a prerequisite for the emergence of fast ripples.

To our knowledge, the direct or indirect involvement of astrocytes in ripple generation has never been discussed. We know that astrocytic \( \text{Ca}^{2+} \) signaling can lead to neuronal depolarization and spontaneous epileptiform discharges (Kang et al. 2005). On the other hand, low external \( \text{Ca}^{2+} \) in hippocampal slices efficiently evokes \( \text{Ca}^{2+} \) oscillations in astrocytes followed by simultaneous \( \text{Ca}^{2+} \) elevations in adjacent neurons (Fellin et al. 2004). Thus, astrocytes might facilitate neuronal synchrony and epileptiform activity. The lack of a prominent frequency band in astrocytic oscillations described in our study raises doubt whether astrocytes are directly involved in generating ripple-like activity in the brain. Likewise, the observed strong local restriction of HFOs does not support the idea of astrocytes being key players in brain oscillatory activity. However, strong interactions between astrocytes and neurons on several functional levels have already been well established. By giving evidence for extracellular voltage fluctuations produced by astrocytes our study adds new colors to the palette of astrocytic potential to modulate and interfere with neuronal function. Obviously, experiments in mixed cultures will be a necessary next step to address the question how and to which extent these voltage fluctuations affect or are affected by neuronal activity.

**HFOs depend on activation of voltage-gated channels**

While the involvement of voltage-gated channels in the emergence of HFOs was clearly shown in our experiments.
with astrocytes cultures, it still raises some questions: In contrast to neurons, astrocytes are thought to be electrically passive. They show a rather linear voltage/current relationship due to the lack of any voltage-gated ion currents (Adermark and Lovinger 2008). However, the expression of voltage-activated channels has been demonstrated in cultured astrocytes (MacVicar 1984; Latour et al. 2003). Whether this is true for astrocytes in slices or in situ still remains under debate. Some authors claim that the presence of voltage-gated channels in non-neuronal brain cells determines a separate class of cells, but a proper nomenclature and classification has not yet been established. These cells are either called complex astrocytes (Adermark and Lovinger 2008), NG2 cells (polydendrocytes; Schools et al. 2003) or OPCs (oligodendrocyte precursor cells; Paez et al. 2009), since they are assumed to be progenitors of oligodendrocytes. OPCs express purinergic receptors in addition to voltage-gated channels, and they respond to glutamate and ATP with a rise of intracellular calcium (Hamilton et al. 2010; Zonouzi et al. 2011).

We cannot make a clear statement whether the cells in our experiments were glial precursor cells or astrocytes that up- or downregulated several genes due to the culturing process. Immunostainings revealed that the vast majority of the cells in our cultures were positive for GFAP (glial fibrillary acidic protein), which is not a feature of NG2 cells in vivo (Nishiyama et al. 2005) but common for astrocytes in vitro.

Strong GFAP expression can be also a sign of reactive astrogliaisis—a response to brain trauma, infection, ischemia, or neurodegenerative disorders where astrocytes strongly proliferate and undergo molecular and morphological changes. An upregulation of L-type Ca\(^{2+}\) channels was reported for astrocytes in several models of brain injury (Westenbroek et al. 1998; Chung et al. 2001; Xu et al. 2007; Willis et al. 2010; Rubio et al. 2013).

### HFOs are highly susceptible to changes in extracellular Ca\(^{2+}\)

Variation in extracellular calcium concentrations affects many signaling compartments, so we cannot directly infer a certain cellular mechanism from the strong calcium dependency. Based on previous studies and observations we can make some assumptions, although. Incubation of hippocampal slices in ACSF with low calcium is an established model of epilepsy. Some researchers claim that in this model seizure-like activity is due to excessive electric coupling, since calcium blocks the transport of molecules through gap junctions (Perez-Velazquez et al. 1994). Uncoupled gap junctions—so called hemichannels—that release or take up substances from the extracellular space are open when external calcium concentrations are low (Quist et al. 2000). Hales et al. (2012) attributed the strong effect of calcium on the observed HFOs in their experiments to the involvement of hemichannels. In addition, low calcium is known to elicit Ca\(^{2+}\) oscillations in astrocytes and to enhance the rise in intracellular Ca\(^{2+}\) induced by ATP, where it is unclear whether gap junctions are involved (Zanotti and Charles 1997). It was also discussed that low extracellular Ca\(^{2+}\) depolarizes the cell membrane possibly by altering the surface charge (Anderson et al. 1995), but membrane depolarization with high extracellular potassium concentrations failed to show the same promoting effect on HFOs as low Ca\(^{2+}\) in our experiments. Furthermore, the ion Ba\(^{2+}\) which blocks K\(^+\) channels—thus leading to a depolarization of the cell—did not enhance but significantly weakened HFOs, which nearly vanished when 1 mmol/L BaCl\(_2\) was added to the bath solution. Hence, it is not very likely that Ca\(^{2+}\) influences HFOs by depolarization of the cell membrane.

### HFOs do not depend on synaptic release

Several studies have used bafilomycin A1 to demonstrate vesicular release of gliotransmitters (Araque et al. 2000; Coco et al. 2003; Bowser and Khakh 2007; Gourine et al. 2010): Araque et al. showed that bafilomycin diminished glutamate release while Coco et al. saw the same effect on ATP release. However, in Coco et al. bafilomycin only partially inhibited the gliotransmitter release and it did not affect the initial calcium rise in the stimulated astrocytes. Bowser and Khakh observed that mechanical stimulation in bafilomycin-treated astrocytes induced a calcium rise in some adjacent astrocytes, but strongly suppressed further propagation of the calcium wave. We only saw a minor effect of bafilomycin on the oscillation power. Since previous publications have shown that vesicular release of gliotransmitters is relevant for the propagation of Ca\(^{2+}\) waves but not necessarily for their initiation, it is possible that HFOs reflect early steps in a signal cascade that may ultimately lead to propagating Ca\(^{2+}\) waves that can spread across a length of up to 200 µm. This explains why HFOs—if they represent early stages of a Ca\(^{2+}\) waves—were not detected by any of the neighboring electrodes at a distance of 200 µm.

### HFOs are strengthened by hyperpolarization and increase in extracellular Na\(^+\)

Due to a large resting conductance for potassium, astrocytes have a lower membrane potential than neurons (−85 mV vs. −65 mV; Magistretti and Ransom 2002). Thus, elevation of extracellular K\(^+\) effectively hyperpolarizes the membrane potential. Assuming an intracellular
potassium concentration of 120 mmol/L, lowering the extracellular potassium from 5 to 2 mmol/L K⁺ should – according to the Nernst equation – result in a drift of the membrane potential from approximately −80 to −100 mV. If HFOs were dependent on vesicular release, hyperpolarization of the membrane should rather inhibit HFO power. The observed increase in HFO power after membrane hyperpolarization thus suggests mechanisms other than vesicular release.

Hyperpolarization increases the driving force for ions with equilibrium potentials in the positive range. Thus, influx of Na⁺ and Ca²⁺, as well as efflux of anions like ATP³– and glutamate is facilitated. Due to the increased driving force for these ions after hyperpolarization, cross-membrane ion currents increase and can contribute to HFOs. On the other hand, K⁺ or Cl⁻ with strongly negative equilibrium potentials are unlikely HFO sources. Possible candidates for channels that could carry such currents are connexin-formed hemichannels, unspecific chloride channels like volume-regulated anionic channels or pore-forming P2X-receptors.

No indication for an involvement of hemichannels and volume-regulated anionic channels in HFO generation

Hemichannels were discussed as possible sites of glutamate efflux from astrocytes (Parpura et al. 2004; Ye et al. 2009). This transport is markedly blocked by divalent cations such as Ca²⁺, Mg²⁺, and Ba²⁺ in the extracellular solution. Ca²⁺ and Ba²⁺ were also very effective in blocking HFOs in our experiments. Carbenoxolone was used in several studies as blocker of gap junctions and hemichannels that eliminates the electrical coupling between neurons (Zsiros and Maccaferri 2005). Here, it partially inhibited the astrocytic HFO. However, recently the specificity of carbenoxolone was called into question (Beaumont and Maccaferri 2011). The VRAC blocker NPPB had no significant effect on HFO power in our study, giving no indication for VRAC involvement. Incidentally, NPPB was shown to act rather unspecifically as well (Cranmer et al. 1995; Ye et al. 2003).

Purinergic ATP receptors play a key role in HFO generation

ATP release from astrocytes after electrical stimulation was first described by Guthrie et al. in 1999; ATP is known to induce autocrine or paracrine ATP release (Anderson et al. 2004), initiating a propagating calcium wave in neighboring astrocytes (Guthrie et al. 1999; Bower and Khakh 2007) followed by further ATP release. In addition to vesicular and hemichannel-mechanisms, ATP release can occur via pores formed by purinergic P2X receptors (Suadicani et al. 2006; Hamilton et al. 2008). Several subtypes of the ionotropic P2X receptor were shown to form membrane pores after long stimulation with high ATP concentrations that are permeable for larger molecules such as NMDG⁺. We observed comparable increase in HFO power by equimolar amounts of Na⁺ or NMDG⁺, arguing for NMDG⁺ entry into the cell. ATP induces ion currents for instance through P2X7 receptors, but they are very small in normal medium containing Ca²⁺ and Mg²⁺. These currents can be greatly increased when the external medium contains a low concentration of these divalent cations (Suadicani et al. 2006; Oliveira et al. 2011). Ca²⁺ is thought to be a negative allosteric modulator of the P2X7R by decreasing the receptor’s affinity to its ligand ATP (Yan et al. 2011). This behavior fits our observations that low Ca²⁺ caused extremely powerful HFOs. The involvement of purinergic receptors in HFO generation was also demonstrated in our study by the dose-dependent inhibition of HFO power when treated with the unspecific P2 receptor blocker PPADS.

Acidic pH boosts HFO power

A decrease in pH is an important pathophysiological factor and can be measured in the brain after ischemia, hypoxia, or stroke. HFOs were dramatically strengthened when the pH of the extracellular solution was slightly changed to more acidic values. The opposite effect was seen when we induced HFOs at basic pH. It was already shown in 1997 that low extracellular pH raises intracellular calcium in astrocytes and therefore interacts with astrocytic Ca²⁺ signaling (Nagaoka 1997).

Single-channel recordings of P2X2 receptor showed that lowering extracellular pH enhanced the affinity for ATP and thereby increased the current through the channel (Ding and Sachs 1999). This fits our data very well, where the effect of ATP (100 μmol/L) on oscillation power grew stronger in an acidic bath solution, while our PPADS measurements rather suggested the P2X7 receptor subtype. We therefore propose that different subtypes of purinergic receptors – in combination with other ion channels – contribute to generating HFOs. P2X receptor interplay could also explain HFO strengthening under Ca²⁺ deficiency: P2X7 receptors that are blocked by Ca²⁺ show a slower desensitization than other receptors (Jones et al. 2000). Decreasing extracellular Ca²⁺ would then raise the amount of open P2X7 receptors with a large conductance that desensitize more slowly than other P2X subtypes that are not inhibited by Ca²⁺.

Our study shows that electrical stimulation triggers fast extracellular voltage oscillations in cultured astrocytes on
MEAs. We propose the following mechanism for these astrocytic high-frequency oscillations: The electrical stimulus leads to Ca$^{2+}$ influx through voltage-gated calcium channels, which mediates ATP release and subsequent activation of P2X autoreceptors. Persistent binding at the P2X receptor causes formation of membrane pores that facilitate influx of Na$^+$ and Ca$^{2+}$.

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**Conflict of Interest**

None declared.

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A voltage-operated Ca(2+)-channel (VOC) is a calcium channel that is activated by changes in the transmembrane potential of the cell membrane. This type of channel allows calcium ions to enter the cell, which can lead to a range of cellular responses, including the propagation of action potentials, the release of neurotransmitters, and the regulation of gene expression. VOCs are important for the proper functioning of excitable cells, such as neurons and muscle cells, and are involved in processes such as neurotransmitter release, synaptic plasticity, and cell proliferation.

The different types of VOCs include voltage-operated Na+ channels, voltage-operated K+ channels, and voltage-operated Ca(2+) channels. Each type of VOC has its own unique subunits and properties, which determine its sensitivity to changes in the membrane potential and the type of action potential it produces. For example, voltage-operated Na+ channels are sensitive to depolarization and cause a rapid increase in the membrane potential, while voltage-operated Ca(2+) channels are sensitive to changes in the membrane potential and are involved in the propagation of action potentials.

The regulation of VOCs is critical for the proper functioning of the nervous system. Dysregulation of VOCs can lead to a range of neurological disorders, such as epilepsy, Alzheimer’s disease, and Parkinson’s disease. Understanding the regulation of VOCs is therefore critical for the development of new treatments for these disorders.

In summary, VOCs are important for the proper functioning of the nervous system and play a critical role in the propagation of action potentials, neurotransmitter release, and synaptic plasticity. Understanding the regulation of VOCs is therefore critical for the development of new treatments for neurological disorders.
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